

## Characterization of anti-silencing factor 1 in *Leishmania major*

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*Anti-silencing factor 1 (ASF1) is a histone chaperone that contributes to the histone deposition during nucleosome assembly in newly replicated DNA. It is involved in chromatin disassembly, transcription activation and in the cellular response to DNA damage. In Leishmania major the ASF1 gene (LmASF1) is located in chromosome 20 and codes for a protein showing 67% of identity with the Trypanosoma brucei TbASF1a. Compared to orthologous proteins, LmASF1 conserves the main residues relevant for its various biological functions. To study ASF1 in Leishmania we generated a mutant overexpressing LmASF1 in L. major. We observed that the excess of LmASF1 impaired promastigotes growth rates and had no impact on cell cycle progress. Differently from yeast, ASF1 overproduction in Leishmania did not affect expression levels of genes located on telomeres, but led to an upregulation of proteins involved in chromatin remodelling and physiological stress, such as heat shock proteins, oxidoreductase activity and proteolysis. In addition, we observed that LmASF1 mutant is more susceptible to the DNA damaging agent, methyl methane sulphonate, than the control line. Therefore, our study suggests that ASF1 from Leishmania pertains to the chromatin remodelling machinery of the parasite and acts on its response to DNA damage.*

Key words: *Leishmania major* - anti-silencing factor 1 - histone chaperone

*Leishmania* spp is a member of the Trypanosomatidae family and a parasitic protozoa and digenetic eukaryote of medical and veterinary relevance. *Leishmania* undergoes several changes to adapt to and survive in different hostile environments, namely the insect vector digestive tract and the vertebrate host's macrophagic phagolysosomes. For that a tight and agile control of gene expression is required.

Differently from most of the eukaryotes, these parasites have polycistronic transcription and control of gene expression occurs primarily at the posttranscriptional level (reviewed by Clayton 2002, Martinez-Calvillo et al. 2010). Likewise other eukaryotes, in these parasites the chromatin structure modulates the access of proteins to the DNA, which ultimately regulates different aspects of gene expression, DNA processing, replication and double-stranded DNA repair (Navarro et al. 1999, Elias et al. 2001, Gontijo et al. 2003, McNairn & Gilbert 2003).

Anti-silencing factor 1 (ASF1) is a histone chaperone that together with chromatin assembly factor (CAF)-1 adds histones H3 and H4 onto newly replicated DNA (Tyler et al. 2001). This suggests a role for ASF1 in chromo-

matin assembly regulation (Loyola & Almouzni 2004, Zhang et al. 2005). ASF1 can be involved either in gene activation or transcriptional repression; it depends on the factors it interacts with (Sutton et al. 2001, Schwabish & Struhl 2006, Adkins et al. 2007, Takahata et al. 2009, Vary et al. 2010). In different eukaryotes deletion of ASF1 alters response to DNA damage and DNA replication blocking agents (Le et al. 1997, Tyler et al. 1999) and leads to gross chromosomal rearrangements (Prado et al. 2004) and even to cell death (Sanematsu et al. 2006).

The increased susceptibility of chromatin-assembly factor mutants to DNA-damaging agents may be an effect of the direct role of these factors in modulating chromatin structure (Mello et al. 2002). In addition, an association between ASF1 and genomic stability has been provided by the identification of a dynamic interaction between ASF1 and the Rad53 DNA damage checkpoint protein. Also, activation of ASF1 may be an important cellular response to DNA damage and replication stress (Emili et al. 2001, Hu et al. 2001). When DNA damage occurs, ASF1 is recruited to the lesion site where it may disrupt the H3/H4 tetramer. This results in nucleosome eviction and allows access of the repair machinery to unencumbered DNA (Canfield et al. 2009).

In *Leishmania* species ASF1 roles have not been investigated, although orthologous genes have been annotated in the sequenced genomes (tritrypdb.org). In another trypanosomatid, *Trypanosoma brucei*, the participation of ASF1 in the machinery that regulates spindle assembly and S-phase progression has been described (Li et al. 2007). Therefore, given that ASF1 is associated with gene expression in other eukaryotes and that chromatin modifications are relevant for controlling

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transcription in trypanosomatids (Green & Almouzni 2003, Figueiredo et al. 2009) we investigated *Leishmania major*'s ASF1. Our aim was to understand which of the known functions of ASF1 from other eukaryotes are shared by this parasite's protein.

## MATERIALS AND METHODS

**Parasites culture and transfection -** *L. major* MHOM/IR/83/IR (LmCC-1) promastigotes were grown at 26°C in M199 medium supplemented with 10% heat inactivated foetal calf serum, HEPES (0.04 M), adenine (0.1 mM), biotin (0.1 %), haemin (0.25 %), penicillin (2.5 U/mL<sup>-1</sup>), streptomycin (2.5 mg/mL<sup>-1</sup>). Transfection experiments were conducted as described elsewhere (Kapler et al. 1990). Mutants overexpressing ASF1, Lm/pX63NEO-ASF1 and Lm/pX63NEO, which bears the vector with no insert, were selected and cultured in the presence of G418. Increasing concentration of drug of selection (G418) was used to raise plasmid copy number and obtain higher levels of expression of episomal genes. G418 LD<sub>50</sub> was determined (1 µg.mL<sup>-1</sup>); we worked on a range of two-100 folds the LD<sub>50</sub>.

**Plasmid constructs -** A 1,246 bp fragment from pUCE8HH5.4, containing *LmASF1* ORF and its 5' and 3'UTR, was obtained after *Aat*II digestion. This plasmid was constructed with a 5.4 kb *Hind*III fragment subcloned from a cosmid, it has been described elsewhere (Pedrosa et al. 2001). pX63NEO-ASF1, which was used to overexpress the *LmASF1* in *L. major*, was obtained by insertion of the 1,246 bp fragment into pX63NEO *Aat*II-digested (LeBowitz et al. 1991).

**RNA extraction and northern blotting hybridization -** RNA from *L. major* in vitro cultures was extracted with Trizol® reagent following the manufacturer's protocol. Twenty micrograms of total RNA of each sample were fractionated on 2.2 M formaldehyde 1.2% agarose gel and transferred to a hybond-N+ nylon membrane. Subsequent steps for probe labelling and northern hybridization were conducted according to established protocols (Feinberg & Vogelstein 1983).

**Flow cytometry analysis and propidium iodide staining -** *L. major* control and ASF1 overexpressor cells were prepared according to Shapiro (Shapiro et al. 1984). The DNA content of PI-stained cells was analyzed with a FACScan analytical flow cytometer using the Cellquest software (BD Biosystems).

**Total protein cell extracts and two-dimensional gel electrophoresis (2D) -** Cells (1 x 10<sup>9</sup>) in late-log phase of axenic growth were pelleted (2,000 g/10 min/4°C), washed twice with 1x phosphate buffered saline (PBS) (pH 7.0) and resuspended in 1 mL 0.1x PBS containing protease inhibitors (Complete, Roche) for cell lysis. Cell extracts were submitted to five freeze-thaw cycles in liquid N<sub>2</sub>/water bath at 30°C and cellular debris were removed by two rounds of centrifugation (10,000 g/10 min, at 4°C). The supernatants were precipitated with 5 mL 0.1x PBS and trichloroacetic acid (TCA) at 4°C for 2 h. Precipitates were collected by centrifugation at 10,000 g followed by two washes with frozen acetone.

Proteins were solubilised in 300 µL of rehydration buffer [7 M urea, 2 M thiourea, 4% CHAPS (w/v)] plus nucleic solution (160 units of DNase and 125 µg RNase) and 10 mM MgCl<sub>2</sub> and incubated for 12 h at room temperature. Protein concentration was determined by the Bradford method (Bradford 1976).

Independent cultures of Lm/pX63NEO and Lm/pX63NEO-ASF1 were used for comparative analysis by 2D gel electrophoresis Immobiline DryStrip gels (13 cm: 4-7, GE-HealthCare) were rehydrated in 700 µg of parasite protein sample. Isoelectric focusing (IEF) was carried out at 20°C with maximum current setting at 50 µA per strip, using an active two-step rehydration system of 2 h at 0 V and of 10 h at 30 V, directly in ceramic strip holders, and then focused on an IPGphor IEF unit (GE HealthCare). Conditions for IEF were: 1 h at 500 V, 1 h at 1,000 V, 1 h at 2,000 V, 1 h at 4,000 and at 8,000 V until voltage reached 55,000 Vh. Focused pH gradient strips were then fractionated in sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) (12.5% polyacrylamide) according to the GE-HealthCare manual. Gel fixation and colloidal Coomassie staining were performed according to (Neuhoff et al. 1988). Gels were stored at 4°C in 25% ammonium sulfate and scanned with ImageScanner III (GE-HealthCare). Comparative analysis of digitized proteome maps was conducted with ImageMaster Platinum v7.0 software (GE-HealthCare). Differences higher than one-and-a-half-fold in the pattern and/or intensity of protein spots of analyzed strains were identified by pairwise comparison using gel images (Supplementary data). The proteins were identified by mass spectrometry using a MALDI TOF-TOF (4700 Proteomics Analyzer, Applied Biosystems).

**Real-time PCR (RT-PCR) - Relative quantification -** RT-PCR was performed using an ABI 7500 Sequence Detection System (Applied-Biosystems) in the presence of SYBR-green. The optimization of the RT-PCR reaction was performed according to the manufacturer's instructions (Applied-Biosystems, User Bulletin 2 applied to the SYBR-Green I core reagent protocol). The reactions were performed as technical and biological triplicates using SYBR Green (SYBRGreen PCR Master Mix, Applied Biosystems). Final concentration of oligonucleotides in the reaction was 200 nM and the identification of the 44 targeted transcripts and primers sequences are provided as Supplementary data. GLucose-6-phosphate dehydrogenase was used as the control of constitutive gene expression. To evaluate the efficiency of amplification reaction of each gene-target, a standard curve was generated using 10-fold dilutions of cDNA. The relative expression was analyzed by 2<sup>-ΔΔCT</sup> method.

**Production of polyclonal antiserum anti-ASF1 and western blotting analysis -** Rabbit serum was obtained by immunization with a recombinant polypeptide corresponding to the *T. brucei* ASF1 gene. This was PCR amplified using Asf-FowNdeI (5'-CATATGAGATA-CAACCAATT) and Asf-RevBamHI (5'-GGATCCT-CATCTGGGTTCAAGTGC) primers from the parasite genomic DNA and cloned in *Nde*I-*Bam*HI sites pET14b.

After induction with 0.5 mM IPTG for 12 h at 37°C, the recombinant protein was extracted with BugBuster Protein Extraction Reagent (Novagen) and purified by affinity chromatography on Ni-Agarose (Qiagen). Specific antibodies were purified with the same ASF1 used for the immunization. The recombinant protein was immobilized in a cyanogen bromide-agarose beads (GE Healthcare) and after washes with 10 column volumes it has been eluted with 0.1 M triethylamine, pH 11. The fractions were neutralized to pH 8 with Tris-HCl 1 M pH 7.4 and dialyzed against PBS.

For western blots, total protein was extracted from 1 x 10<sup>7</sup> parasites. Samples were fractionated in 12.5% SDS-PAGE, transferred to nitrocellulose membranes. Total proteins were visualized by staining with Ponceau-S and incubated with primary antibody in blocking solution. Antibody dilutions used were 1:500 to 1:1,000. The membrane was washed three times for 10 min with PBS, 0.1% Tween-20 and incubated with protein A 1:10,000 or anti-rabbit IgG (1:20,000) both conjugated to HRP (Invitrogen) for 45 min. The membrane was treated and bond antibodies developed following ECL kit instructions (ECL, Millipore).

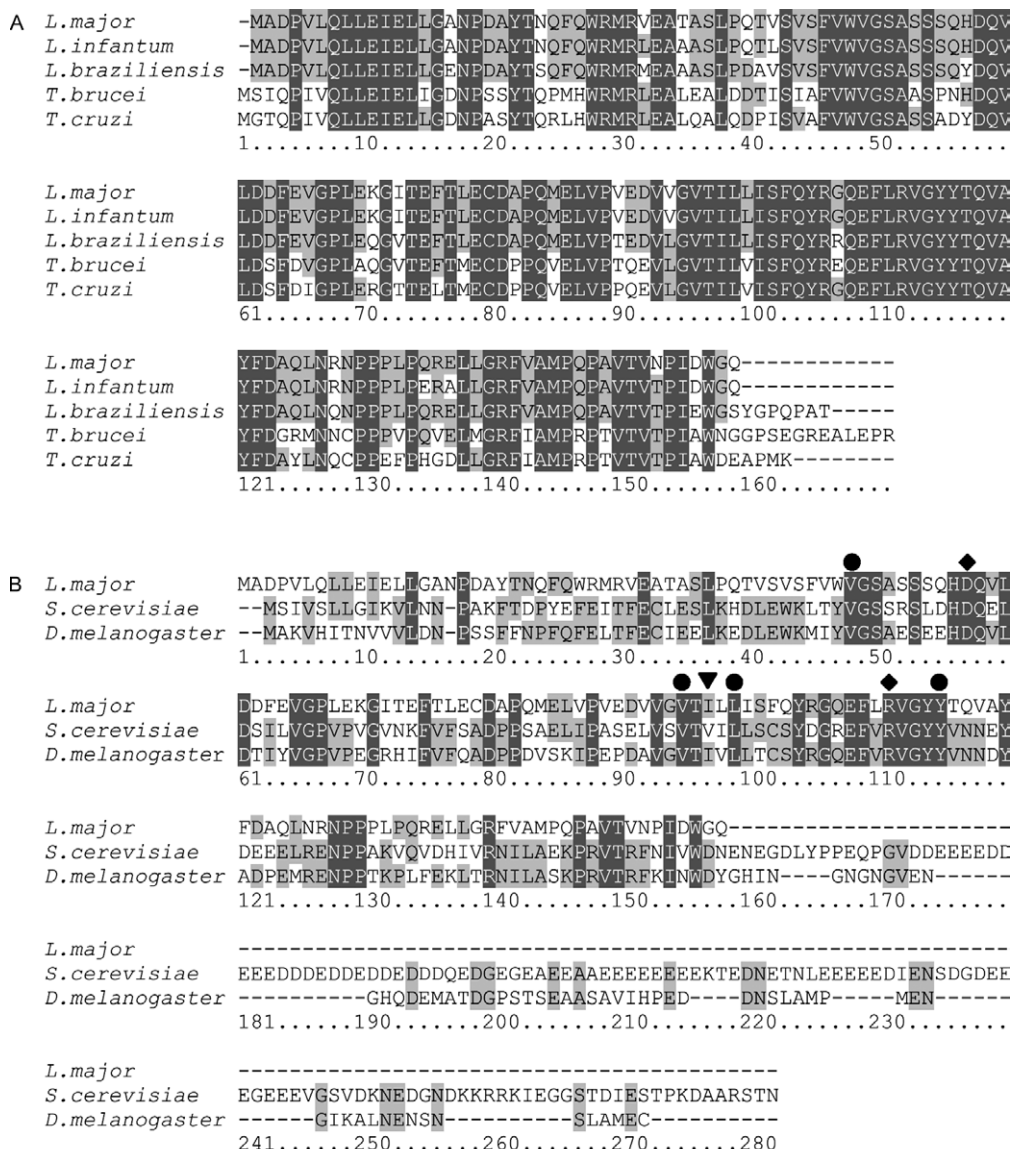


Fig. 1: conservation of anti-silencing factor 1 (ASF1) sequence. A: alignment of ASF1 from Trypanosomatidae showing high conservation in aminoacid residues (~67%); B: alignment of *Leishmania major*, *Saccharomyces cerevisiae* and *Drosophila melanogaster* ASF1 showing conserved residues of N-terminal region. Identical residues are shaded dark grey while conserved residues are shaded light grey. The symbol (●) means the amino acids V45, V92, L96 and Y112 and (◆) means the residues D54 and R108. The substitution V94I is indicated by the arrow head. The accessions of ASF1 sequences are: *L. major*, AAK28282.1, *Leishmania infantum*, CAM67383.1, *Leishmania braziliensis*, XP\_001564694.1, *Trypanosoma brucei*, CAJ15961.1, *Trypanosoma cruzi*, XM\_810961.1, *S. cerevisiae*, AAC37512.1 and *D. melanogaster*, AAF15354.1. The aminoacid positions correspond to *D. melanogaster* ASF1 sequence.

**Western blot signal quantification by ImageJ** - We selected the band indicated by “*LmASF1*” (at the right end of panel C) for quantification. ImageJ square tool was selected and used to determine the number of pixels per band using the tool named “plot profile”. The number of pixels obtained per band was subtracted from background pixels. The relative expression was calculated by dividing the value obtained for the signal detected in the transfectant by that observed in the control strain. Normalization of the values was conducted using the corresponding Ponceau-stained membrane.

## RESULTS

***L. major* ASF1 sequence conservation** - The ASF1 primary sequence is highly conserved throughout evolution, as it is shown by the in silico comparative analysis of the ASF1 homologues among trypanosomatids, *Saccharomyces cerevisiae* and *Drosophila melanogaster* (Fig. 1A, 1B). *L. major* ASF1 (LmjF20.0050) comprises 158 amino acids and displays 67% sequence identity with the *T. brucei* ASF1a (Tb927.1.630). The N-terminal domain is highly conserved between *Leishmania*, *S. cerevisiae* and *D. melanogaster*, but BLAST analysis revealed that the ASF1 C-terminal region from trypanosomatids lacks the long poly-acidic tract found in yeast (Fig. 1B).

**Effects of *LmASF1* overexpression over growth rate and cell cycle** - The detection of *LmASF1* transcript in promastigotes by northern blotting was only possible when at least 20 µg of total RNA was used, indicating that its endogenous level is low. We observed that the *LmASF1* transfectant (*Lm/pX63NEO-ASF1*) has higher levels of *LmASF1* transcript as compared to other transcripts. A positive correlation between drug concentration and increment of transcript levels was detected (Fig. 2A). The larger RNAs hybridizing with the ASF1 probe are probably read-through transcripts, commonly

found in episomal transcription in *Leishmania*. Relative expression of the *LmASF1* transcript in log phase promastigotes is 90.5 times higher in the overexpressor as compared to the control line (Fig. 2B). Interestingly, the protein level seeing as 18 KDa band detected by western blot using an anti-*T. brucei* ASF1 increased, but the increment did not seem so high (Fig. 2C). Image quantification revealed that ASF1 was at least 3.0 fold augmented in the overexpressor as compared to the control line *Lm/pX63NEO*. Quantification was conducted with the western band signals from cells kept at 100 x LD<sub>50</sub> to be comparable to the quantitative RT-PCR estimation of the transcript (Fig. 2C).

In addition overexpression of *LmASF1* decreased slightly the growth rate of promastigotes in axenic culture (Fig. 3A). The doubling time of the overexpressor mutant is 8.67 h as compared to 6.84 h of the control line. We did not witness significant changes of the cell cycle progression after submitting non-synchronized populations of ASF1 overexpressor as compared with the control transfectants (Fig. 3B).

**Changes in gene expression pattern in the *LmASF1* overexpressor** - Because the *S. cerevisiae*'s ASF1 is an ASF that releases the expression of genes located at the very end of the chromosomes, we investigated whether a similar role is played by its *L. major* orthologue, the *LmASF1*. We evaluated the transcript levels of 42 genes located in various chromosomes by comparing *Lm/pX63NEO-ASF1* and *Lm/pX63NEO*, most of them from the chromosomes' extremities (see distance of each gene from its telomeric end) (Table I). We found that expression of only five genes was affected by ASF1 overexpression (Table I). Among the few modulated genes, only one, the second annotated gene on chromosome 4 (LmjF04.0020), is less than 16 kb far from a telomere, as shown on Table I.

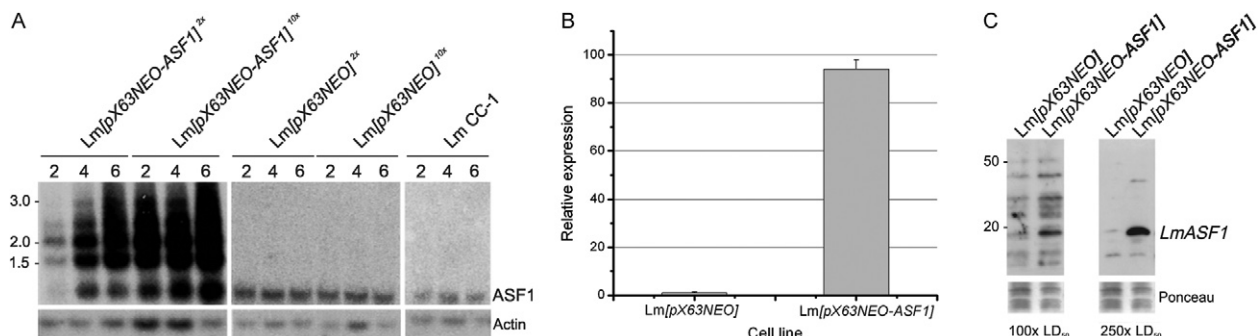


Fig. 2: anti-silencing factor 1 (ASF1) overexpression in *Leishmania major*. A: total RNA was extracted from promastigotes after two, four and six days in axenic culture (at the top of panel A: 2, 4 and 6, respectively) and 20 µg was fractionated in denaturing agarose gel, transferred to nylon membranes and hybridized with radiolabeled ASF1 probe. The left panel shows increasing amounts of *LmASF1* transcript in log phase overexpressor in two conditions of drug pressure (*Lm/pX63NEO-ASF1*<sup>2x</sup> and *Lm/pX63NEO-ASF1*<sup>10x</sup>) as compared to control lines (control transfectant *Lm/pX63NEO*<sup>2x</sup>, *Lm/pX63NEO*<sup>10x</sup> and *LmCC-1*) which are presented in mid and right panels, respectively. A fragment of actin gene was radiolabeled and used as a control of RNA loading (strip shown at the bottom of panels). Numbers shown in superscript mode refer to the drug pressure used in each experiment (times the LD<sub>50</sub> of G418); B: relative expression of *LmASF1* revealed by real-time polymerase chain reaction. The parasites were kept in media containing 100x LD<sub>50</sub> of G418; C: immunoblotting of cell extracts from promastigotes of *L. major* overexpressor, *Lm/pX63NEO-ASF1*, and control line *Lm/pX63NEO* in two conditions of drug pressure (100x LD<sub>50</sub>, left panel and 250x LD<sub>50</sub>, right panel) was probed with affinity purified antiserum to *Trypanosoma brucei* ASF1.

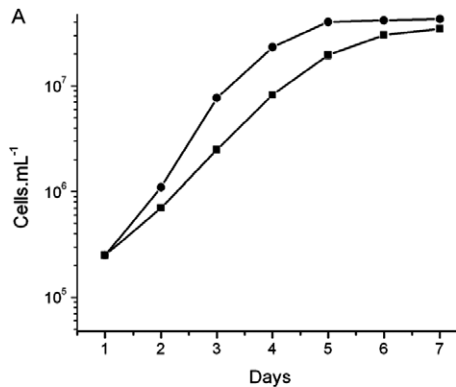


Fig. 3: growth and cell cycle of *LmASF1* overexpressor. A: growth curves of *Leishmania major* anti-silencing factor 1 (ASF1) over-expressor *Lm[pX63NEO-ASF1]<sup>100x</sup>* (■) and control transfectant *Lm[pX63NEO]<sup>100x</sup>* (●). Each point in the curves shown represents the mean and standard error from triplicates obtained from one representative experiment; B: FACS analysis of *L. major* ASF1 overexpressor (*Lm[pX63NEO-ASF1]<sup>100x</sup>*) and control line (*Lm[pX63NEO]<sup>100x</sup>*). Cells were rescued on early logarithmic phase of culture and stained with PI. The bars represent the averages of at least five independent experiments. The error bars indicate the standard deviation. The statistic test used was two-way analysis of variance (GraphPad Prism 4.00) and the variation between the cell lines was not significant. Numbers shown in superscript mode refer to the applied drug pressure (times the LD<sub>50</sub> of G418). Gray, white and black bars correspond to the cell cycle phases G1, S and G2, respectively.

We also evaluated the overall protein profile modifications triggered by the overexpression of ASF1 in the *L. major* ASF1 mutant. From three independent cultures of each cell line, protein was extracted and fractionated by two-dimensional gel electrophoresis and subsequently identified by mass spectrometry (Supplementary data). The number of identified spots in the 2D gel triplicates varied between 638-672 for the *Lm[pX63NEO]* and 725-990 for *Lm[pX63NEO-ASF1]*. We detected 19 proteins that were differentially expressed. Nine proteins were upregulated in the mutant and 10 were downregulated (Table II). Noteworthy, when we grouped the differentially expressed genes by Gene Ontology website terms (GO) (geneontology.org) we observed that six of them were either classified as heat-shock (HS), HS-related proteins or stress induced proteins (LmjF11.0350, LmjF26.1240, LmjF30.2460,

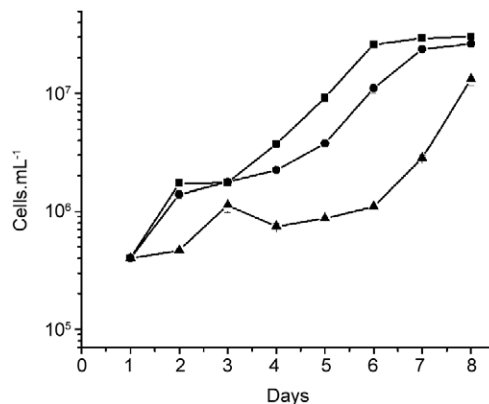


Fig. 4: growth impairment of *Leishmania* anti-silencing factor 1 (ASF1) overexpressors under methyl methane sulfonate (MMS) treatment. *Leishmania major* parental line (LmCC-1) (■), ASF1 overexpressors *Lm[pX63NEO-ASF1]<sup>100x</sup>* (●) and *Lm[pX63NEO-ASF1]<sup>50x</sup>* (▲) were maintained in the presence of MMS (0.005%) and monitored for cell growth for eight days. Numbers shown in superscript mode refer to the drug pressure used (times the LD<sub>50</sub> of G418). Each point in the curves shown represents the mean and standard error from triplicates obtained from one representative experiment.

Lmj28.2770, LmjF36.2030, LmjF08.1110), one protein is involved in proteolytic processes (LmjF33.2770), two have been identified as putative nucleosome assembly proteins (coded by genes LmjF29.2340 and LmjF19.0440) and three are involved in oxidoreductase activity (LmjF23.0040, LmjF23.0360, LmjF07.0430) (Table II). The other seven proteins identified were classified in diverse groups of GO categories (Table II).

**ASF1 mutant is oversensitive to methyl methane sulfonate (MMS)** - To verify whether ASF1 overexpression modifies the response to DNA damaging agents, we evaluated the parasites resistance to an alkylating agent. Parasites were submitted to sub-lethal concentration of MMS (0.005%) and promastigotes multiplication rates were estimated. There was a visible difference between the growth rate of parental cells and ASF1 overexpressor in the presence of MMS. In the *LmASF1* overexpressor the lag phase was extended for five days and only on the sixth day of culture it reached the logarithmic phase of growth, which in the parental line occurred on the third day (Fig. 4).

## DISCUSSION

The eukaryotic ASF1 protein is highly conserved throughout evolution (Daganzo et al. 2003, Mousson et al. 2005, Tamburini et al. 2005). Homologues of the *ScASF1* have been found in the genomes of trypanosomatids (tritrypdb.org) and the involvement of the *T. brucei* homologue, TbASF1a, in the control of spindle assembly and S-phase progression has been shown (Li et al. 2007). Nevertheless, no functional studies have been reported with *Leishmania* ASF1 homologues. This is the first study that presents an initial characterization of *L. major* ASF1.

*Leishmania* ASF1 amino acid sequence shows a high degree (~40%) of identity with homologues of a wide

variety of organisms, from yeast to man (Fig. 1) (ncbi.nlm.nih.gov/). In fact, the trypanosomatid's ASF1 corresponds to the 155-conserved amino-terminal sequence of budding yeast ASF1. Interestingly, we observed conservation among residues V45, V92, L96 and Y112 (Fig. 1B), which are the ones flanking the hydrophobic groove, a potential binding site for histones and other interacting proteins (Daganzo et al. 2003, Mousson et al. 2005). The central residue in this groove is located at position 94.

Whereas in yeast this residue is a valine in *Drosophila* and in trypanosomatids it is an isoleucine. The hydrophobicity of the region is maintained by polar and charged residues such as D54 and R108, which are conserved in trypanosomatids. We observed that in *LmASF1* all substitutions of residues that occurred in this groove and surrounding regions are conserved (Fig. 1). This region is crucial for the establishment of electrostatic interactions between the chaperone and the histone core (Mous-

TABLE I  
Transcript differential expression of 42 genes by real-time polymerase chain reaction

ID	Product	Distance of telomere (Kb)	Relative expression Lm[pX63NEO-ASF1]/Lm[pX63NEO]
LmjF01.0010	Hypothetical protein conserved	3.7	-
LmjF01.0020	Hypothetical protein unknown function	5.8	-
LmjF01.0810	Calcium/potassium channel, putative	24.75	-
LmjF01.0820 <sup>a</sup>	Potassium channel subunit-like protein	16.6	2.3x
LmjF02.0010	Phosphoglycan beta 1,3 galactosyltransferase 3	6.87	-
LmjF02.0180	Phosphoglycan beta 1,2 arabinosyltransferase	92.8	-
LmjF02.0190	Phosphoglycan beta 1,3 galactosyltransferase like-protein	97.1	-
LmjF03.0080	Glycerol 3 phosphate acyltransferase	18.8	-
LmjF04.0010	Calcium translocating P-type ATPase organelle-type calcium ATPase	4.6	-
LmjF04.0020 <sup>a</sup>	Pteridine transporter (truncated) putative	9.8	1.6x
LmjF04.0030	Camp-specific phosphodiesterase, putative	12.7	-
LmjF04.0060	DNA topoisomerase type IB small subunit	27.7	-
LmjF04.1160	Fructose 1,biphosphatase, cytosolic putative	43.9	-
LmjF04.1190	tRNA nucleotidyltransferase putative	18.3	-
LmjF04.1220	Hypothetical protein conserved	6.7	-
LmjF08.0080	ATP-dependent DEAD/H RNA helicase putative	21.0	-
LmjF09.0070	RNA helicase putative	17.2	-
LmjF11.0010	Hypothetical protein conserved	1.9	-
LmjF11.0020	Hypothetical protein conserved	2.9	-
LmjF11.0030	Hypothetical protein conserved	4.6	-
LmjF11.1360	Hypothetical protein conserved	10.6	-
LmjF11.1370	Hypothetical protein conserved	7.1	-
LmjF11.1380	Hypothetical protein conserved	4.0	-
LmjF18.0010	Hypothetical protein conserved	4.7	-
LmjF18.0020	Diphosphomevalonate decarboxylase, putative	7.9	-
LmjF18.0030	Hypothetical protein conserved	9.7	-
LmjF18.1660	Gamma-glutamylcysteine synthetase	10.0	-
LmjF18.1670	Hypothetical protein, unknown function	4.0	-
LmjF20.0020	Hypothetical protein conserved	7.5	-
LmjF20.0030	Hypothetical protein conserved	9.4	-
LmjF20.0040	Hypothetical protein conserved	13.2	-
LmjF20.0050 <sup>a</sup>	Anti-silencing protein 1-like protein	15.1	90.5x
LmjF20.0060	Hypothetical protein conserved	19.6	-
LmjF20.0070	Hypothetical protein conserved	21.3	-
LmjF20.0080 <sup>a</sup>	Hypothetical protein conserved	24.9	0.57x
LmjF20.0090 <sup>a</sup>	Hypothetical protein conserved	30.3	0.62x
LmjF20.160	Hypothetical protein conserved	3.2	-
LmjF22.1670	Hypothetical protein conserved	32.8	-
LmjF23.0290	Hypothetical protein conserved	108.2	-
LmjF23.0300	Tryptophanyl-tRNA synthase, putative	110.1	-
LmjF23.0310	Hypothetical protein conserved	112.8	-
LmjF23.0320 <sup>a</sup>	Hypothetical protein conserved	118.2	0.48x
LmjF23.0330	Hypothetical protein conserved	120.0	-

<sup>a</sup>: genes affected by anti-silencing protein 1 (ASF1) overexpression; ID: gene identification as annotated in the *Leishmania major* genome.

son et al. 2005). It has been shown (Daganzo et al. 2003) that this core domain is sufficient to confer resistance to DNA damage agents and to guarantee interaction of ASF1 with its partners, H3/H4 tetramer and Rad53. This conservation of the N-terminal region of *LmASF1* indicates that the hydrophobicity of this concave groove and its surrounding regions is maintained in *L. major*, which suggests that interaction of ASF1 with these partners is maintained in *Leishmania*.

We generated *Leishmania* parasites that produces higher amounts of ASF1 transcripts (90.5 times higher) and protein (at least 3 times higher) when compared to the control. The difference between ASF1 transcript and protein levels in the mutant is remarkable. This finding led us to speculate that this protein at high levels is deleterious to the parasite and that degradation of the excess of protein is necessary for the maintenance of cell homeostasis. Alternatively, the mRNA produced is degraded before being translated. These hypotheses are supported by other observations made in the course of this study. One of them is the fact that the ASF-over-expressing parasite presented a delayed growth, with a doubling time of 8.67 h compared to 6.84 h observed in the control line (Fig. 3A). Another possibility to explain the growth rate decrease is that cells have a delay in the S phase, a finding compatible with the fact that Asf1A is a substrate for Tousled-like kinases implicated in DNA replication control in *T. brucei* (Li et al. 2007). Nevertheless, we have not observed more than a subtle apparent increase on S-phase non synchronized cells, which was not confirmed in experimental triplicates (Fig. 3B).

ASF1 was originally identified as a disrupter of silencing of specific loci in *S. cerevisiae* (Le et al. 1997, Singer et al. 1998). In yeast, position-dependent gene silencing, including telomere silencing, involves many factors responsible for chromatin/nucleosome assembly, for instance ScAsf1p (Sharp et al. 2001). The fact that telomere proximal domains are less accessible to the transcriptional machinery, as compared to other chromosome regions, offers protection to these regions (Gottschling 1992). In several organisms, including *T. brucei*, this is achieved by repression of nearby genes (Levis et al. 1985, Gottschling et al. 1990, Zomerdijk et al. 1991, Horn & Cross 1995), with the control of chromatin packaging being essential for the strict control of the variant surface glycoproteins expression in these parasites (Rudenko 2010). To search for a potential role for *LmASF1* over telomere-proximal genes transcriptional control, we quantified the transcript level of 42 genes of *Leishmania*, mostly from the telomere proximal domains. We found that the augmented levels of ASF1 did not change the expression levels of any of the telomere-proximal genes analyzed (Table I), suggesting that differently than *S. cerevisiae*'s ASF1, the parasitic homologue is not involved in the expression control of telomeric genes. Nevertheless, we cannot rule out that a different factor plays this function in *Leishmania*.

Through proteomic analysis we investigated the potential changes that overexpression of ASF1 induced in gene expression. Interestingly, we observed changed levels of two proteins related to proteolytic activity (Table

II). These proteins are the ubiquitin conjugating enzyme (*Lmj33.2770*) and a metalloprotease (*LmjF05.0960*). In addition, there is a third protein that might be also involved in proteolysis; it is classified as an ATPase beta subunit protein (*LmjF25.1170*) that might be a member of the AAA ATPase family, which participates in a variety of cellular processes, including the shuttle of a subset of ubiquitinated substrates to the proteasome for degradation (Ye 2006). An indication that the ASF1 mutant was under stress comes from the overexpression of genes involved in physiological stress processes (*LmjF11.0350*, *LmjF26.1240*, *LmjF30.2460*, *Lmj28.2770*, *LmjF36.2030*, *LmjF08.1110*) (Table II, Supplementary data). Furthermore, the detection of altered levels of three proteins involved in the oxidoreductase activity (*LmjF23.0040*, *LmjF23.0360*, *LmjF07.0430*) suggests that cell redox metabolism is affected in the mutant.

Because ASF1 has a wide spectrum of functions related to chromatin metabolism, control of ASF1 levels must be critical for cell viability. ASF1 post-translational modifications were shown to be relevant for the control of its own cellular levels. It has been shown that phosphorylation of human and *Drosophila* ASF1 by tousled-like kinases (TLK) positively regulates protein levels, because it hampers the proteasome-dependent degradation of ASF1 (Pilyugin et al. 2009). In *T. brucei*, both ASF1 isoforms interact and are phosphorylated by TLK1 (Li et al. 2007). Currently, we have no information on the occurrence of post-translational modifications of *LmASF1*, nor whether or not it interacts with the *L. major* TLK. Assuming that the post-translational process happens similarly to that in *T. brucei*, we speculate that the endogenous LmTLK is unable to phosphorylate the excess of ASF1 produced by the mutant and that the unstable ASF1 is ubiquinated and shuttled to the proteasome to recover the organism homeostasis.

Finally, consistent with the *LmASF1* role in the control of chromatin dynamics, which interacts with other CAFs, we found in the comparative proteomic analysis (Table II, Supplementary data) that two of the proteins found in higher levels in the ASF1 mutant are involved in chromatin remodelling (*LmjF29.2340* and *LmjF19.0440*). Interaction of these proteins with ASF1 must be further investigated.

It has been shown that in other organisms ASF1 contribution to DNA repair is also linked to its effect on assembly/disassembly of chromatin, which is similar to what has been shown for its DNA replication or control of transcription (Mello et al. 2002, Gontijo et al. 2003, Green & Almouzni 2003, Canfield et al. 2009). The remarkable decrease in multiplication rates of ASF1 mutant in the presence of MMS indicates that this *LmASF1* isoform modifies the response to DNA damage. It is possible that high levels of ASF1 affect the chromatin structure, making the overexpressor's DNA more susceptible to the genotoxic effect of MMS. Alternatively, the delayed growth rate of the ASF1 overexpressor may be the result of the parasite attempt to maintain cellular homeostasis combined to the deleterious effects of the genotoxic agent.

TABLE II  
Identification of proteins differentially expressed in Lm/pX63NEO-ASF1/ as compared to Lm/pX63NEO/

MS ID	Differentially expressed proteins spot identification	Accession	GO ID	GO term name	Gene Ontology (GO)		Normalized volume <sup>d</sup>		
					theo/exp <sup>a</sup>	2nd D (MWkDa) theo/exp <sup>b</sup>	Mascot score <sup>c</sup>	Lm /pXN63NEO/	Lm /pX63NEO-ASF1/
25	No identified	-	-	-	?/6.4	?/16	-	0.133	-
28	MaoC family dehydratase-like protein from <i>Leishmania major</i>	LmjF07.0430	0008152	Metabolic process	5.1/5.1	16/17	137	0.055	0.109 (+1.55)
44	Hypothetical protein from <i>L. major</i>	LmjF29.2340	0006334	Nucleosome assembly	4.6/4.5	21/20	102	0.113	0.199 (+1.50)
54	Tryptaredoxin peroxidase from <i>L. major</i>	LmjF23.0040	0045454	Oxidoreductase activity	4.4/5.5	25/22	194	1.251	0.865 (-1.25)
73	Ubiquitin-conjugating enzyme from <i>L. major</i>	LmjF33.2770	0006508	Protein modification process/proteolysis	7.0/6.3	25/24	119	-	0.254
133	Chaperonin Hsp60, mitochondrial precursor from <i>L. major</i>	LmjF36.2030	0006457	Cellular protein metabolic process/protein folding	5.3/5.3	59/33	286	0.344	0.198 (-1.62)
175	NADP-dependent alcohol dehydrogenase from <i>L. major</i>	LmjF23.0360	0008152	Oxidoreductase activity	5.8/6.3	38/38	201	0.376	0.845 (+1.83)
194	Heat shock 70-related protein 1, mitochondrial from <i>L. major</i>	LmjF30.2460	0006457	Protein folding/unfolded protein binding	5.5/6.4	68/40	171	0.155	0.306 (+1.67)
222	Nucleosome assembly protein from <i>L. major</i>	LmjF19.0440	0006334	Nucleosome assembly	4.6/4.7	39/7/45	89	0.141	0.238 (+1.54)
252	ATPase beta subunit from <i>L. major</i>	LmjF25.1170	0015986	ATP synthesis coupled proton transport	5.1/5.1	56/50	157	0.040	0.079 (+1.56)
266	Dihydrolipoamide acetyltransferase precursor from <i>L. major</i>	LmjF36.2660	0008152	Acyltransferase activity/ protein binding	7.0/6.4	49/53	87	0.060	0.134 (+1.57)
300	Protein antigen LmSTII from <i>L. major</i>	LmjF08.1110	none	None	5.9/6.2	62/61	87	0.310	0.137 (-1.89)
356	Heat shock protein 70-related protein from <i>L. major</i>	LmjF26.1240	0005524	ATP binding	5.0/4.8	71/39	91	-	0.069
405	Alpha tubulin from <i>Leishmania infantum</i>	LmjF13.0280	0007017	Microtubule-based movement	6.8/5.5	36/21	100	0.075	-
407	Peroxidoxin; tryparedoxin peroxidase from <i>Leishmania</i>	LmjF23.0040	0045454	Oxidoreductase activity	6.4/5.3	25/22	83	0.136	-
411	“14-3-3 protein” from <i>L. major</i>	LmjF11.0350	0019904	Protein domain Specific binding	5.3/5.4	29-30/28	85	0.122	-
414	Hypothetical protein from <i>L. major</i>	LmjF25.2020	none	None	5.5/5.7	32/29	152	0.089	-



MS ID	Differentially expressed proteins spot identification	Gene Ontology (GO)				Normalized volume <sup>d</sup>			
		Accession	GO ID	GO term name	1st D (pI) theo/exp <sup>a</sup>	2nd D (MWkDa) theo/exp <sup>b</sup>	Mascot score <sup>c</sup>	Lm [pXN63NEO]	Lm [pX63NEO-ASF1] <sup>e</sup>
426	Activated protein kinase c receptor (LACK) from <i>L. major</i>	LmjF28.2740	0032465	Regulation of cytokinesis/ protein binding	6.0/6.4	34/36	234	0.077	-
478	Heat-shock protein 70 from <i>L. major</i>	LmjF28.2770	0005524	ATP binding	5.3/5.5	71-75/63	84	0.084	-

<sup>a</sup>: the theoretical (theo) pI value over the experimental (exp) pI found for each given protein; <sup>b</sup>: the theo molecular weight (MW) for each given protein over the encountered MW values in the experiment; <sup>c</sup>: quantification of the identification probability of the fragment match. A definition and interpretation of the meaning of the Mascot score can be found at the website matrixscience.com. We have included only statistically significant Mascot score results ( $p < 0.05$ ). Protein spots with statistically non significant Mascot scores remained unidentified; <sup>d</sup>: it corresponds to the amount of protein detected in each spot, in each strain. It is expressed as volume of the spot relative to the total volume of protein loaded in the gel. Therefore, the estimation of the difference between the volume of the spot in the compared strains is precisely expressed and allows to estimate the intensity of each signal comparatively in a given gel. When the spot was not detected in one strain a (-) signal indicates it; <sup>e</sup>: positive numbers (+) represent the fold difference of the upregulated proteins in the mutant strain and negative numbers (-) represent those proteins that are downregulated in the mutant, comparatively to the control strain; ASF1: anti-silencing protein 1; NADP: nicotinamide adenine dinucleotide phosphate.

Overall, our results indicate that the *Leishmania* ASF1 is not involved in controlling the expression of telomeric genes. On the other hand, altered levels of nucleosome assembly proteins (putative partners of ASF1) and the delayed growth rate of ASF1 overexpressor when submitted to a clastogenic agent indicate that ASF1 is involved in chromatin assembly/disassembly. Changed levels of proteins potentially involved in physiological stress conditions and in proteolytic processes, combined with the observed impaired growth rates, indicate that overexpression of ASF1 is deleterious to the parasite. Chromatin assembly dynamics is relevant for controlling transcription in trypanosomatids (Green & Almouzni 2003, Figueiredo et al. 2009) and is central for the DNA repair machinery. Investigating the roles and interactions of *Leishmania's* ASF1 is a route to improve understanding of how this important biological process may occur in trypanosomatids.

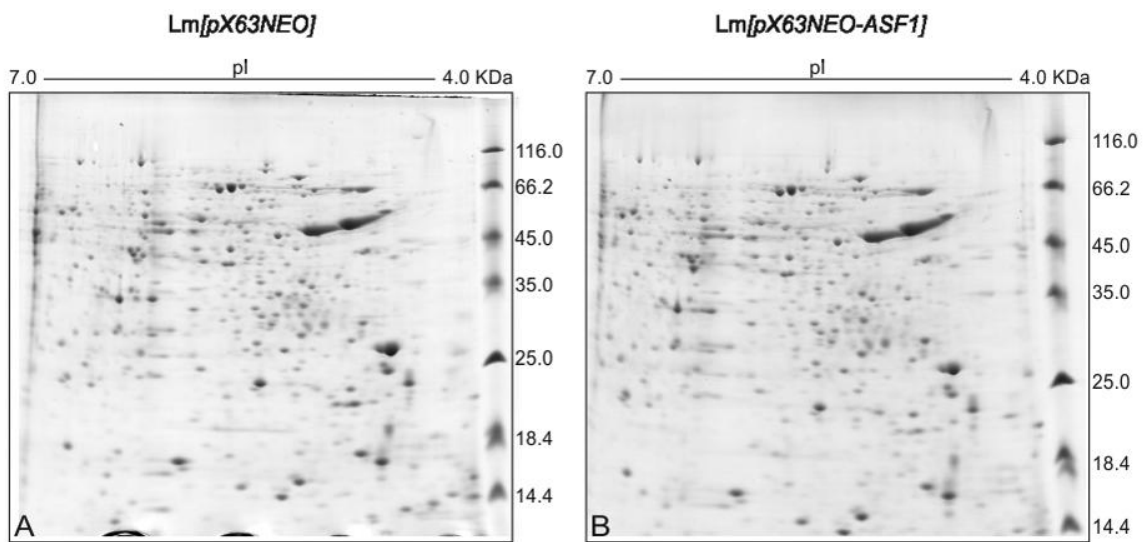
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Protein expression profiles of Lm[*pX63NEO*] (A) and Lm[*pX63NEO-ASF1*] (B). This panel represents a 2D gel electrophoretic fractionation of proteins from each cell lines of the three independent experiments. Equal amounts (700 µg) of proteins were fractionated on the 13 cm pI 4-7 linear gradient focused pH gradient strips and by 12% sodium dodecyl sulfate polyacrylamide gel electrophoresis.

Supplementary data

List of genes and primers used in the quantitative real-time polymerase chain reaction

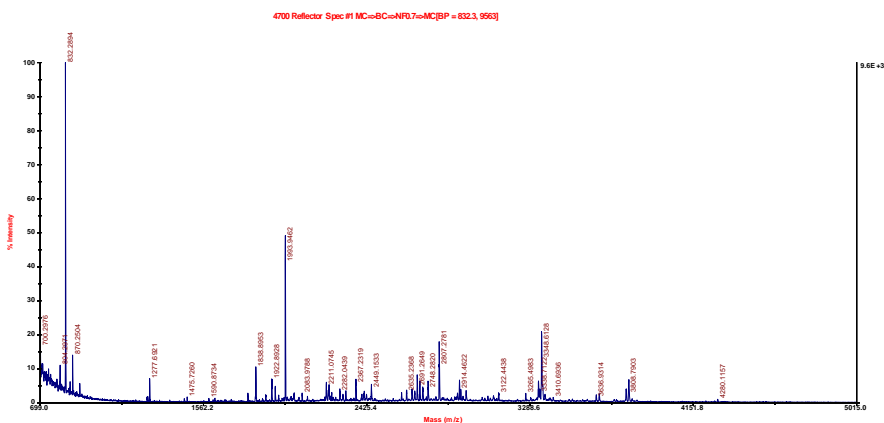
Oligonucleotide ID <sup>a</sup>	Gene ID	Sequence
LmjF01.0010 (R)	LmjF01.0010	ATGGGAACGTTCGTTGCACCTTCG
LmjF01.0010 (F)	LmjF01.0010	CACACGTACAAGTACTGCGGCAAA
LmjF01.0020 (R)	LmjF01.0020	GTTGTACGTGATGCCCTTCGCCCTT
LmjF01.0020 (F)	LmjF01.0020	GGCTCACAAGAACACAACAGCA
LmjF01.0810 (R)	LmjF01.0810	TCTGTGGCGAAGGCAGATAATAA
LmjF01.0810 (F)	LmjF01.0810	TGGTCCGCTACTTTATTCCGGCTGT
LmjF01.0820 (R)	LmjF01.0820	AGTAGCTGAAGCCGACACCAATGA
LmjF01.0820 (F)	LmjF01.0820	ACATGATGCTGCTGCTTCCTCTCGT
SCG3 (F)	LmjF02.0010	GCGCGTGTGCTACATGGC
SCG3 (R)	LmjF02.0010	AGAATCTGGTGCGGCCGG
LmjF02-SCA (F)	LmjF02.0180	CACCACTGCTGTGCCATACT
LmjF02-SCA (R)	LmjF02.0180	TATGTTGGTGCTCGTCTCGT
LmjF02.0190 (F)	LmjF02.0190	TAGGCCAGGATGAAGACACCG
LmjF02.0190 (R)	LmjF02.0190	CGCGGTGAAGGCTGGAAT
LbrM03-0090 (F)	LmjF03.0080	ATCGGTGGCATTATCAA
LbrM03.0090 (R)	LmjF03.0080	GAGTAGAGGAAGACGATGAACG
LmjF04.0010 (F)	LmjF04.0010	TCGACAAACAAGTGGCTATTGCG
LmjF04.0010 (R)	LmjF04.0010	AGTCCGTGAAATCAGTAGGCAGCA
LbrM04.0040 (F)	LmjF04.0020	AGGGGTGTGAGAGCACGATGT
LbrM04.0040 (R)	LmjF04.0020	AGGTTGCTGAAGTCGCAC
LmjF04.0030 (F)	LmjF04.0030	ATTTCAGAGCTAACGTCTCGC
LmjF04.0030 (R)	LmjF04.0030	TGTGTAGCGAGGCGAAGAAGGAAA
LmjF04.0060 (F)	LmjF04.0060	ACTCAACC CGC GAGGAGAACATCAT
LmjF04.0060 (R)	LmjF04.0060	TCTTCTGGATGGTGC GGAGAAGA
LmjF04.1160 (F)	LmjF04.1160	TGGCGACTTCACGCTCTTGATGAT
LmjF04.1160 (R)	LmjF04.1160	AGATCACATCCAGCTTCGCCTGAT
LmjF04.1190 (F)	LmjF04.1190	AATACAGCAGGCTACGAGCGACTT
LmjF04.1190 (R)	LmjF04.1190	AGCAGCGITGTTCCAACGTTCTTG
LmjF04.1220 (F)	LmjF04.1220	ACTCTTATGTAACACGCTGGCGA
LmjF04.1220 (R)	LmjF04.1220	AGTGCTGGAGACGTAAGCTTCT
LbrM08-0080 (F)	LmjF08.0080	TGGGCTACGCAAACCGC
LbrM08-0080 (R)	LmjF08.0080	ATATCCAACCGCGCTCTGC
LbrM09.0090 (F)	LmjF09.0070	TCGCACTGAGGTGGAGAATA
LbrM09.0090 (R)	LmjF09.0070	AGTGTGGCAGGTGACAAGTA
LmjF11.0010 (F)	LmjF11.0010	TGGCTGCTGCGAATAATCTCCTGA
LmjF11.0010 (R)	LmjF11.0010	GAGCATCTGCGAAAGCGGATTGT
LmjF11.0020 (F)	LmjF11.0020	AGATGTTCCCTCGACAAAGGGCT
LmjF11.0020 (R)	LmjF11.0020	AGACGATTGCAAGGCATAAACGGG
LmjF11.0030 (F)	LmjF11.0030	AGGCGATAGTCACGGGAAACACAA
LmjF11.0030 (R)	LmjF11.0030	AAACGAGCACGACGCAATATCAGC
LmjF11.1360 (F)	LmjF11.1360	TCGCGAGITGGACCCGTTATATT
LmjF11.1360 (R)	LmjF11.1360	CCAATTCTCTTGCGCTTTCGCCA
LmjF11.1370 (F)	LmjF11.1370	ACCTCAGCTGAAACGGCTTATCA
LmjF11.1370 (R)	LmjF11.1370	GAAAGCAAAGGCAGCCATGTCGAA
LmjF11.1380 (F)	LmjF11.1380	ACAAGAGAGCAGTACCGACAGCTT
LmjF11.1380 (R)	LmjF11.1380	TAACGGCGTTGCCTTCTTGATGC

LmjF18.0010 (F)	LmjF18.0010	AGATGCCACTCTGCTGAACGT
LmjF18.0010 (R)	LmjF18.0010	TGACGATAGAGTCACACGTTGCGT
LmjF18.0020 (F)	LmjF18.0020	CGCGAATTGCTTCATGTTACCCCT
LmjF18.0020 (R)	LmjF18.0020	TGCACCTCTCCAAGAGGTCTGCAT
LmjF18.0030 (F)	LmjF18.0030	TGCAGCGGTGTCATCTCCT
LmjF18.0030 (R)	LmjF18.0030	GTAGCCACGTCGAATTGCAAGCTTT
LmjF18.1660 (F)	LmjF18.1660	GTGACGATGCAAGCTGGACAACATT
LmjF18.1660 (R)	LmjF18.1660	TGTCGCAAAGAACGACCTCTGGA
LmjF18.1670 (F)	LmjF18.1670	TGTCGCAAAGAACGACCTCTGGA
LmjF18.1670 (R)	LmjF18.1670	AATTGATGGCTGTCCCTGGGACT
LmjF20.0020 (F)	LmjF20.0020	TGTTTCTTGCAGGAGATGGTATGC
LmjF20.0020 (R)	LmjF20.0020	TGATGGGAGAGTAAGCCAAAGCGT
LmjF20.0030 (F)	LmjF20.0030	ACCCATCTCGAGGCACAGTCAA
LmjF20.0030 (R)	LmjF20.0030	ACGGCGTAGGAAAGTACACAGCA
LmjF20.0040 (F)	LmjF20.0040	CAGCAAAGAGAGCAACTGGCTTGT
LmjF20.0040 (R)	LmjF20.0040	TTGGGCAAACCAGCACTGTCAAAG
LmjF20.0050 (F)	LmjF20.0050	AAACTGTGTCCGCTCCCTCGTGT
LmjF20.0050 (R)	LmjF20.0050	AGGTGAACTCAGTGATGCCCTTCT
LmjF20.0060 (F)	LmjF20.0060	TCATGCAGATGTACAACGCGGAGA
LmjF20.0060 (R)	LmjF20.0060	AGAGCTCATGTACAGCACCAACCAA
LmjF20.0070 (F)	LmjF20.0070	TGACTTCTCGAGACGACCATGA
LmjF20.0070 (R)	LmjF20.0070	CAGCACACCAATGAGAAAGGCAGT
LmjF20.0080 (F)	LmjF20.0080	AGATAACACCGCGGGAGATGAAGT
LmjF20.0080 (R)	LmjF20.0080	AGCGGATTGTTCTTCGACGCC
LmjF20.0090 (F)	LmjF20.0090	ACACTCGCCTTATGATGGTCGT
LmjF20.0090 (R)	LmjF20.0090	AGCTTGAAGAGCGCTGTGACTGT
LmjF20.1620 (F)	LmjF20.1620	AGTGGTCTTGTAAATCAGGCTGCA
LmjF20.1620 (R)	LmjF20.1620	TGGTACACTTCAGCTCGCCTTCAT
LbrM22.1580 (F)	LmjF22.1670	AGGATAGGATGGCGGAGATT
LbrM22.1580 (R)	LmjF22.1670	TCAAGTTGTTGGCGAGAAG
LmjF23.0290 (F)	LmjF23.0290	AACTGAAGAACACGGCAGCCGATA
LmjF23.0290 (R)	LmjF23.0290	TGAAGTGCTCGCGTAGATGAGAA
LmjF23.0300 (F)	LmjF23.0300	AGCTGATTGCAAGAACATCAAGG
LmjF23.0300 (R)	LmjF23.0300	CAAGCTGCGACATTATCGCTGT
LmjF23.0310 (F)	LmjF23.0310	CAATCGCGATCTTCAGCAGCACAA
LmjF23.0310 (R)	LmjF23.0310	AGCGTACAAGATGCGAGCCATAGT
LmjF23.0320 (F)	LmjF23.0320	TGCTGCGTACAAGTCCGACACTAA
LmjF23.0320 (R)	LmjF23.0320	TCACCGCTTTGCTGTCGCTGTAA
LT 197	LmjF23.0330	AGGTGGTTGTTGGCTTCAAC
LT 198	LmjF23.0330	AACAAACGACATGGCGAAGACCAC
Lm G6PDH (F)	LmjF34.0080	ACCGCATTGACCAACTACCTC
Lm G6PDH (R)	LmjF34.0080	GATGTTGTTCGAGTTCCAC

*a:* some primers were designed to amplify the corresponding gene fragment from the *Leishmania braziliensis*.  
ID: ?; R: reverse; F: forward.

Supplementary data  
Mass spectrometry of spots obtained from 2D gels. Identification of the proteins are  
presented in Table II

Sample: 25  
Mass Spectrum (positive reflector)

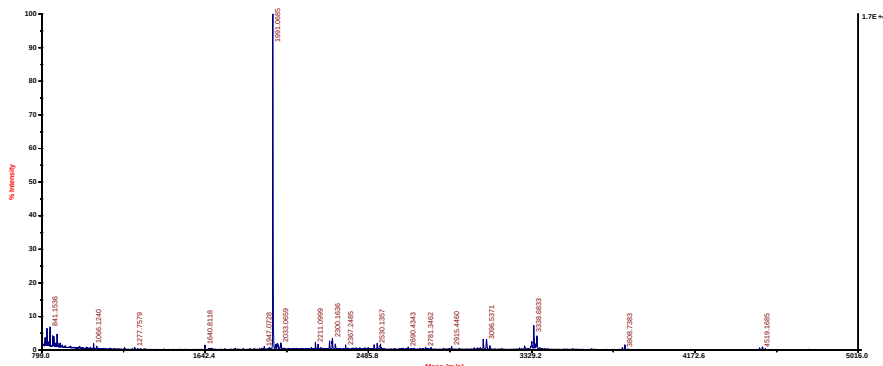


### Conclusion:

- The spectrum shows only m/z values corresponding to keratin tryptic peptides and trypsin autolytic fragments.
- With this data could **not identify** in a significant manner any protein in **sample 25**.

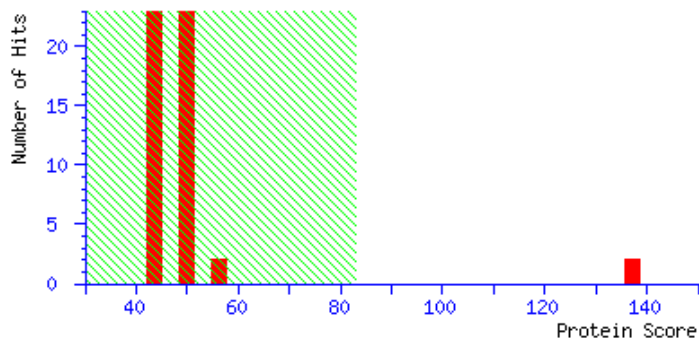
Sample: 28  
Mass Spectrum (positive reflector)

4700 Reflector Spec #1 MC=>BC=>NP0.7=>MC [BP = 1992.1, 16887]



## Mascot Score Histogram

Protein score is  $-10 \times \log(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p < 0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Match to: [gi|157864476](#) Score: 137 Expect: 2.2e-07 **maoc** family dehydratase-like protein [Leishmania major] Nominal mass ( $M_r$ ): **16974**; Calculated pI value: **5.14** NCBI BLAST search of [gi|157864476](#) against nr Unformatted sequence string for pasting into other applications Taxonomy: [Leishmania major strain Friedlin](#) Links to retrieve other entries containing this sequence from NCBI Entrez: [gi|68124241](#) from [Leishmania major strain Friedlin](#) Variable modifications: Carbamidomethyl (C),Oxidation (M) Cleavage by Trypsin: cuts C-term side of KR unless next residue is P Sequence Coverage: **35%**

Matched peptides shown in **Bold**:

1 MSSETAPI·VI RVGSRATRTV HISQENVNT GRVIDDHNPI HSDPEAARAA

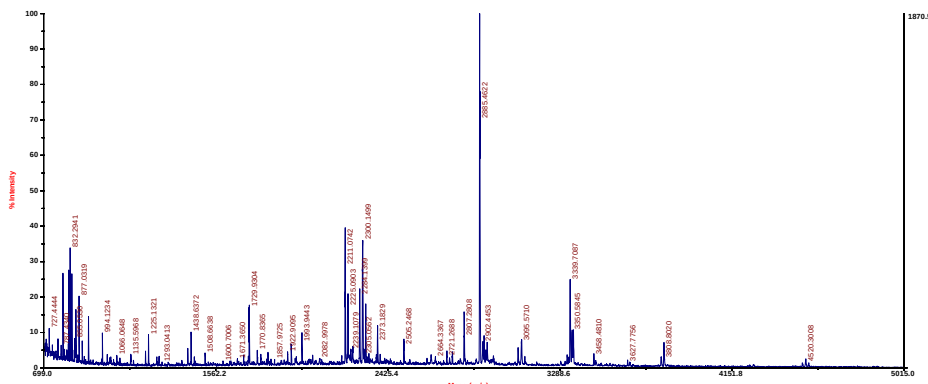
**51 GEPTTICYGM YAGSLFSSGLM AKEMPGPGTV YLSONLRTA PVFVGDPPLDV**

**101 IVEVREFRKD KGLISLSNIV QKTDPTGKT TVCVEGSAVV MNKTLQFEGE**

151 SEWSVTK

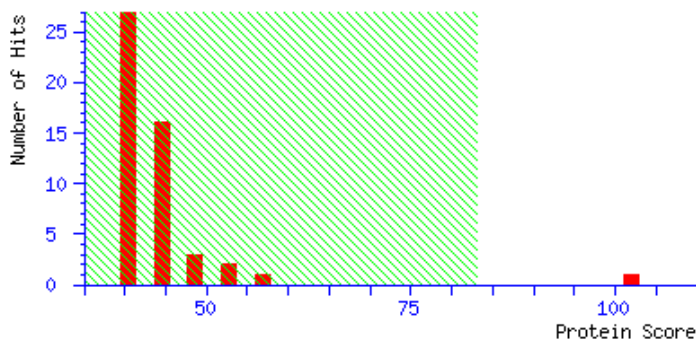
Sample: 44  
 Mass Spectrum (positive reflector)

4700 Reflector Spec #1 MC=>BC=>NP0.7=>MC [BP = 2886.5, 1871]



### Mascot Score Histogram

Protein score is  $-10 \times \text{Log}(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p < 0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.

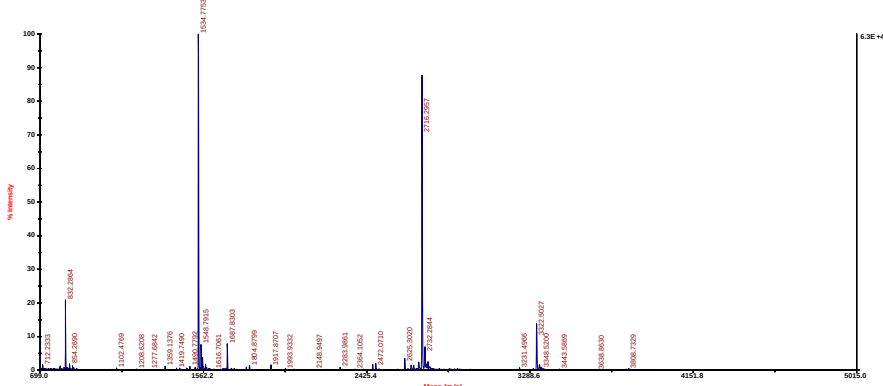


Match to: [gi|73536961](#) Score: 102 Expect: 0.0007 hypothetical protein [Leishmania major strain Friedlin] Nominal mass ( $M_r$ ): 21157; Calculated pI value: 4.61 NCBI BLAST search of [gi|73536961](#) against nr Unformatted sequence string for pasting into other applications Taxonomy: [Leishmania major strain Friedlin](#) Links to retrieve other entries containing this sequence from NCBI Entrez: [gi|70799775](#) from [Leishmania major strain Friedlin](#) Variable modifications: Oxidation (M) Cleavage by Trypsin: cuts C-term side of KR unless next residue is P Sequence Coverage: 40% Matched peptides shown in **Bold**:

**1** MELSQCATQA ISKFTDDLES IDRKCEEETT ALRLEYRSKM EPLL MERREL  
**51** LKGVTNFWSG VLSSPETPIA ALLNGTIDPK IIRAVTDFQV VTRISENKLY  
**101** RKIIFTFRQN MFVEEGEISR EVDTDMETTS LHPVKWKGST ERARTDSFFA  
**151** FFTENFSSDP EDVMEVTEAL DTIYQNPFLA VEAD

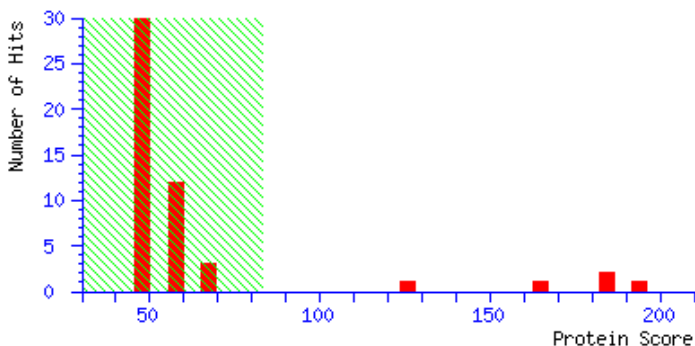
Sample: 54  
Mass Spectrum (positive reflector)

4700 Reflector Spec #1 MC=>BC=>NF0.7=>MC[BP = 1534.8, 62637]



## Mascot Score Histogram

Protein score is  $-10 \cdot \log(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p < 0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



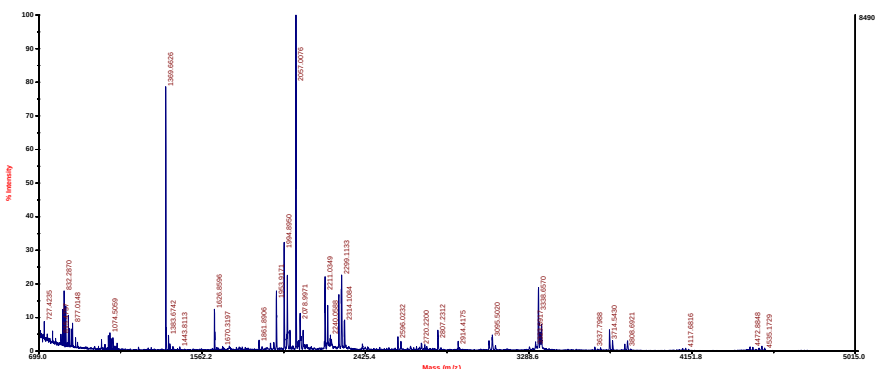
Match to: [gi|157869550](#) Score: **194** Expect: **4.4e-13** peroxidoxin; tryparedoxin peroxidase [Leishmania major] Nominal mass ( $M_r$ ): **25326**; Calculated pI value: **6.43** NCBI BLAST search of [gi|157869550](#) against nr Unformatted sequence string for pasting into other applications Taxonomy: [Leishmania major strain Friedlin](#) Links to retrieve other entries containing this sequence from NCBI Entrez: [gi|68126391](#) from [Leishmania major strain Friedlin](#) Variable modifications: Oxidation (M),Carbamidomethyl (C) Cleavage by Trypsin: cuts C-term side of KR unless next residue is P Sequence Coverage: **52%**

Matched peptides shown in **Bold**:

1 MLRRRLSTSSF LKRAQFRGFA ATSPLLNLDY QMYRTATVRE AAPQFSGQAV  
51 VNGAIKDINM NDYKGKYIVL FFYPMDFTV CPTEIIAFSD RHADFEKLNT  
101 QVVAVSCDSV YSHLAWVNTP RKKGGLGEMH IPVLADKSME IARDYGVLIE  
151 ESGIALRGLF IIDKKGILRH STINDLPVGR NVDEALRVLE AFQYADENGD  
201 AIPCGWKPGQ PTLDTTKAEE FFEKNM

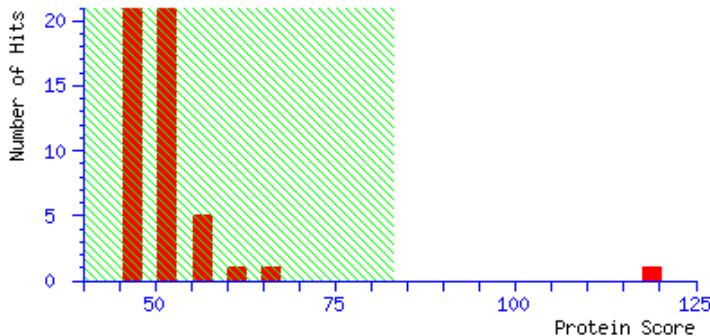
Sample: 73  
 Mass Spectrum (positive reflector)

4700 Reflector Spec #1 MC=>BC=>NP0.7=>MC|BP = 2050.0, 8490|



### Mascot Score Histogram

Protein score is  $-10 \times \text{Log}(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p < 0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.

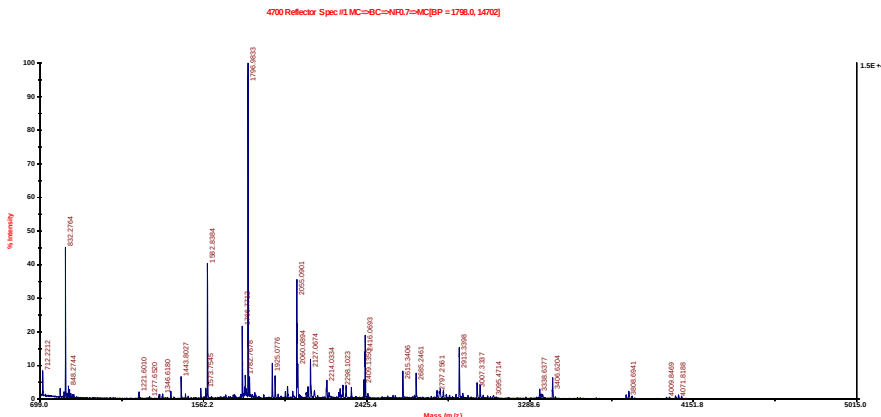


Match to: [gi|157875282](#) Score: 119 Expect: 1.4e-05 ubiquitin-conjugating enzyme [Leishmania major strain Friedlin] Nominal mass ( $M_r$ ): 25934; Calculated pI value: 7.01 NCBI BLAST search of [gi|157875282](#) against nr Unformatted [sequence string](#) for pasting into other applications Taxonomy: [Leishmania major strain Friedlin](#) Links to retrieve other entries containing this sequence from NCBI Entrez: [gi|68129113](#) from [Leishmania major strain Friedlin](#) Variable modifications: Oxidation (M), Carbamidomethyl (C) Cleavage by Trypsin: cuts C-term side of KR unless next residue is P Sequence Coverage: 22%

Matched peptides shown in **Bold**:

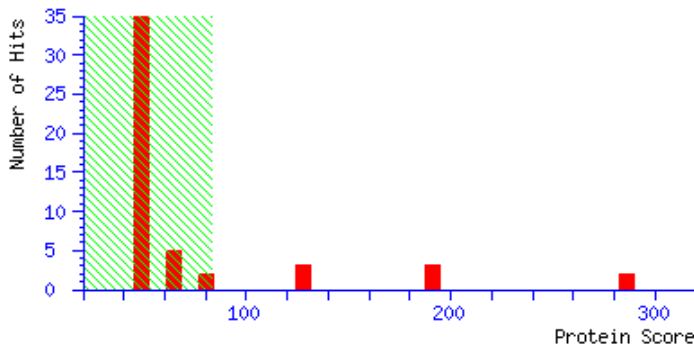
**1** MPLTAAGMRT LMRQMQEVE HPVDGVQVRP SDSMNEYHFD VDGPEGTPFA  
**51** AGRFHVVLIF **DEQYYPEVPPK** GFFRTKIFHP NISERGDICV NTLKRDWSPM  
**101** LGLRHILVVI RCLLIEPNPE SALNEEAGRL **I**LEYYAAYER KAAMYTAINA  
**151** ARPNGVPRFT LPEVKAEKAS GSVTACASAG AAADSGASAT AAAAAASASS  
**201** MANGGAPSTA TRGLTLSEAN CHPGSATAAE KAEKSKKKAL RRI

Sample: 133  
Mass Spectrum (positive reflector)



### Mascot Score Histogram

Protein score is  $-10 \cdot \log(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p < 0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.

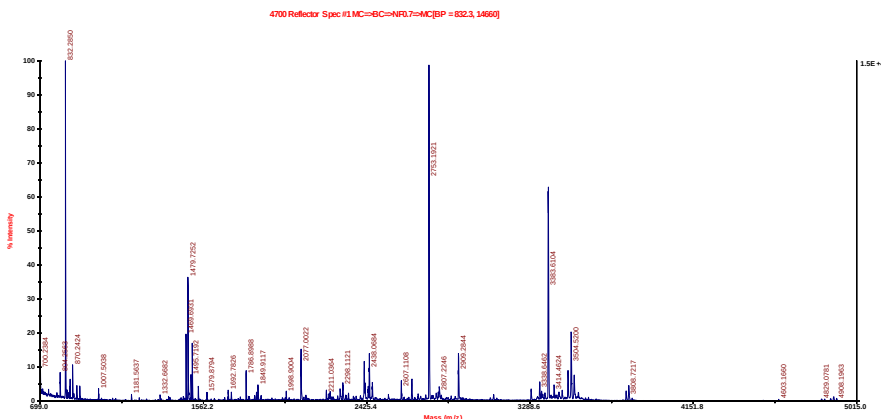


Match to: [gi|157876872](#) Score: **286** Expect: **2.8e-22** chaperonin Hsp60, mitochondrial precursor [Leishmania major] Nominal mass ( $M_p$ ): **59281**; Calculated pI value: **5.33** NCBI BLAST search of [gi|157876872](#) against nr Unformatted sequence string for pasting into other applications Taxonomy: [Leishmania major strain Friedlin](#) Links to retrieve other entries containing this sequence from NCBI Entrez:[gi|68129853](#) from [Leishmania major strain Friedlin](#) Variable modifications: Oxidation (M),Carbamidomethyl (C) Cleavage by Trypsin: cuts C-term side of KR unless next residue is P Sequence Coverage: **24%**

Matched peptides shown in **Bold** :

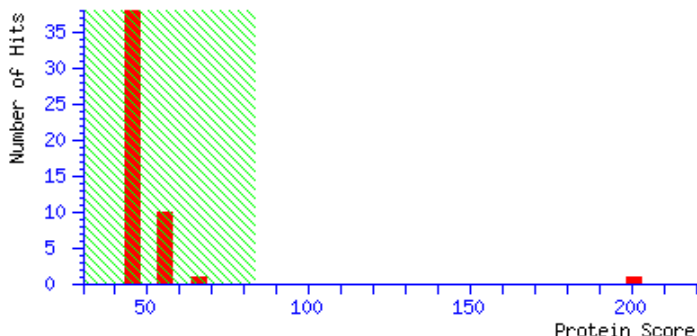
**1 MLRSAVCLAG KDVRFSDKAR RSMQKGVTRA VAAVATTLGP KGRNVIIIEQA**  
**51 YGAPKITKDG VTVAKAIEFK DRFENMGAQL VRQVCNKTND LAGDGTTS**  
**101 VLVDSIFGEG LKCIAQGTNP IDMKGMDRA VEVILKSVAK QSRPIKGSED**  
**151 IVQVATISAN GDKEIGEMIG QAMDKVGRDG VITAQDGKTM ATELEVVEGM**  
**201 SVDRGYVSPY FVTDAKAQKA ELEDALVLMS AKKIQSIIHSL LPALNHVVRS**  
**251 GRPLLIADD VESEALTTLI FNKLQGKLKV CCVKAPGFGD NKTAMLQDMS**  
**301 VFTGAQLVGD EGTGLELDAE NFDPSILGSV KKVTVTKDDT VMLNGGGDAA**  
**351 AVKERVELLR ELIKNETSDY NRDKLKERLA KLSGGVAVIK VGGGSEVEVN**  
**401 EKKDRIEDAL CSTRAAVQEG IVAGGGTALL RASKELEALA NDGSLTRDQR**  
**451 TGVTVRNAI RLPAMTIAAN AGKEGAVIVE KVLEASEEST GYDAQNDKYV**  
**501 NMFEAGIIDP TRVVRAVAISD ATSVASLMMT AEAAIVELPK DDSAAAPMGD**  
**551 MGGMGGMGGM GF**

Sample: 175  
 Mass Spectrum (positive reflector)



### Mascot Score Histogram

Protein score is  $-10 \times \text{Log}(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p < 0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.

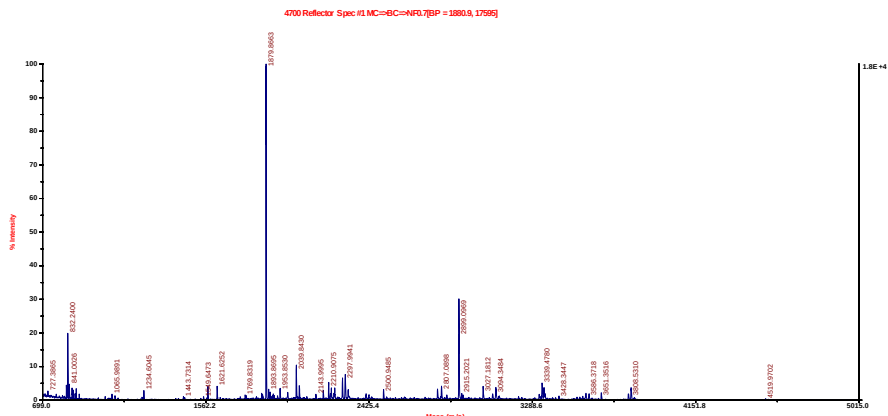


Match to: gi|157869618 Score: 201 Expect: 8.8e-14 NADP-dependent alcohol dehydrogenase [Leishmania major strain Friedlin] Nominal mass ( $M_r$ ): 38345; Calculated pI value: 5.78 NCBI BLAST search of gi|157869618 against nr Unformatted sequence string for pasting into other applications Taxonomy: Leishmania major strain Friedlin Links to retrieve other entries containing this sequence from NCBI Entrez:gi|68126425 from Leishmania major strain Friedlin Variable modifications: Oxidation (M), Carbamidomethyl (C) Cleavage by Trypsin: cuts C-term side of KR unless next residue is P Sequence Coverage: 41%

Matched peptides shown in **Bold**:

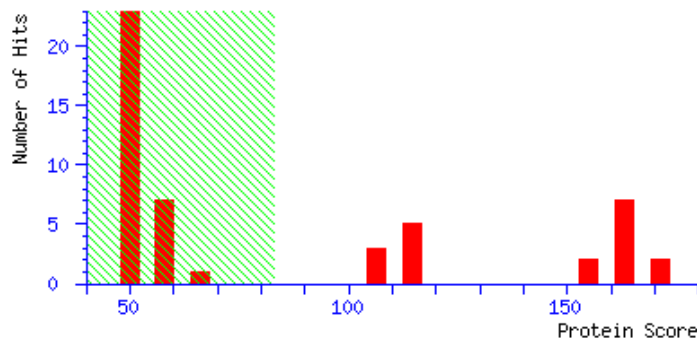
**1** MPTEAHGWAA LSAKSKLEPF AFQRRDVGPD DVVIAIAYCG VCHSDVHQAR  
**51** DEWDGSTFPM VPGHEIVGHV TKVGSEVTKY KAGDRVGVGC MVDSCMKCRQ  
**101** CERGLEQYCV NGASFTYNST QQDKKKPTFG GYSDHVVVRE HFVVSIPDNL  
**151** DLCAAAPLLC AGVTTFSPLR YWGVKKGTRV GVGVLGLGH MAVKLANAMG  
**201** AEVTVFTRSS NKVEEAENLG AHVVNTNNE QEMNSIQGTL DVIVDTVGMS  
**251** HDLRPYMMTL DIDGKLALVG MPEHAHPPLD PRRIIASRQC VGGSNIAAGMP  
**301** ETQELLNFCG EHNITATVEK IGIEYINEAY ERILASDVRY RFVIDMASLK  
**351** KE

Sample: 194  
 Mass Spectrum (positive reflector)



### Mascot Score Histogram

Protein score is  $-10 \times \log(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p < 0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



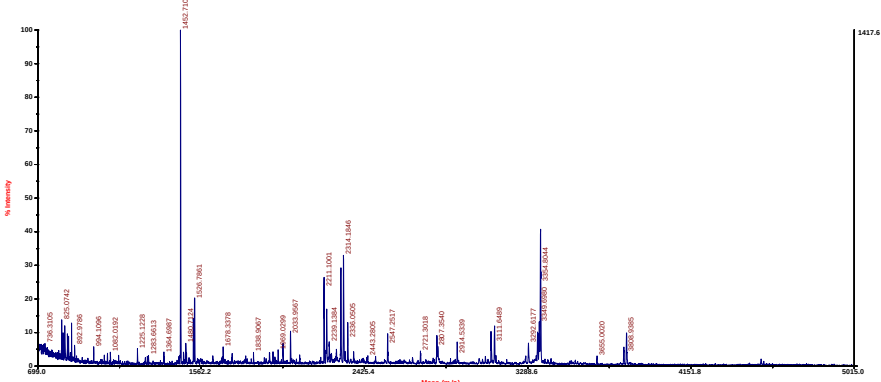
Match to: gi|57015345 Score: 171 Expect: 8.8e-11 RecName: Full=Heat shock 70-related protein 1, mitochondrial; Flags: Precursor Nominal mass ( $M_r$ ): 68288; Calculated pI value: 5.52 NCBI BLAST search of gi|57015345 against nr Unformatted sequence string for pasting into other applications Taxonomy: Leishmania major Links to retrieve other entries containing this sequence from NCBI Entrez: gi|50299857 from Leishmania major Variable modifications: Oxidation (M), Carbamidomethyl (C) Cleavage by Trypsin: cuts C-term side of KR unless next residue is PSequence Coverage: 18%

Matched peptides shown in **Bold**:

**1** MFARRVCGSA AASAACLARH ESQKVQGDVI **GVDLGTTYS**C VATMDGDKAR  
**51** VLENSEGFRTPSVVAFKGS EKLVGLAAKR QAITNPQSTF YAVKRLIGRR  
**101** FEDEHIQKDI KNVPYKIVRA GNGDAWVQDG NGKQYSPSQI GAFVLEKMKE  
**151** TAENFLGHKV SNAVVTCPAY **FNDAQRQATK** DAGTIAGLVN IRVVNEPTAA  
**201** ALAYGMDKTK DSLIAVYDLG GGTFDISVLE IAGGVFEVKA **TNGDTHLGGE**  
**251** **DFD LALS DYI LEEFRKTSGI DLS KER MALQ RVREA AEKAK CELSS AMETE**  
**301** VNLPFITANA DGAQHQIQMRI SRSKFEGITQ RLIERSIAPC KQCMKDAGVE  
**351** LKEINDVVLV **GGMTRIRSGG GGEEVLPEGP VRGVNPDEAV ALGAATLGGV**  
**401** LRGKASDLIL VDVTPSLGT SVVGDFVPI IPKNTTIPCM RSHIFTTVDD  
**451** GQTAIKFKVF **QGEREIASEN QIRGEFDLSG IPPAPRGVPQ VEVTFDIDAN**  
**501** GICHVTAKDK ATGKTQNITI TANGL SKEQ IEQMIRDSEQ HAEADRVKRE  
**551** LVEVRNNAET QLT TAERQLG EWKYVSDAEK ENVKTLVAEL RKAMENPNVA  
**601** KDDLAAATDK LQKAVMECGR TEYQQAAAAN SGQC

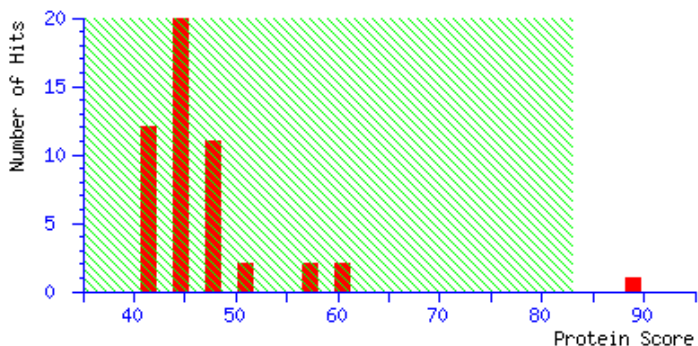
Sample: 222  
Mass Spectrum (positive reflector)

4700 Reflector Spec #1 MC=>BC=>NP0.7=>MC [BP = 1452.7, 1418]



## Mascot Score Histogram

Protein score is  $-10 \log(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p < 0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.

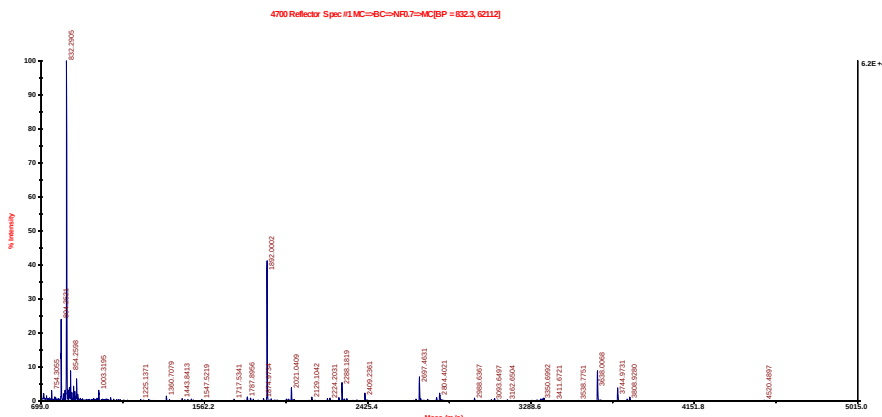


Match to: **gi|157868136** Score: **89** Expect: **0.014**nucleosome assembly protein [Leishmania major strain Friedlin] Nominal mass ( $M_r$ ): **39727**; Calculated pI value: **4.61**NCBI BLAST search of gi|157868136 against nrUnformatted sequence string for pasting into other applications Taxonomy: Leishmania major strain FriedlinLinks to retrieve other entries containing this sequence from NCBI Entrez:gi|68126076 from Leishmania major strain Friedlin Fixed modifications: Carbamidomethyl (C)Variable modifications: Oxidation (M)Cleavage by Trypsin: cuts C-term side of KR unless next residue is PSequence Coverage: **15%**

Matched peptides shown in **Bold**:

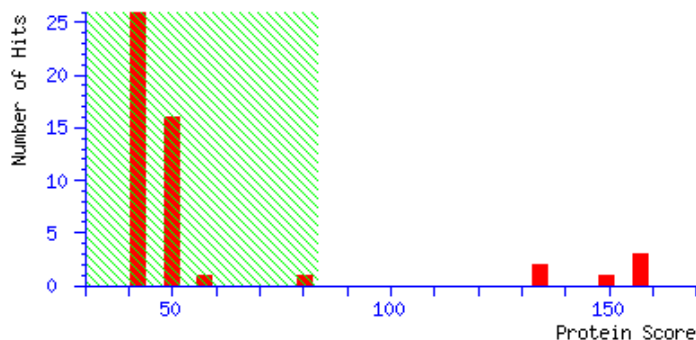
1 MPPKNQRDAP VVIPDEDDEG DMMEMGNMLD FQKYLDPDFS KNFMASLPEK  
51 IRQRAQVLSA YDKDLSAQQK AYKAKEMDIL RRYDALFEPL LQRRKEIVTG  
101 AAVSDEEVKK GMPEEHVNVI SVEVDETDEA AKAADAFGLE GFWLRVLRHH  
151 AVIDSTIEPH DEDVLKHLVD IRSSVAEHEY GSFQVIFTFS PNDFEEETI  
201 TATVSIKDDK SELTVSPITW KPGKNVMIHT VTKKQRAKRT GQVRTTTRDV  
251 PQLSFFWLFK KKTEAVGDDD AEKDGEEDDE EQRISMLEVL HTCIVPNAVR  
301 YYTGEAPNGF SDVDEEDEED EEEEEEEEEEI IPRGRGGNRG GRGGRGGRGI  
351

Sample: 252  
Mass Spectrum (positive reflector)



### Mascot Score Histogram

Protein score is  $-10^{\ast}\log(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p<0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



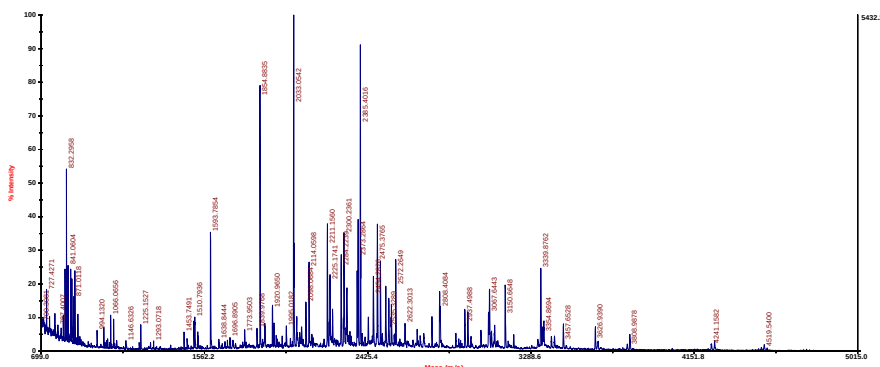
Match to: **gi|157870644** Score: **157** Expect: **2.2e-09** ATPase beta subunit [Leishmania major strain Friedlin] Nominal mass ( $M_r$ ): **56541**; Calculated pI value: **5.14** NCBI BLAST search of gi|157870644 against nr Unformatted sequence string for pasting into other applications Taxonomy: Leishmania major strain FriedlinLinks to retrieve other entries containing this sequence from NCBI Entrez:gi|68126939 from Leishmania major strain Friedlin Fixed modifications: Carbamidomethyl (C) Variable modifications: Oxidation (M) Cleavage by Trypsin: cuts C-term side of KR unless next residue is P Sequence Coverage: **31%**

Matched peptides shown in **Bold**:

**1** MLSRVQSA MI RRAAGVRA AS SAVAAAAAKP AEHKGRVGYV SQVIGAVVDV  
**51** HFSEGVPPL TALDVTEDLG **RDEPLTLEIV QHLDANTGRC IAMQTTDLLK**  
**101** LKSKVVSTGG NISVPVGRET LGRIFNV LGD **AIDQRGPVGE KMRMAIHAEA**  
**151** PKLADQAAED TILT GIKVI DLIL PYCKGG KIGLF GGAGV GKTVI MELI  
**201** NNVAKGHGGF **SVFAGVGERT REGTDLYLEM MQSKVIDLKG ESKC VLVY GQ**  
**251** MNEPPGARAR VAQSALTMAE YFRDVEGQNV **LLFIDNIFRF TQANSEVSAL**  
**301** LGRIPAAVGY QPTLAEDLGM LQERITSTTK GSITSVQAVY VPADDITDPA  
**351** PATTFSHLDA TT VLDRAVAE SG IYP AVNPL ECASRIMDPD VIDVDHYNVA  
**401** QDIVQMLTKY KELQDIIA VL **GIDELSEEDK VVVDRARKVT RFLSQPFQVA**  
**451** EVFTGMTGHY VQLVDTVESF SG LLMGSYDQ IP EM AFY MVG GIKSV LEKAK  
**501** AMAEEAAAME KQRRARQAAN ASDSQ

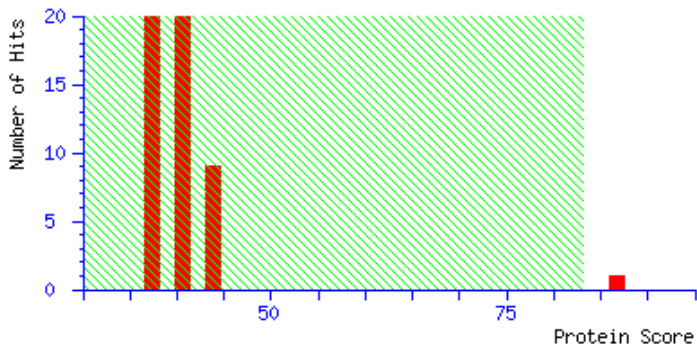
## Sample: 266 Mass Spectrum (positive reflector)

4700 Reflector Spec #1 MC=>BC=>NF0.7=>MC [BP = 2034.1, 5432]



## Mascot Score Histogram

Protein score is  $-10 \times \text{Log}(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p < 0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.

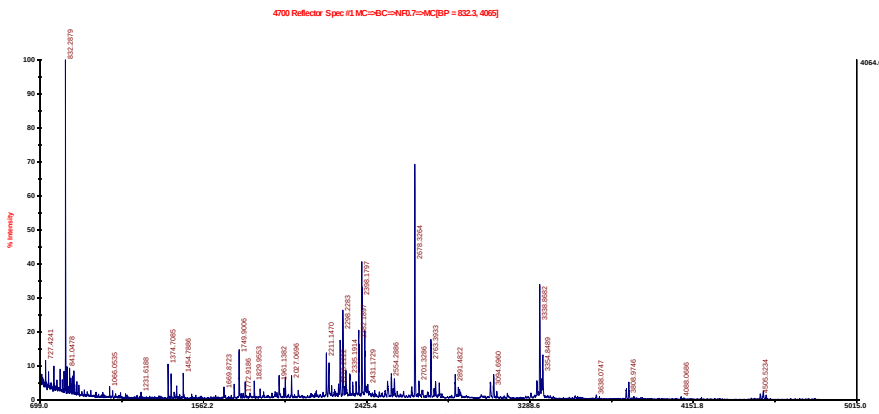


Match to: gi|157877007 Score: 87 Expect: 0.023 dihydrolipoamide acetyltransferase precursor [Leishmania major strain Friedlin] Nominal mass ( $M_r$ ): 49062; Calculated pI value: 7.01 NCBI BLAST search of gi|157877007 against nr Unformatted sequence string for pasting into other applications Taxonomy: Leishmania major strain Friedlin Links to retrieve other entries containing this sequence from NCBI Entrez:gi|68129918 from Leishmania major strain Friedlin Fixed modifications: Carbamidomethyl (C) Variable modifications: Oxidation (M) Cleavage by Trypsin: cuts C-term side of KR unless next residue is P Sequence Coverage: 17%

Matched peptides shown in **Bold**:

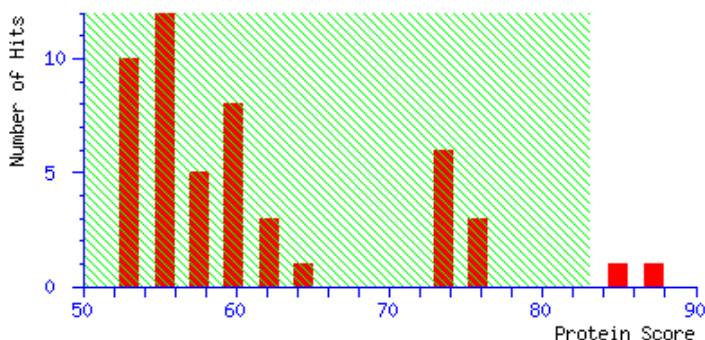
**1** MLRCRAVSKL ATLAALRFLT ITPIPMPALS PTMEKGKITE WCKQPGDFIR  
**51** PGDTFCNIET DKAVVSYDNA TEEGFFARVI TSPGEETVVG QTVCCLIVDEK  
**101** EGVHSDEVKN WKPEAAEAPA AAAEEAAPAAP AATTPVAAAP VAASGDRVKA  
**151** SPYARKMAAE KNVSLRGIKG TGGGVGRITS KDVAAAVASG TASSAAEVAA  
**201** PAKTAATAAL AAPAKPAAAK **GTPPANPNFT DIPVTTMRSV IAKRLHQSKN**  
**251** LEIPHYYLF DCRVDNMLAL IKQLNAKGNG EYKITVNDYI VKAVARANTL  
**301** VPEVNSSWQG DFIRQYATVD VSVAVATPTG LITPIIRNAQ AKGLVEISKE  
**351** TKALAKKARD GTLQPSEFQG GTCSVNLGA TGIPGFTAII NPPQAMILAV  
**401** GSAKPRAEIV KSEETGEFEM TGRVENVVSF **SASF DHRIVD GALGAKWFQH**  
**451** FHADAMENPLSLLL

Sample: 300  
 Mass Spectrum (positive reflector)



### Mascot Score Histogram

Protein score is  $-10 \cdot \log(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p < 0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.

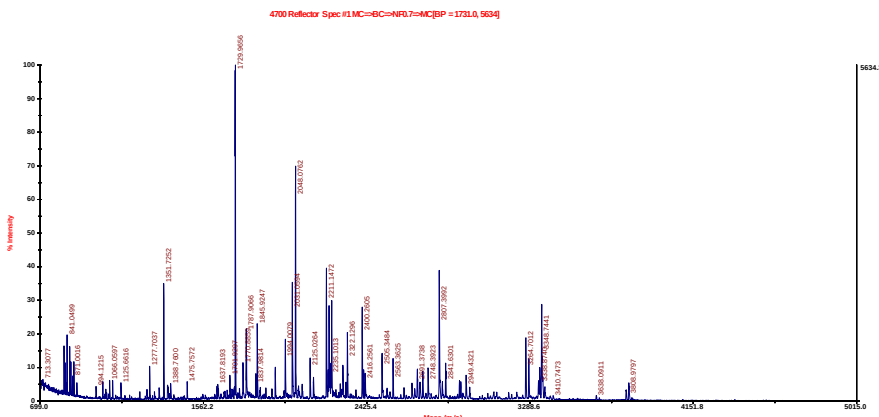


Match to: gi|1698880 Score: 87 Expect: 0.021 protein antigen LmSTI1 [Leishmania major] Nominal mass ( $M_r$ ): 62507; Calculated pI value: 5.91 NCBI BLAST search of gi|1698880 against nr Unformatted sequence string for pasting into other applications Taxonomy: Leishmania major Fixed modifications: Carbamidomethyl (C) Variable modifications: Oxidation (M) Cleavage by Trypsin: cuts C-term side of KR unless next residue is P Sequence Coverage: 18%

Matched peptides shown in **Bold**:

**1** DATELKNKGN EEF SAGRYVE AVNYFSKAIQ **LDEQNSVLYS NRSACFAAMQ**  
**51** KYKDALDDAD KCISIKPNWA KGYVRRGAAL HGMRRYDDAI AAYEKGLKVD  
**101** PSNSGCAQGV KDVQVAKARE ARDPIARVFT PEAFRKIQEN PKLSLLMLQP  
**151** DYVKMVDTVI RDPSQGRILYM EDQRFALTLM **YLSGMKIPND GDGEEEERPS**  
**201** AKAAETAKPK EEKPLTDNEK EALALKEEGN KLYLSKKFEE ALTKYQEAQV  
**251** KDPNNNTLYIL NVSavyFEQG **DYDKCIAECE HGIEHGRENH CDYTIIAKLM**  
**301** TRNALCLQRQ RKYEAAIDLY KRALVEWRNP DTLKKLTECE KEHQKAVEEA  
**351** YIDPEIAKQK KDEGNQYFKE **DKFPEAVAAY TEAIKRNPAAE HTSYSNRAAA**  
**401** YIKLGAFNDA LKDAEKCIEL KPDFVKGYAR KGHAYFWTKQ YNRALQAYDE  
**451** GLKVDPSENAD CKDGRYRTIM KIQEMASGQS ADGDEAARRA **MDDPEIAAIM**  
**501** QDSYMQQLVLK EMQNDPTRIQ EYMKDSGISS KINKLISAGI IRFGQ

Sample: 356  
 Mass Spectrum (positive reflector)

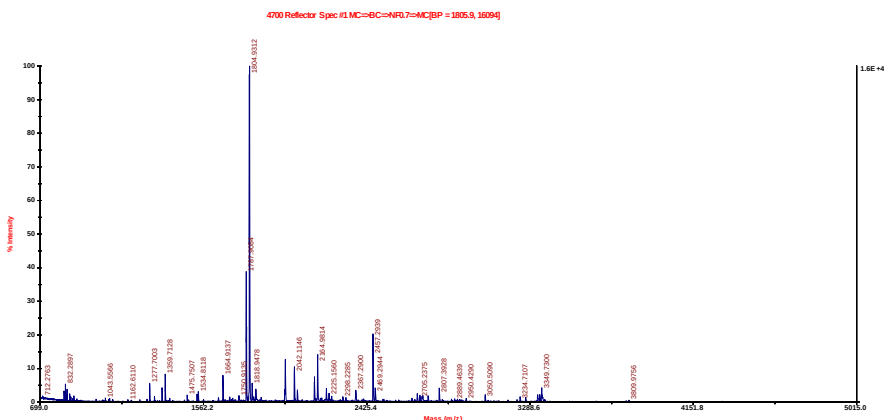


Match to: **gi|157871157** Score: 91 Expect: 0.008 heat shock protein 70-related protein [Leishmania major] Nominal mass (M<sub>r</sub>): 71001; Calculated pI value: 5.04 NCBI BLAST search of gi|157871157 against nr Unformatted sequence string for pasting into other applications Taxonomy: Leishmania major strain FriedlinLinks to retrieve other entries containing this sequence from NCBI Entrez:gi|68127196 from Leishmania major strain Friedlin Fixed modifications: Carbamidomethyl (C) Variable modifications: Oxidation (M) Cleavage by Trypsin: cuts C-term side of KR unless next residue is PSequence Coverage: 19%

Matched peptides shown in **Bold**:

**1** MSSTNAIGID LGTTYSVGV FKNEQVDIIA NDQGNRTTPS YVAFTETERL  
**51** IGDAAKNQVA MNPSNTVFDA KRMIGRKFDD PDLQSDMKHW PFKVIVKDGGK  
**101** PVISVEYQNQ TKTFFPPEIS AMVLQKMKET AEAYLGTTVK DAVITVPAYF  
**151** NDSQRQATKD AGSIAGLNVL RIINEPTAAA IAYGMDRKGD KGEKNVLIFD  
**201** LGGGTDFVTLL LTIESGVFEV KATAGDTHLG GEDFDNLRLVD YFATELKMR  
**251** GKDCRGNARA TRRLRTACER VKRTLSSSTT ANIEIDALYE GNDFFSKITR  
**301** ARFEEMCRDQ FEKCLEPVKK VLADADMKPQ DVDDVVLVGG STRIPKIQQI  
**351** VSQFFGGKEL NRSINPDEAV AYGAAVQAHI LAGGHSSKTD GLLLL DVTP  
**401** SLGVETSGGV MSLIPRNST MPVQKTQYNS NNADNQRNVV IKVYEGERPL  
**451** VSQCQLCLGF TLTDIPPMPR GKARINVTFD VNTDGILIVS AVEESGGRKE  
**501** AITIQNDTGR LSKEQIESMV REAEKFAED RMNSERVEAR NTLENYTFSM  
**551** RATLDDPDVQ NGITQGDRQQ IQDAVNAASS WLEKNREATK EEYMEQTCLI  
**601** EGIAHPILSE FYKKRVMEAP PSAGPKDGAP SDEVPHAEDEV

Sample: 405  
 Mass Spectrum (positive reflector)

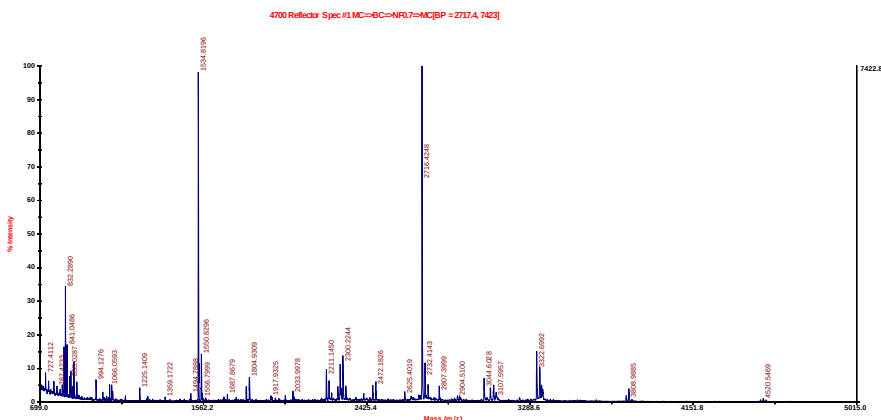


Match to: **gi|146080653** Score: 100 Expect: 0.0011alpha tubulin [Leishmania infantum JPCM5] Nominal mass (M<sub>r</sub>): 36306; Calculated pI value: 6.78 NCBI BLAST search of gi|146080653 against nr Unformatted sequence string for pasting into other applications Taxonomy: Leishmania infantum JPCM5 Links to retrieve other entries containing this sequence from NCBI Entrez:gi|134068142 from Leishmania infantum Variable modifications: Oxidation (M),Carbamidomethyl (C),Pyro-carbamidomethyl (N-term C) Cleavage by Trypsin: cuts C-term side of KR unless next residue is P Sequence Coverage: 18%

Matched peptides shown in **Bold**:

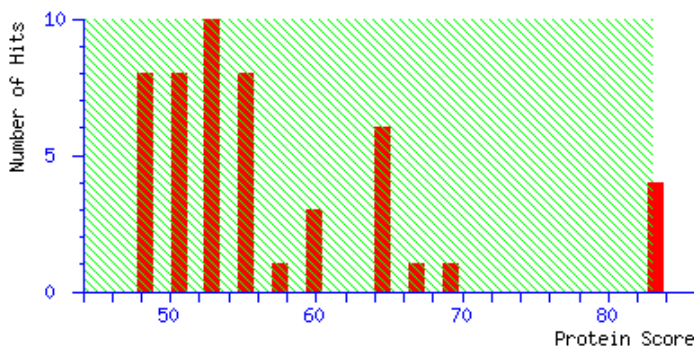
**1** MHTDTHAHTP PSSRSAALLL LLLPPCAGRL PRVSSPHLAL SLPPPLPSWR  
**51** TQNMRDAHTR TPTEKKTRSS SLSFFEQTPL NRLLTPLSSF SAMREAICIH  
**101** IGQAGCQVGN TCWELFCLEH GIQPDGSMPS DKCIGVEDDA FNTFFSETGA  
**151** GKHVPRCIFL DLEPTVDEV RTGYRQLFN PEQLVSGKED AANNYARGHY  
**201** TIGKEIVDLA LDRIRKLADN CTGLQGFMVF RESPFLPLS LPLPLPLFPP  
**251** LPREGNTKEK IAKSDVYMM RHKDEINPPM PPPLPSLPPC VCVRVFFLLL  
**301** SVAVVPLLFP FYVLVCLLFS PFASLFL

Sample: 407  
 Mass Spectrum (positive reflector)



### Mascot Score Histogram

Protein score is  $-10 \times \text{Log}(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p < 0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



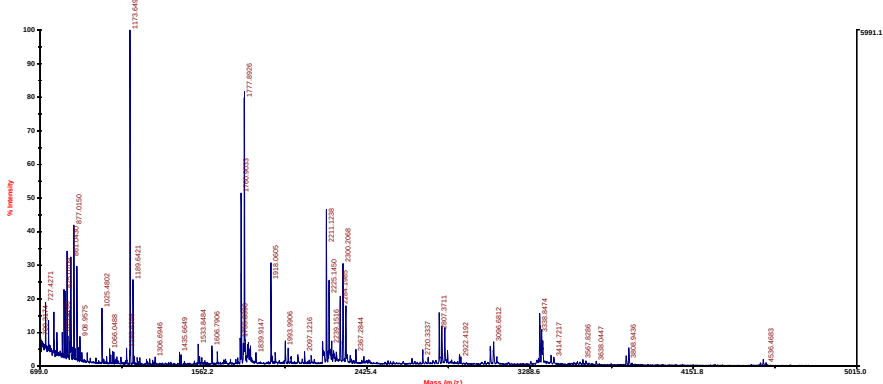
Match to: gi|146086967 Score: 83 Expect: 0.053 peroxidoxin [Leishmania infantum JPCM5] Nominal mass ( $M_r$ ): 25354; Calculated pI value: 6.43NCBI BLAST search of gi|146086967 against nr Unformatted sequence string for pasting into other applications Taxonomy: [Leishmania infantum JPCM5](#) Links to retrieve other entries containing this sequence from NCBI Entrez:gi|16751316 from [Leishmania infantum](#)gi|134069782 from [Leishmania infantum](#) Variable modifications: Carbamidomethyl (C),Oxidation (M)Cleavage by Trypsin: cuts C-term side of KR unless next residue is P Sequence Coverage: 16%

Matched peptides shown in **Bold**:

**1** MLRRRLPTSCF LKRSQFRGFA ATSPLLNLDY QMYRTATVRE AAPQFSGQAV  
**51** VNGAIKDINM NDYKGKYIVL FFYPMDFTFV CPTEIIAFSD RHADFEKLNT  
**101** QVVAVSCDSV YSHLAWVNTP RKKGGLGEMH IPVLADKSME IARDYGVLIE  
**151** ESGIALRGLF IIDKKGILRH STINDLPVGR NVDEALRVLE AFQYADENGD  
**201** AIPCGWPKPGQ PTLDTTKAGE FFEKNM

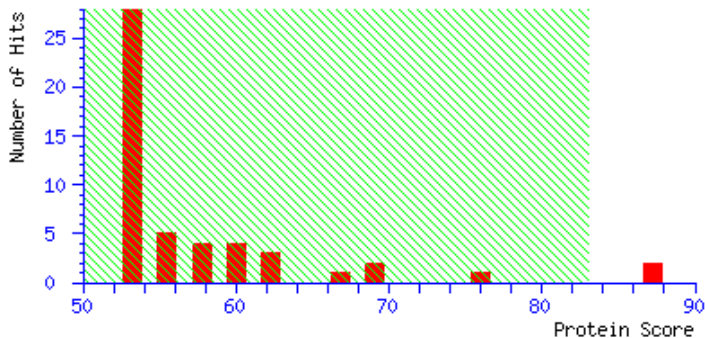
## Sample: 411 Mass Spectrum (positive reflector)

4700 Reflector Spec #1 MC=>BC=>NF0.7=>MC [BP = 1173.6, 5991]



## Mascot Score Histogram

Protein score is  $-10 \times \text{Log}(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p < 0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



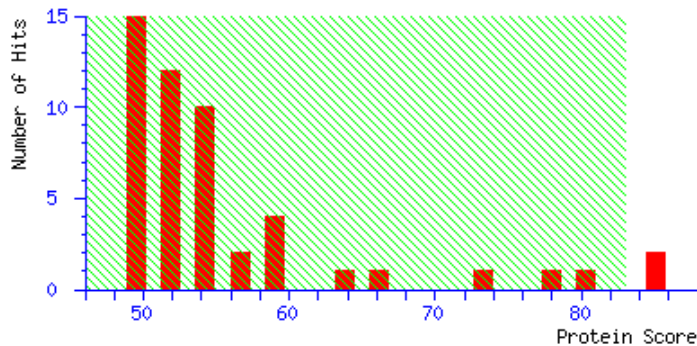
Match to: gi|157865592 Score: 87 Expect: 0.02114-3-3 protein [Leishmania major strain Friedlin] Nominal mass (M<sub>r</sub>): 29126; Calculated pI value: 5.19 NCBI BLAST search of gi|157865592 against nr Unformatted sequence string for pasting into other applications Taxonomy: [Leishmania major strain Friedlin](#) Links to retrieve other entries containing this sequence from NCBI Entrez:gi|68124800 from [Leishmania major strain Friedlingi|257900393](#) from [Leishmania major](#) Variable modifications: Carbamidomethyl (C),Oxidation (M) Cleavage by Trypsin: cuts C-term side of KR unless next residue is P Sequence Coverage: 27%

Matched peptides shown in **Bold**:

**1 MTNVFKVPEK REELVYTAKI AEQCERHDEI LFCMKRAVKM NPrLSSeERN  
51 LLSAAKYII SARRACWRSM SSMAHKEDSH KGKTASLFNG FQHQVEKELA  
101 EICSDILELL DKYLIPAADN DESKVYYYKL KGDYHRYFAE VESGSDTQKN  
151 LALEAYKKAS EFTTSLKPTS PIRLGLALNF SVFYYEILRS PDKGCQLARQ  
201 AFEEALSDPE VLDEEQHKES ALIMQLLRDN LALWTEDSRP EGQDDGTAME  
251 ELE**

## Mascot Score Histogram

Protein score is  $-10 \cdot \log(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p < 0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.

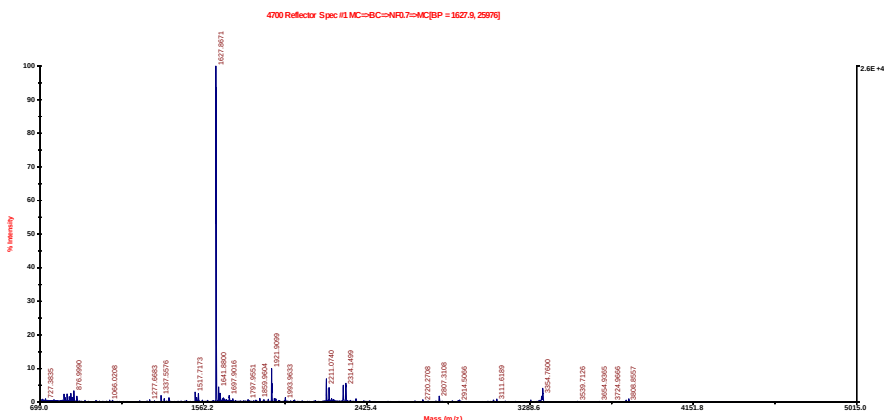


Match to: gi|157864162 Score: **85** Expect: **0.039** hypothetical protein [Leishmania major strain Friedlin] Nominal mass ( $M_r$ ): **30868**; Calculated pI value: **5.34** NCBI BLAST search of gi|157864162 against nr Unformatted [sequence string](#) for pasting into other applications Taxonomy: [Leishmania major strain Friedlin](#) Links to retrieve other entries containing this sequence from NCBI Entrez:gi|68124086 from [Leishmania major strain Friedlin](#) Variable modifications: Carbamidomethyl (C),Oxidation (M)Cleavage by Trypsin: cuts C-term side of KR unless next residue is P Sequence Coverage: **14%**

Matched peptides shown in **Bold**:

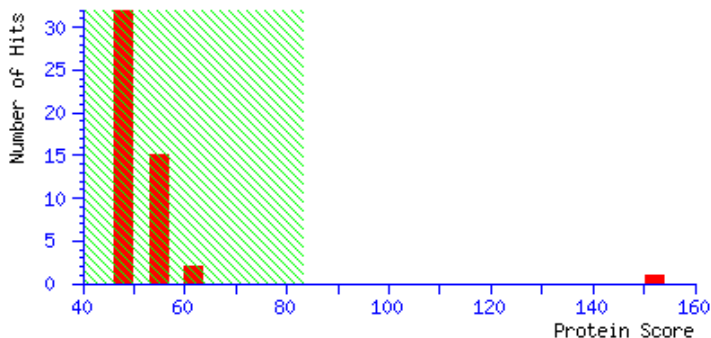
**1** MDPLAGLTTG SRNVIDDDFFR WSRLSLDKFK RDLTVEASAF QAGNLWREQY  
**51** TKAеваQMLN GQLALMLGVV DTQMEGTTAA SAELMRAVLR EADRQRVTLD  
**101** IDVKSVLNNA GGVNAMGEHG RQLMSGPTGR LAPLTVEEAS GEAARQLAEA  
**151** NAEVKRLQAK LQQVTDAYTQ VMRGRSSDTE NLLSMQDAMN DKERLAAELA  
**201** RRCQQQDPSL AATVQELRAE VAEAKRELAS RLHQSTQYQQ VKHLLAQRN

Sample: 414  
 Mass Spectrum (positive reflector)



### Mascot Score Histogram

Protein score is  $-10 \times \text{Log}(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p < 0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.

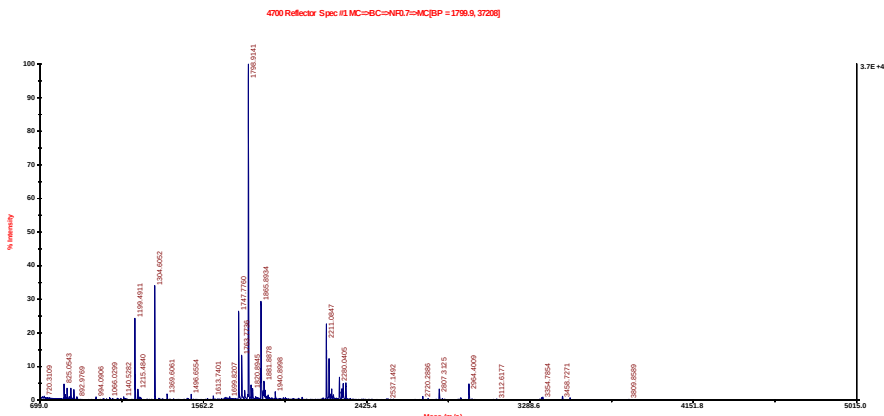


Match to: [gi|157870822](#) Score: **152** Expect: **7e-09** hypothetical protein [Leishmania major strain Friedlin] Nominal mass (M<sub>r</sub>): **32835**; Calculated pI value: **5.52** NCBI BLAST search of gi|157870822 against nr Unformatted [sequence string](#) for pasting into other applications Taxonomy: [Leishmania major strain Friedlin](#) Links to retrieve other entries containing this sequence from NCBI Entrez:[gi|68127028](#) from [Leishmania major strain Friedlin](#) Variable modifications: Carbamidomethyl (C),Oxidation (M)Cleavage by Trypsin: cuts C-term side of KR unless next residue is P Sequence Coverage: **35%**

Matched peptides shown in **Bold**:

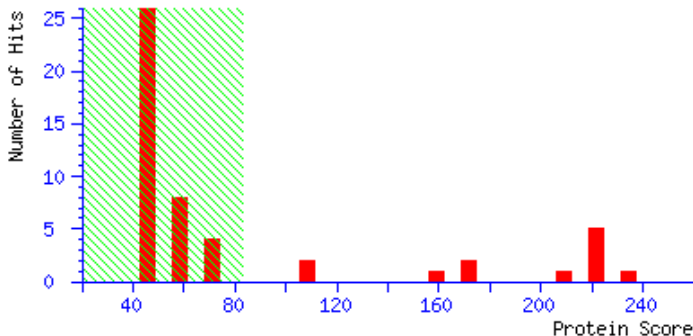
**1** MQKTAILQILR EQDDASSSCV LALLKGNAF QCLSKSEGED ILKRDNEDAE  
**51** KLSDGIRIAI RKGIVSGKPD VSPVNKLNVV GLSAQEVTDK IVAALPSKNG  
**101** NVIILQGLSG TGKGTTVSKL KDALPKCVTW SNGNIFRSYT YLCNEVLAAQ  
**151** GKEITEESLT DELLSSVEKR VTFVKCEDGM FHTMLDGTRR VADVQNTLLK  
**201** TPLISSKVPT VAQQTQGEVI RFGAGAVKQL SADGYNVILE GRSQTLDYIS  
**251** SPLRFELIIP DVEVLGQRRA AQRVMAKALE MLDDSASDEE VSKVLVKAVE  
**301** TLQ

Sample: 426  
 Mass Spectrum (positive reflector)



### Mascot Score Histogram

Protein score is  $-10 \cdot \log(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p < 0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.

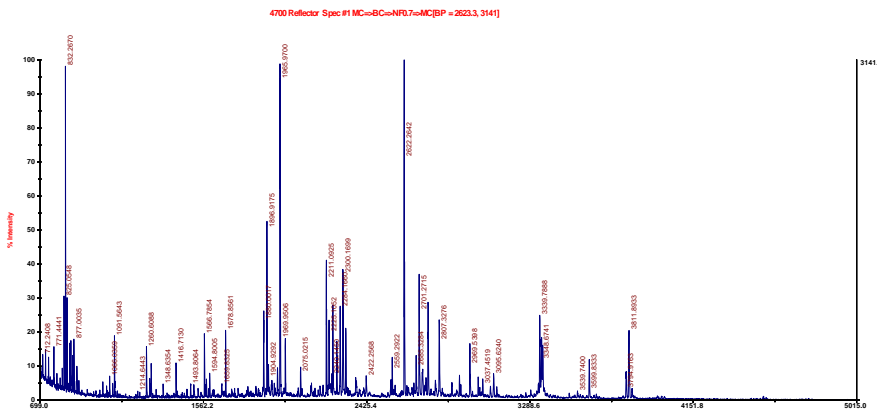


Match to: gi|157872022 Score: 234 Expect: 4.4e-17 activated protein kinase c receptor (LACK); guanine nucleotide-binding protein beta subunit-like protein [Leishmania major] Nominal mass ( $M_r$ ): 34378; Calculated pI value: 6.05 NCBI BLAST search of gi|157872022 against nr Unformatted [sequence string](#) for pasting into other applications Taxonomy: [Leishmania major strain Friedlin](#) Links to retrieve other entries containing this sequence from NCBI Entrez:gi|157872024 from [Leishmania major strain Friedlin](#)gi|68127629 from [Leishmania major strain Friedlin](#)gi|68127630 from [Leishmania major strain Friedlin](#) Variable modifications: Carbamidomethyl (C),Oxidation (M)Cleavage by Trypsin: cuts C-term side of KR unless next residue is P Sequence Coverage: 50%

Matched peptides shown in **Bold**:

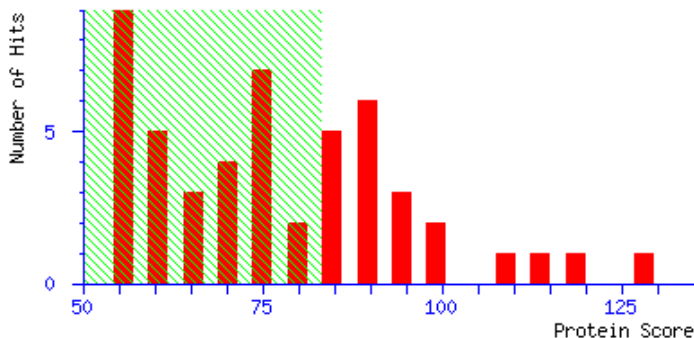
**1** MNYEGHLKGH RGWVTSACP QQAGSYIKVV STSRDGTAIS WKANPDR**HSV**  
**51** DSDYGLPNHR LEGHTGFVSC VSLAHATDYA LTASWDRSIR MWDLRNGQCQ  
**101** RKFLKHTKDV LAVAFSPDDR LIVSAGRDNV IRVWNVAGEC MHEFLRDGHE  
**151** DWVSSICFSP SLEHPIVVSG SWDNTIKVWN VNGGKCERTL KGHSNYVSTV  
**201** TVSPDGSLCA SGGKDGAALL WDLSTGEQLF KINVESPINQ IAFSPNRFWM  
**251** CVATERSLSV YDLESKAVIA ELTPDGAKPS ECISIAWSAD GNTLYSGHKD  
**301** NLIRVWSISD AE

Sample: 478  
Mass Spectrum (positive reflector)



## Mascot Score Histogram

Protein score is  $-10 \times \log(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p < 0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



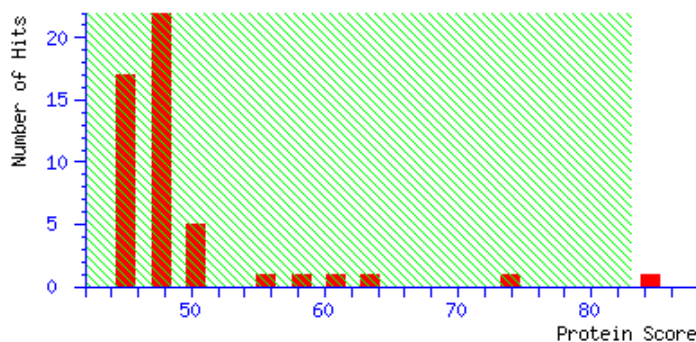
Match to: gi|157872028 Score: 128 Expect: 1.8e-06 heat-shock protein hsp70 [Leishmania major strain Friedlin] Nominal mass ( $M_r$ ): 71608; Calculated pI value: 5.30 NCBI BLAST search of gi|157872028 against nr Unformatted [sequence string](#) for pasting into other applications Taxonomy: [Leishmania major strain Friedlin](#) Links to retrieve other entries containing this sequence from NCBI Entrez:gi|157872030 from [Leishmania major strain Friedlin](#)gi|68127632 from [Leishmania major strain Friedling](#)gi|68127633 from [Leishmania major strain Friedlin](#) Variable modifications: Carbamidomethyl (C),Oxidation (M)Cleavage by Trypsin: cuts C-term side of KR unless next residue is P Sequence Coverage: 29%

### Matched peptides shown in **Bold**:

**1 MTFDGAIGID LGTTYSCVGV WQNERVDIIA NDQGNRTTPS YWAFTDSERL  
51 IGDAAKNQVA MNPHNTVFDA KRLIGRKFND SVVQSDMKHW PFKVTTKGDD  
101 KPVISVQYRG EEKFTFPEEI SSMVLLKMKE TAEAYLGKQV KKAVVTVPAY  
151 FNDSQRQATK DAGTIAGLEV LRIINEPTAA AIAYGLDKGD DGKERNVLIF  
201 DLGGGTFDVT LLTIDGGIFE VKATNGDTHL GGEDFDNRLV TFFFTEEFKRK  
251 NKGKNLASSH RALRRLRTAC ERAKRTLSSA TQATIEIDAL FENIDFQATI  
301 TRARFEELCG DLFRSTIQPV ERVLQDAKMD KRSVHDVVVLV GGSTRIPKVQ  
351 SLVSDFFGGK ELNKSINPDE AVAYGAAVQA FILTGGKSKQ TEGLLLLDVT  
401 PLTLGIETAG GVMTALIKRN TTIPTKKSQI FSTYADNQPG VHIQVFEGER  
451 AMTKDCCHLLG TFDLSGIPPA PRGVPQIEVT FDLDANGILN VSAEKGTCGK  
501 RNQITITNDK GRLSKDEIER MVNDAMKYEE DDKAQRDRVE AKNGLENYAY  
551 SMKNTLSDSN VSGKLDDSDK ATLNKEIDAA LEWLSSNQEAE TKEEYEHRQK  
601 ELESVCNPIM TKMYQSMGGA AGGMPGGMPG GMPDMSGMSG GAGPAGGASS  
651 GPKVEEVYD**

## Mascot Score Histogram

Protein score is  $-10^{\ast}\log(P)$ , where P is the probability that the observed match is a random event. Protein scores greater than 83 are significant ( $p<0.05$ ). Protein scores are derived from ions scores as a non-probabilistic basis for ranking protein hits.



Match to: gi|157864052 Score: 84 Expect: 0.04 dipeptidyl-peptidase III; metallo-peptidase, Clan M-, Family M49 [Leishmania major strain Friedlin] Nominal mass ( $M_r$ ): 75920; Calculated pI value: 5.30 NCBI BLAST search of gi|157864052 against nr

Unformatted sequence string for pasting into other applications Taxonomy: [Leishmania major strain Friedlin](#) Links to retrieve other entries containing this sequence from NCBI Entrez:gi|68223784 from [Leishmania major strain Friedlin](#) Variable modifications: Carbamidomethyl (C),Oxidation (M)Cleavage by Trypsin: cuts C-term side of KR unless next residue is P Sequence Coverage: 26%

Matched peptides shown in **Bold**:

**1** MSHNTLYITP RAVPYCILSI ANAFKDLTPK QRHYAHHMMA AGWCGAPVVA  
**51** EQLSPESLPL LRLFFKVLSA QPLDTFKANS VNAGVDPDEV DQFLEYFAMV  
**101** YSNMGNYISF GDTKFVPSIP KERFAKIVAS ADGSPAVDAQ LLDAIYNLDD  
**151** DKLTLDFPPK GLTRYYSPNV TREDAAVAND FLASKKMDGV NTRVFKEEDG  
**201** SLVIRVAAAT ERTVPAEKFN GRTIAIYYGD YKEEMARVVA ELRKAQPYAE  
**251** NETEVVMLNH YIEHFQYGDV EAHKESQKEW VRDVGPTVET **NIGFIESYRD**  
**301** PSGVRAEWEG FVAVVNKEQS KMYGALVAQG EKFIAQLPWG KAFEKDVFSS  
**351** PDFTSLDVLG FASSGIPAGI **NIPNYDDIRQ** TVGFKNVYLS NVVSAMTFKD  
**401** KLNYYITEADW ELYKASILAA TSVNVGIHEL **LGHGTGKLLS** ENSDGTNFND  
**451** EKTVDPLSGK PVATWYKPGD TYSSVFGGLG SSYEECRAEA VSYYLCLLPE  
**501** LLEIFNIQTA KEQQDVYIVC WLNMVVRAGLV **GLEFYTPEKQ** QWRQAHMRAR  
**551** FCILQALARA PNPIVQITEN AKEGVLTLD RERIATDGRQ **AIGDLLVNLN**  
**601** VNKATADAKR **GSAYFENMTV** VSDQYVHYRD IIMARRKPRK QYVQPHTFIS  
**651** GDTVEVLEFA GSVEGVVESF VTRHREIPL