The iron-dependent mitochondrial superoxide dismutase SODA promotes *Leishmania* virulence

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Bidyottam Mittra, Maria Fernanda Laranjeira-Silva, Danilo Ciccone Miguel, Juliana Perrone Bezerra de Menezes, and Norma W. Andrews

*From the Department of Cell Biology and Molecular Genetics, University of Maryland, College Park, Maryland 20742-5815*

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Leishmaniasis is one of the leading globally neglected diseases, affecting millions of people worldwide. *Leishmania* infection depends on the ability of insect-transmitted metacyclic promastigotes to invade mammalian hosts, differentiate into amastigotes, and replicate inside macrophages. To counter the hostile oxidative environment inside macrophages, these protozoans contain anti-oxidant systems that include iron-dependent superoxide dismutases (SODs) in mitochondria and glycosomes. Increasing evidence suggests that in addition to this protective role, *Leishmania* mitochondrial SOD may also initiate $\text{H}_2\text{O}_2$-mediated redox signaling that regulates gene expression and metabolic changes associated with differentiation into virulent forms. To investigate this hypothesis, we examined the specific role of SODA, the mitochondrial SOD isoform in *Leishmania amazonensis*. Our inability to generate *L. amazonensis* SODA null mutants and the lethal phenotype observed following RNAI-mediated silencing of the *Trypanosoma brucei* SODA ortholog suggests that SODA is essential for trypanosomatid survival. *L. amazonensis* metacyclic promastigotes lacking one SODA allele failed to replicate in macrophages and were severely attenuated in their ability to generate cutaneous lesions in mice. Reduced expression of SODA also resulted in mitochondrial oxidative damage and failure of SODA/$\Delta$ sodA promastigotes to differentiate into axenic amastigotes. SODA expression above a critical threshold was also required for the development of metacyclic promastigotes, as SODA/$\Delta$ sodA cultures were strongly depleted in this infective form and more susceptible to reactive oxygen species (ROS)-induced stress. Collectively, our data suggest that SODA promotes *Leishmania* virulence by protecting the parasites against mitochondrion-generated oxidative stress and by initiating ROS-mediated signaling mechanisms required for the differentiation of infective forms.

Leishmaniasis affects an estimated 12 million people worldwide, with an estimated 350 million at risk of infection (1). Depending on the *Leishmania* species, symptoms vary from self-healing skin lesions to a visceralizing form that can be lethal. The absence of efficacious and cost-effective drugs, combined with the emergence of drug resistance, accentuates the critical need for new therapeutic targets.

The life cycle of *Leishmania* spp. alternates between non-virulent promastigotes inside the insect vectors and virulent amastigotes inside mammalian hosts. Transmission of the parasites to mammals occurs through sand fly bites (2). Amastigotes are adapted to survive and replicate inside acidic parasitophorous vacuoles of macrophages. After ingestion by sand flies during a blood meal, amastigotes transform into promastigotes, which replicate in the fly’s digestive tract. As nutrients become depleted, promastigotes cease to replicate and move up the sand fly gut toward the proboscis, where they mature into infective metacyclic forms. Metacyclics are reintroduced into new hosts during the next feeding cycle and enter host macrophages where they transform into amastigotes. To adapt to the rapidly changing environmental conditions during its life cycle, *Leishmania* undergoes extensive morphological and metabolic changes orchestrated at the post-transcriptional and post-translational levels (3–5). Despite significant progress in characterizing these life cycle-associated developmental changes, the molecular pathways that initiate differentiation in *Leishmania* are still poorly understood.

Over the last 2 decades, a role for reactive oxygen species (ROS)\(^2\) as regulators of physiological and biological responses (redox biology) has emerged (6–8). Two of the most abundant ROS generated as by-products of mitochondrial respiration or as end products of metabolic reactions, $\text{O}_2^-$ and $\text{H}_2\text{O}_2$, have been linked to cell fate determination. Subtle increases in intracellular $\text{O}_2^-$ levels promote cell proliferation, whereas low-level accumulation of $\text{H}_2\text{O}_2$ can inhibit cell growth and initiate differentiation (9–11). Recent studies specifically implicate the mitochondrion-generated ROS in intracellular signaling (12–14). $\text{O}_2^-$, the highly toxic ROS resulting from reduction of $\text{O}_2$ by complexes I, II, and III of the mitochondrial electron transport chain (ETC) (15), is rapidly converted by cytosolic or mitochondrial dismutases (SOD) into $\text{H}_2\text{O}_2$. Because of its higher stability, membrane diffusibility, and

\(1\) To whom correspondence should be addressed: Dept. of Cell Biology and Molecular Genetics, 2134 Bioscience Research Bldg., University of Maryland, College Park, MD 20742-5815. Tel.: 301-405-8418; Fax: 301-314-9489; E-mail: andrewsn@umd.edu.

\(2\) The abbreviations used are: ROS, reactive oxygen species; SOD, superoxide dismutase; ASL, adenosuccinate lyase; TEM, transmission electron microscopy; SEM, scanning electron microscopy; BMM, bone marrow-derived macrophage; FDA, fluorescein diacetate; ETC, electron transport chain; FD, forward; RV, reverse.
ability to promote target-specific thiol modifications, \( \text{H}_2\text{O}_2 \) is considered a primary intracellular signaling ROS molecule (7, 10).

Recent evidence indicates that \( \text{H}_2\text{O}_2 \) generated within mitochondria through the action of SOD plays a central role in regulating differentiation of \( L. \text{amazonensis} \) promastigotes into infective amastigotes (16, 17). Given that SOD enzymes in trypanosomatid protozoa exclusively utilize iron as an essential co-factor (18), these enzymes have emerged as important links between iron and ROS-regulated differentiation pathways in \( L. \text{amazonensis} \). Notably, the role of \( \text{H}_2\text{O}_2 \) in triggering promastigote to amastigote differentiation was directly demonstrated during studies of iron uptake in \( L. \text{amazonensis} \) promastigotes (16). Subsequent studies with \( L. \text{amazonensis} \) lines defective in mitochondrial iron import (\( \text{LMIT1}/\Delta\text{lm}11 \)) implicated mitochondrial iron-dependent SOD (SODA) and mitochondrial-generated ROS in the generation of virulent forms (17).

Further supporting a role for SODA and \( \text{H}_2\text{O}_2 \) in the development of virulence in \( L. \text{amazonensis} \), two cues known to trigger in vitro amastigote differentiation, low pH and high temperature (19), are also effective triggers of ROS generation (20–22). Exposure to elevated temperature is thought to lead to hyperpolarization of mitochondria, increased respiratory rate, and a ROS surge that is not only tolerated by \( L. \text{amazonensis} \) but is actually used as a differentiation signal. Stationary phase promastigotes show increased resistance to oxidative stress and enhanced SOD activity as they differentiate into amastigotes. In contrast, low SOD activity induces logarithmic phase promastigotes to accumulate higher levels of ROS and to undergo apoptosis, a process that can be reversed by overexpression of mitochondrial SODA (21).

Here, we confirm the prediction that iron import into mitochondria is required for activity of \( L. \text{amazonensis} \) SODA. Furthermore, by targeting the SODA gene for allelic knock-out and examining the phenotype of mutant lines, we conclude that SODA expression is required for maintaining mitochondrial redox balance and also for the development of parasite virulent forms.

**Results**

**Amastigote differentiation is associated with mitochondrial iron import and activation of mitochondrial SODA**

In earlier work, we identified LMIT1, an \( L. \text{amazonensis} \) mitochondrial iron transporter, and we suggested that the reduced ability of \( \text{LMIT1}/\Delta\text{lm}11 \) promastigotes to import iron into mitochondria and activate the iron-dependent SODA might be responsible for their failure to differentiate into amastigotes (17). This hypothesis was largely based on the reduced SODA activity observed in total parasite extracts and on morphological evidence for mitochondrial oxidative damage in \( \text{LMIT1}/\Delta\text{lm}11 \) parasites subjected to the low pH/high temperature protocol for axenic amastigote differentiation. To directly investigate the specific role of SODA in the signaling process leading to amastigote differentiation, we developed antibodies capable of distinguishing the mitochondrial SODA from glycosomal SODB, and we compared SOD activity in mitochondrial fractions of wild-type and \( \text{LMIT1}/\Delta\text{lm}11 \) parasites undergoing low pH/high temperature-induced axenic differentiation.

Specific polyclonal antibodies against mitochondrial SODA and SODB were generated using purified recombinant proteins (supplemental Fig. S1). The immunofluorescence localization of anti-SODA antibodies was identical to the staining pattern of MitoTracker Red, a mitochondrion-specific dye (Fig. 1A). This result demonstrates the mitochondrial localization of endogenous SODA, as proposed earlier based on overexpression of GFP-tagged SODA (23).

To obtain mitochondrion-enriched fractions free of glycosomal contaminants, subcellular fractionation of promastigotes expressing a 3×FLAG-tagged form of the LMIT1 mitochondrial iron transporter was carried out with increasing concentrations of digitonin, as described previously (17). Western blotting detection of organelle-specific markers (adenosuccinate lyase for cytosol (24), LMIT1-FLAG for mitochondria (17), and arginase for glycosomes (25)) showed that pellets obtained after 1 mg/ml digitonin treatment contained ~98% of the total LMIT1-FLAG and less than 5% of the total cellular arginase, reflecting mitochondrial enrichment in fractions mostly devoid of glycosomal contaminants (Fig. 1B). Immunoblot with the specific antibodies against SODA or SODB confirmed that

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**Figure 1. SODA localizes to mitochondria.** A, immunolocalization of SODA in \( L. \text{amazonensis} \) promastigotes was performed using polyclonal antibodies against SODA (green), and mitochondria were stained with MitoTracker Red (red). Merging the two images (merge) confirmed the mitochondrial localization of SODA. Bar, 4 μm. B, SODA is enriched in mitochondrial fractions. Subcellular fractionation of \( L. \text{amazonensis} \) promastigotes expressing FLAG-tagged LMIT1 was performed using increasing concentrations of digitonin as indicated. Proteins in the supernatant (S) and pellet (P) fractions were detected by Western blotting using antibodies against cytoplasmic adenosuccinate lyase, mitochondrial FLAG-tagged LMIT1, and glycosomal arginase. SODA and SODB were detected with specific polyclonal antisera and LMIT1–3×FLAG with an anti-FLAG monoclonal antibody.
higher during early differentiation (6 h), it was ~2.5-fold lower than WT at the 48-h time point when most WT parasites had already assumed the rounded morphology typical of amastigotes (Fig. 2A). Western blot analysis showed comparable amounts of SODA protein and only trace levels of SODB in both WT and LMIT1/Δmit1 mitochondrial fractions (Fig. 2, B and C), indicating that the reduced SOD activity in LMIT1/Δmit1 cells is likely to reflect SODA inactivation as a result of impaired LMIT1-mediated iron import into mitochondria. Importantly, we also observed down-regulation (of about 2-fold) in the expression of ascorbate-dependent peroxidase (APX) in LMIT1/Δmit1 mitochondrial lysates, when compared with WT (Fig. 2, B and C). Expression of APX, a mitochondrial protein required for the enzymatic breakdown of H$_2$O$_2$ in Leishmania, is known to be up-regulated in response to H$_2$O$_2$ accumulation (26). Collectively, these results demonstrate that differentiation of L. amazonensis promastigotes into infective amastigotes is associated with activation of the mitochondrial iron-dependent SODA and accumulation of H$_2$O$_2$, a ROS previously shown to directly trigger differentiation (16).

**SODA is an essential gene in L. amazonensis and in Trypanosoma brucei procyclids**

To further understand how mitochondrial SODA regulates the ROS signaling pathway leading to amastigote differentiation, we proceeded to generate SODA null mutants. Knock-out constructs carrying drug resistance gene cassettes flanked by 5’- and 3’-UTR regions of the SODA gene were generated and used for gene replacement through homologous recombination (supplemental Fig. S2A). Replacement of a single SODA allele with HYG or PHLEO drug resistance gene cassettes was possible, and allelic integration into the desired locus was confirmed in each case by PCR and Southern blot analysis (supplemental Fig. S2B). SODAΔsodA promastigotes showed ~53% reduction in SODA protein levels as the parasites reached early stationary growth, when compared with WT (Fig. 3). During the log phase of growth, both SODA and SODB were expressed at lower levels, and no significant difference was observed between WT and SODAΔsodA (Fig. 3, A and B). Repeated attempts to generate a SODA null line by replacing the second allele were unsuccessful. Similar to our previous experience with LMIT1, a SODA ORF was consistently detected in lines resistant to both phleomycin and hygromycin, even when the intended in situ integration of the targeting drug resistance markers was confirmed. Thus, SODA appears to be essential for the viability of L. amazonensis promastigotes. Providing further evidence that SODA is an essential gene in trypanosomatid parasites, tetracycline-inducible RNAi-mediated ablation of SODA in T. brucei procyclids (supplemental Fig. S3A) markedly impaired the parasites’ ability to sustain replication after 72 h in culture (supplemental Fig. S3B). T. brucei procyclids were reported to be partially dependent on mitochondrial metabolism (27, 28), suggesting that the phenotype we observed may be related to gradual mitochondrial damage as accumulated ROS reach toxic levels in the absence of SODA.
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In addition to the low pH/high temperature stimuli, in previous work we showed that up-regulation of the iron uptake machinery in response to iron deprivation also triggers differentiation of *L. amazonensis* promastigotes into amastigotes in a process dependent on mitochondrial H$_2$O$_2$, the product of SODA (16, 17). Thus, we also examined the effects of iron deprivation on the growth and differentiation of SODA/ΔsodA promastigotes. When cultured in iron-deficient medium, both WT and SODA-deficient parasites were initially able to replicate but reached a maximum cell density (~3 × 10$^7$/ml) that was lower than what is normally observed in complete growth medium (Fig. 5A, compare with WT in Fig. 6A). However, in contrast to WT, the SODA/ΔsodA promastigote count steadily declined after reaching the peak density (Fig. 5A), resembling the “population crash” we previously reported for Δlit1/Δlit1 (lacking the ferrous iron transporter LIT1) and LMIT1/Δlmit1 (partially deficient in the mitochondrial iron importer LMIT1) parasites grown in iron-depleted medium (16, 17). Microscopic analysis of the cultures showed that on day 5 of iron deprivation >55% of WT parasites lacked a visible flagellum and had assumed the rounded amastigote-like morphology, whereas only 20% of viable SODA/ΔsodA cells were able to undergo this transformation (Fig. 5B). Complementation with episomally expressed SODA (SODA/ΔsodA + SODA) partially restored both the growth pattern and the parasite’s ability to transform into amastigote-like forms (Fig. 5, A and B). Taken together, the results described in this section show that mitochondrial SODA is important for the differentiation of *L. amazonensis* promastigotes into amastigotes after both types of stimulation, low pH/high temperature and iron deprivation.

Stationary phase SODA/ΔsodA promastigotes are more susceptible to ROS stress and are impaired in metacyclogenesis

We next examined whether a single copy of SODA was sufficient to sustain growth and development during the promastigote stage. When grown in complete promastigote culture medium (containing iron and 10% FBS), SODA/ΔsodA promastigotes grew at a similar rate as WT during the early logarithmic phase of growth but entered stationary phase earlier at a density of 5–6 × 10$^7$ cells/ml (day 5). In contrast, WT parasites kept growing until reaching stationary phase at 7–8 × 10$^7$ cells/ml (day 7) (Fig. 5A). Moreover, after day 5, the SODA/ΔsodA parasite population showed a steady decline, a process that was partially reverted in the complemented SODA/ΔsodA + SODA line. These results suggest that SODA activity is also important for promastigote viability during the stationary phase of growth.

To investigate whether an impaired ability to detoxify mitochondrial ROS products due to SODA deficiency might cause promastigote death, we exposed WT, SODA/ΔsodA, and SODA/ΔsodA + SODA mid-log promastigotes to increasing concentrations of menadione, a drug that induces O$_2^-$ generation in mitochondria. SODA/ΔsodA parasites showed markedly higher sensitivity to the drug, with a drop in viability from >60% in WT to <30% in SODA/ΔsodA after exposure to 4 μM menadione. As expected, complemented SODA/ΔsodA + SODA promastigotes were more resistant to menadione toxicity when compared with...
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Figure 4. SODA/ΔsodA parasites fail to transform into amastigotes following the low pH/high temperature differentiation cue. Wild-type (WT), SODA single knock-out (SODA/ΔsodA), and complemented SODA single knock-out (SODA/ΔsodA + SODA) promastigote cultures were shifted to low pH (4.5)/high temperature (32 °C) culture conditions (0 h). Samples were taken at the indicated time intervals, assessed for viability by FDA staining, and for differentiation into the amastigote form by light microscopy morphological examination (loss of long flagella). A, number of viable cells in WT, SODA/ΔsodA, and SODA/ΔsodA + SODA cultures estimated at different time intervals following induction of low pH/high temperature mediated differentiation. B, percentage of viable rounded forms with a shortened flagellum in WT, SODA/ΔsodA, and SODA/ΔsodA + SODA cultured for 48 h in amastigote growth conditions. At least 200 FDA-stained parasites were counted in each sample. The data represent the mean ± S.D. of triplicate determinations and are representative of three independent experiments (**, p < 0.009). C, SEM analysis of parasite morphology in WT, SODA/ΔsodA, and SODA/ΔsodA + SODA cultures. Bars, 10 μm. D, TEM micrographs of WT, SODA/ΔsodA, and SODA/ΔsodA + SODA parasite sections. m = mitochondria; k = kinetoplast, and n = nucleus. Bars, 1 μm.

SODA/ΔsodA parasites (Fig. 6B). Collectively, our data suggest that normal levels of SODA expression are required for protection against the accumulation of mitochondrion-generated ROS during the late-log and stationary phases of promastigote growth.

Environmental stress (e.g. nutrient and oxygen deprivation) experienced by Leishmania promastigotes attached to the sand fly midgut is thought to be important for their transformation into infective metacyclic forms (29). However, very little is known about the actual signaling pathway involved in Leishmania metacyclogenesis in vivo. Considering our evidence for a role of SODA-mediated redox signaling in the promastigote-amastigote transition, and the increased SODA protein levels...
observed in stationary phase promastigotes in culture, we investigated whether SODA was also required for metacyclic promastigote development. The number of metacyclic forms was quantified on day 6 WT, SODA/ΔsodA, and SODA/ΔsodA + SODA stationary promastigote cultures after selective agglutination with an L. amazonensis promastigote-specific antibody. SODA/ΔsodA cultures showed ~3-fold reduction in the yield of metacyclics compared with WT, a phenotype partially restored by SODA complementation (SODA/ΔsodA + SODA) (Fig. 7A).

Scanning electron microscopy (SEM) analysis revealed a normal promastigote morphology for all three lines during the logarithmic phase of growth (day 3), but during stationary phase (day 7) the elongated and slender forms characteristic of metacyclic forms were only observed in WT and SODA/ΔsodA + SODA (Fig. 7B).

To assess mitochondrial function, we treated promastigotes with JC-1, a lipophilic cationic dye whose rate of accumulation inside mitochondria is directly dependent on the maintenance of an active mitochondrial membrane potential (ΔΨm) (17, 30). The shift of JC-1 from a monomeric form under low concentrations to an aggregated form inside mitochondria is detected as a shift from green (emission 530 nm) to red (emission 590 nm) fluorescence. Thus, the 590 nm/530 nm ratio in this assay provides an accurate quantification of the ΔΨm-dependent amount of dye imported into mitochondria. Increase in the 590 nm/530 nm fluorescence ratio was observed for all three lines on days 2 and 3 of culture, indicating healthy mitochondrial activity during the logarithmic phase of growth. However, SODA/ΔsodA parasites showed a significant reduction in the 590 nm/530 nm fluorescence ratio on day 7, an effect reversed by SODA complementation (Fig. 7C). This result suggests that SODA expression from a single allele is not sufficient to maintain normal mitochondrial function as the parasites enter the stationary phase of growth, when differentiation into virulent metacyclic forms is initiated.

Deletion of one SODA allele reduces mitochondrial SOD activity in stationary phase promastigotes

Whole-cell extracts from wild type, SODA/ΔsodA, and SODA/ΔsodA + SODA were prepared at different phases of promastigote growth and assayed for SOD activity. A reduction in the total SOD activity was observed in SODA/ΔsodA parasites in late-log (day 5) and stationary phase (day 7) (Fig. 8A). Because this biochemical activity assay does not distinguish between the mitochondrial SODA and the glycosomal SODB, it is important to note that SODA is the major SOD activity in Leishmania parasites. Since the SOD activity in WT was significantly higher than in SODA/ΔsodA, the observed reduction in SOD activity in SODA/ΔsodA confirms that SODA, and not SODB, is the major SOD isoenzyme in L. amazonensis.

To further investigate the role of SODA in the regulation of mitochondrial function, we used a mitochondrial-specific dye, MitoTracker Green, which stains live mitochondria with increased membrane potential. In WT promastigotes, the MitoTracker Green fluorescence intensity was higher, indicating a higher mitochondrial membrane potential compared to SODA/ΔsodA and SODA/ΔsodA + SODA (Fig. 8B). This suggests an important role for SODA in maintaining mitochondrial membrane potential during the stationary phase of growth.
we analyzed the same parasite extracts by immunoblotting with antibodies specific for each of the enzymes. As observed for the total SOD activity, SODA protein expression in the WT and SODA/ΔsodA + SODA parasites was similar and increased gradually as the cells entered stationary phase, whereas significantly less SODA was detected in SODA/ΔsodA extracts at the same time points (days 5 and 7) (Fig. 8B). In contrast, the levels of SODB protein were more stable and comparable between the three lines, showing only slightly elevated levels on days 5 and 7 (Fig. 8B). These results suggest that the abnormal mitochondrial function observed in SODA/ΔsodA promastigotes is likely to be a consequence of reduced SODA expression during the stationary phase of growth. This finding is in agreement with the mitochondrial localization of SODA (Fig. 1A) and with a lack of involvement of the glycosomal SODB isoform.

Consistent with this view, progression into the stationary phase of growth was associated with a reduction in both the total SOD activity and the amount of SODA protein in mitochondrial fractions isolated from SODA/ΔsodA parasites, when compared with WT (Fig. 9, A and B). As expected, only trace amounts of SODB were detected in the same mitochondrial fractions. Slightly elevated levels of SOD activity were observed in mitochondrial extracts from SODA/ΔsodA on day 3, possibly as a response to elevated ROS stress in SODA-deficient parasites during the logarithmic phase of growth.

Mitochondrial ROS generation is required for development of virulent forms

The data discussed above suggested that the inability of stationary phase SODA/ΔsodA promastigotes to efficiently differentiate into metacyclic forms might be due to inadequate generation of H$_2$O$_2$, which is known to function as an amastigote differentiation signaling molecule in L. amazonensis (16). However, an alternative explanation is that the differentiation defect
of electrons from the mitochondrial respiratory chain, thereby inhibiting production of all ROS, including $O_2^-$, $H_2O_2$, and peroxynitrite (31). Accordingly, we found that treatment of both WT and SODA/ΔsodA promastigotes with MitoTempo improved parasite survival during the stationary phase of growth (Fig. 10A). MitoTempo also restored the $\Delta\psi_m$ of the SODA/ΔsodA stationary phase promastigotes to wild-type levels (Fig. 10B). These effects on promastigote survival and mitochondrial membrane potential were observed with both low (15 $\mu M$) and high (50 $\mu M$) concentrations of MitoTempo. Interestingly, when we examined the yield of metacyclic forms, an improvement was observed after treatment of SODA/ΔsodA parasites with 15 $\mu M$ but not with 50 $\mu M$ MitoTempo (Fig. 10C). We observed a similar concentration-dependent effect in the expression of the $H_2O_2$ reporter protein APX; 15 $\mu M$ MitoTempo restored APX expression to WT levels, but such an effect was not seen after treatment with 50 $\mu M$. In WT parasites, both concentrations of MitoTempo caused >2-fold reduction in both the yield of purified metacyclics (Fig. 10C) and in APX protein levels (Fig. 10D). Thus, although mitochondrial dysfunction was restored in SODA/ΔsodA promastigotes treated with 50 $\mu M$ MitoTempo, this treatment did not increase the yield of metacyclic stage differentiation, suggesting that mitochondrial dysfunction is not the primary cause for the inability to differentiate SODA-deficient parasites. In WT parasites, where a normal complement of SODA may reduce the availability of $O_2^-$, MitoTempo may not stimulate $H_2O_2$ production, as indicated by the lower levels of APX expression under these conditions (Fig. 10D). Collectively, the results of these experiments reinforce the view that SODA-mediated $H_2O_2$ production plays a central role in the development of Leishmania virulent forms.

**Deletion of one SODA allele results in strong loss of virulence**

Metacyclic promastigotes from WT, SODA/ΔsodA, and SODA/ΔsodA + SODA promastigotes were purified from day 5 stationary cultures, and their viability was assessed with the dye FDA and compared for their ability to establish infections in mouse bone marrow macrophages (BMMs). Because of the reduced ability of SODA/ΔsodA to undergo metacyclogenesis, larger culture volumes were used to obtain comparable numbers of viable metacyclic forms. BMM infection with WT parasites progressed normally, and after the typical 24-h lag period...
the parasites replicated intracellularly as amastigotes, progressively increasing in number (Fig. 11A). In contrast, a steep decline in the number of intracellular parasites was observed with the SODA/sodA line between 3 and 24 h after BMM infection, with only very few intact parasites being detectable after 72 h. Episomal expression of SODA enhanced the ability of the parasites to survive intracellularly, as indicated by a markedly increased number of parasites detected after infection of BMM with the SODA/sodA line. The SODA complemented parasites were also able to replicate intracellularly, albeit at a slower rate when compared with the WT line (Fig. 11A). These results indicate that normal levels of SODA expression are necessary for L. amazonensis to establish successful infections in host macrophages.

We also examined the role of SODA in the parasite’s ability to induce cutaneous lesions in mice. Purified viable metacyclic forms were injected into the footpads of C57BL/6 mice, and the progression of cutaneous lesions was quantified over a period of 9 weeks (Fig. 11B). A steady growth in lesion size was recorded for WT L. amazonensis up to week 9. In contrast, mice infected with SODA/sodA metacyclics showed no evidence of lesion formation. Quantification of the parasite tissue load after 9 weeks showed a $>10^4$-fold higher number of parasites in the footpad tissues of mice injected with WT parasites, when compared with the group infected with the SODA/sodA line. This phenotype was also partially restored in the complemented SODA/sodA line (Fig. 11C). The inability to fully restore virulence was not unexpected, considering that lack of robust complementation is commonly observed in transgenic Leishmania (17, 34–38). These findings demonstrate that L. amazonensis must maintain normal levels of expression of the mitochondrial enzyme SODA to replicate inside host macrophages and establish cutaneous infections in vivo.

**Discussion**

Leishmania spp. are among the few organisms that can survive and replicate in the hostile environment of macrophage phagolysosomes, where degradative enzymes and ROS function as effective mechanisms of protection against pathogens. $O_2^-$ and ‘NO are two key ROS molecules generated by macrophages to neutralize invading microorganisms (39). To successfully evade this onslaught, Leishmania species have developed multiple adaptive features that include an antioxidant defense repertoire that includes trypanthione/trypanthione reductase,
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Figure 11. Deletion of a single SODA allele results in a marked loss of virulence that is partially restored by complementation with epoximally expressed SODA. The ability of metacyclic promastigotes purified from wild-type (WT), SODA single knock-out (SODA/ΔsodA), or SODA single knock-out complemented (SODA/ΔsodA + SODA) stationary phase cultures to infect macrophages (BMM) and generate cutaneous lesions in mice was compared. A, BMMs isolated from C57BL/6 mice were infected with metacyclics for 3 h, fixed immediately (3 h), or after further incubation for 24, 48, or 72 h, and the number of intracellular parasites was determined microscopically. The results are expressed as the percentage of infected macrophages counted at each time point, with the number of infected macrophages in the WT group set at 100%. The values obtained from five individual mice in each group. B, WT, SODA/ΔsodA, or SODA/ΔsodA + SODA macrophages were infected with metacyclics for 3 h, and lesion size was determined microscopically. The data correspond to the mean ± S.D. of triplicate determinations and are representative of more than three independent experiments. C, female C57BL/6 mice were inoculated in their left hind footpad with viable WT, SODA/ΔsodA, or SODA/ΔsodA + SODA metacyclics, and lesion development was determined by weekly caliper measurements. The results are expressed as the percentage of infected macrophages counted at each time point, with the number of infected macrophages in the WT group set at 100%. The values obtained from five individual mice in each group.

peroxidases, and three iron-dependent SODs (39–41). The Leishmania glycosomal iron-dependent SOD isoforms SODB1 and SODB2 are developmentally regulated, as indicated by reports of elevated SODB1 transcripts in amastigotes and of SODB2 in promastigotes, respectively (42, 43). Leishmania chagasi SODB1 null mutants are not viable, and parasite lines lacking one SODB1 allele have markedly reduced viability inside macrophages (42). The mitochondrial SOD isoform SODA was previously proposed to protect Leishmania mitochondria from oxidative stress (21, 44), but prior to this study its physiological function had not been investigated. Here, we investigated the physiological role of L. amazonensis SODA in light of recent evidence implicating this iron-dependent enzyme in ROS generation inside mitochondria, a process proposed to play a key role in the development of the virulent life cycle stages of Leishmania (17).

We previously identified and functionally characterized the L. amazonensis mitochondrial iron importer LMIT1 (17). Similar to what we report here for SODA, LMIT1 null mutants are not viable, consistent with the importance of iron for the assembly of Fe-S cluster proteins and ETC function. We also found that LMIT1/Δlmit1 promastigotes partially impaired in mitochondrial iron import are markedly defective in the ability to differentiate into infective amastigote forms. LMIT1/Δlmit1 parasites also showed a significant drop in mitochondrial iron content, a decrease in aconitase (a Fe-S cluster protein) and SOD activity, and extensive damage to mitochondria following oxidative stress. Based on these findings, we hypothesized that mitochondria is the major site where O$_2^*$ is generated and then converted by the iron-dependent SODA to H$_2$O$_2$, previously shown to act as a signal for differentiation (17). However, our initial studies of the role of SOD in L. amazonensis differentiation did not distinguish between the mitochondrial SODA and the glycosomal SODB1 and SODB2 isoforms. In this study, by measuring SOD activity in SODA–enriched mitochondrial fractions with little or no contaminating SODB, we clearly demonstrate that the >50% reduction in SOD activity observed in LMIT1/Δlmit1 L. amazonensis promastigotes in response to a differentiation stimuli can be attributed to SODA.

Our inability to generate SODA null mutants suggests that SODA is also essential for the long-term survival of L. amazonensis. This conclusion is consistent with the presence of functioning mitochondria in both promastigote and amastigote forms, and the well-established role of SOD in detoxifying ROS generated through electron leakage from the respiratory chain (6, 8). The critical role played by SODA in protecting mitochondria from endogenous ROS accumulation is further evident from the lethal phenotype we observed following RNAi-mediated knockdown of the SODA ortholog Tb927.5.3350 in procyclic forms of T. brucei. No phenotype was observed following RNAi-mediated silencing of the Tb927.5.3350 in bloodstream forms (18), in agreement with the dependence on active mitochondrial metabolism of T. brucei procyclics but not bloodstream forms, which possess only rudimentary mitochondria (27, 28).

SODA expression in L. amazonensis promastigotes increased progressively during culture reaching maximal levels on the stationary phase, presumably as a direct consequence of build up in oxidative stress. Such priming of the antioxidant defense system was proposed to facilitate virulence development, by preparing Leishmania parasites for the invasion of host macrophages (17, 26, 42). The ~50% reduction in both SODA activity and SODA protein levels following entry into stationary phase is probably responsible for the sudden decline in cell viability we observed in SODA/ΔsodA promastigote cultures. In addi-
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... tion, the slight but reproducible increase in SOD activity observed in early log phase cultures of SODA/ΔsodA parasites is consistent with a putative surge in O$_2^-$ levels during the initial phase of promastigote growth. In this scenario, impaired conversion of O$_2$ into H$_2$O$_2$ by SODA might explain the faster generation. Thus, the high mortality rate we...
RNAi-mediated knockdown

dsRNA synthesis was induced by the addition of 1 μg/ml tetracycline to cultures of clonal cell lines at 1 × 10⁶/ml starting concentration. Cells growing in the presence or absence of tetracycline were counted daily using a hemocytometer and diluted to the initial starting concentrations. SODA knockdown was confirmed by performing Western blot analyses of whole-cell lysates at different time points following tetracycline addition, using anti-SODA rabbit polyclonal antibodies raised against Leishmania SODA.

Generation of L. amazonensis SODA single knock-out cell lines

The L. amazonensis SODA ORF was genetically targeted for replacement with gene deletion constructs containing the hygromycin resistance gene or neomycin phosphotransferase through homologous recombination, as described earlier (17, 52). Sequences upstream and downstream of the SODA ORF were cloned using the following primers containing SfiI restriction enzyme sites (underlined): LamSODA 5’SfiI-A, FD (GGAGCCACCTAGGCCCGAAAGAGGGAGTTGTG), and Lam-SODA 5’SfiI-B, RV (GAGGCCCAGCGGCCGAGTAGTGTAGGTGCAGTTTTT), to amplify the 5’ sequence; LamSODA 3’SfiI-C FD, GAGGCTCTGTTGGCTGTGGTGTGGCAACCGC, and LamSODA 3’SfiI-D RV, and GAGGCCT-GACTGGCCGCTGACAACTGCACG for 3’ UTR. Four-part ligation using the PCR-amplified 5’- and 3’-flanking sequences, drug resistance cassettes, and the plasmid backbone was carried out. Positive clones were identified by analyzing SfiI restriction digests of plasmid DNA samples and confirmed by sequencing with specific primers as described (52). The targeting fragment used to transfect L. amazonensis promastigotes was released by PacI digestion, gel-purified, and used for electroporation. Leishmania clones with a single SODA allele deletion (SODA/Δsoda) were isolated based on the ability of transformants to grow on agar plates containing hygromycin (100 μg/ml) or neomycin (50 μg/ml) and analyzed by Southern blotting and PCR to verify integration of the drug cassette in the desired location.

For generation of a rescue plasmid expressing Leishmania SODA with C-terminal hemagglutinin tag, a two-step PCR amplification strategy was employed. In the first round, a 693-bp fragment of the SODA ORF was amplified with primers SODA HA, FD, AACCCGGGACATAATGTTCGGCCGT-GTCTCGATG (Small site underlined), and SODA HA, RV, CTGGGACGTCTGATTTGGTACACGCTTGTCGG, that allowed for removal of the endogenous stop codon and introduction of an in-frame HA tag. The PCR product was used as template in a second round of amplification using SODA-HA, FD, as sense and HA TAG2, RV, TTTGATCTTTAAGCG-TAGTCTGGGACGTCTGATG (BamHI site underlined), as antisense primers. The final PCR products were digested with BamHI and Smal and cloned into pXG-SAT (courtesy of Prof. S. Beverley, Washington University). Transfected Leishmania clones were selected in plates containing 50 μg/ml nourseothricin (Jena Biosciences), and expression of HA-tagged SODA was confirmed by Western blotting.

Expression of recombinant SODA, SODB, and APX proteins and generation of antibodies

To produce recombinant histidine-tagged SOD proteins, the genes encoding SODA, SODB, and APX were PCR-amplified from L. amazonensis genomic DNA using primers SODAexp, FD (GGCATATGTTCCGCTTGTCGTC), and SODAexp RV, (GGAAAGCTTCTCTGTGGCCTTTTC) for SODA; SODBexp FD (AAATATGCGTTTCGCTTCCAGCCTGCT), and SODBexp RV (TTAAGCTTTCAGATCTGTGGCAGTGA-TGG) for SODB; and APXexp FD (CAGACATGTTCG-GACCTTCGCG) and APXexp RV (TTCAAGCTTGGCT-CGGACCGCGG) for APX. Forward primers for SODA and SODB contained an Ndel restriction site, and the APX gene contained a PciI site, contiguous with the start codon. Reverse primers were engineered to include a HindIII restriction site that removed the endogenous stop codon and allowed for synthesis of a six-histidine tag. The resulting PCR products were cloned in the pET28b (+) expression plasmid (Novagen) using Ndel and HindIII restriction sites, and the resulting plasmids were used to transform Escherichia coli BL21(DE3)pLysS (Novagen). Expression of His₆-tagged SODA, SODB, and APX protein in soluble form was achieved by inducing transformed E. coli strains with 0.1 mM isopropyl 1-thio-β-ß-galactosidase (overnight at room temperature in media supplemented with 2% ethanol). His₆-tagged SODA, SODB, or APX proteins were purified from bacterial cell extracts by nickel column chromatography with His-60 nickel superflow resin (Clontech) followed by elution with 0.3 M imidazole buffer, according to the manufacturer’s protocol. Homogeneity of the purified proteins was ascertained by SDS-PAGE.

Polyclonal antibodies were raised against purified SODA, SODB, and APX proteins by periodic injection of rabbits with purified protein samples (the Pocono Rabbit Farm and Laboratory). Specificity of the antisera was assessed by Western blotting.

Isolation of mitochondrial fractions

Mitochondrial enrichment was performed as described previously (17) using 5 × 10⁶ promastigotes from stationary phase cultures. The cells were washed three times with MES buffer (20 mM MOPS (pH 7.0), 250 mM sucrose, and 3 mM EDTA) and resuspended in 500 μl of MES supplemented with 1 mg/ml digitonin and protease inhibitor mixture (Roche Applied Science). Following a 5-min incubation at room temperature, the cell suspensions were centrifuged for 5 min (10,000 × g at 4 °C), and the supernatant was collected as the cytoplasmic fraction. The pellet was washed once with MES buffer and used for further analysis as the mitochondrial fraction or stored at −80 °C until further use.

Determination of SOD activity in whole-cell and mitochondrial extracts

SOD activity was determined in whole-cell extracts as described earlier (16, 17). To estimate mitochondrial SOD activity, mitochondrial fractions obtained following subcellular fractionation as described above were resuspended in buffer A and lysed by sonication. SOD activity in the lysates was measured using the SOD Assay Kit-WST (Dojindo Molecular
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Technologies, Inc.) according to the manufacturer's protocol. Standard curves were generated using known concentrations of horseradish SOD (Sigma). Protein content was determined using BCA™ protein assay kit (Thermo Fisher Scientific).

Localisation of SODA by immunofluorescence microscopy

Immunolocalization of SODA was performed as described previously (17). To confirm the mitochondrial localization of SODA, promastigotes were incubated with MitoTracker Red CMXRos (Invitrogen) followed by fixation with 4% paraformaldehyde and attachment to poly-l-lysine-coated slides (multitest 8-well; MP Biomedicals). Following treatment with 50 mM NH₄Cl, the cells were permeabilized with 0.1% Triton X-100 in PBS, blocked with PBS 5% horse serum and 1% bovine serum albumin (BSA) for 1 h at room temperature, and incubated with anti-SODA rabbit polyclonal antibodies (1:10,000 dilution in PBS 1% BSA) for 1 h followed by anti-rabbit IgG AlexaFluor 488 (Invitrogen) 1:5000 dilution in PBS 1% BSA for 1 h, and staining with 2 μg/ml DAPI for 1 h. Slides were mounted with ProLong Gold antifade reagent (Invitrogen); images were acquired through a Deltavision Elite Deconvolution microscope (GE Healthcare) and processed using Volocity Suite (PerkinElmer Life Sciences).

Assays for mitochondrial activity

Mitochondrial membrane potential (ΔΨm) was estimated using the MitoProbe JC-1 assay kit (Invitrogen). 1 x 10⁷ promastigotes were incubated with 10 μM JC-1 for 15 min at 27 °C, washed, and resuspended in PBS. Fluorescence measured at 530 and 590 nm using a SpectraMax M5® microtiter plate reader (Molecular Devices) was used to determine the ΔΨm (530:590 ratio).

To visualize the mitochondrial staining pattern, promastigotes were plated in glass-bottom dishes (MatTek Corp.) for live imaging on a Nikon Eclipse Ti inverted microscope with a C9100-50 camera and mCherry and FITC filters. Acquired images were analyzed with the Volocity Software Suite (PerkinElmer Life Sciences).

Electron microscopy

For scanning EM, parasites fixed in 2.5% (v/v) glutaraldehyde in 0.1 M sodium cacodylate buffer (pH 7.4) for 60 min and attached to poly-l-lysine-coated coverslips were rinsed briefly with PBS, fixed with 0.1 M cacodylate buffer (pH 7.4), treated with osmium tetroxide for 1 h, acetone-dehydrated, and critical point dried from CO2. After sputter coating with Au/Pd, the preparations were imaged in an Amray 1820D scanning electron microscope. For transmission EM, fixed parasite cells were post-fixed with osmium tetroxide, and cell sections were prepared as described before (38). Final images were obtained using a Zeiss EM10CA electron microscope.

Quantification of Leishmania intracellular growth in macrophages

Macrophage infection assays were carried out as described previously (16, 17, 37, 38). A total of 1 x 10⁶ BMMs from C57/BL6 mice (Charles River Laboratories), plated on glass coverslips in 3-cm dishes 24 h prior to the experiment, were infected with a 1:5 multiplicity of infection with metacyclic forms purified from stationary phase promastigote cultures (5-day-old) using the m3A1 monoclonal antibody (53). After allowing 3 h for invasion, BMMs were washed three times in PBS and incubated for the indicated times at 34 °C. Coverslips were retrieved after 3 (initial infection) 24, 48, and 72 h of incubation, fixed in 4% PFA, permeabilized with 0.1% Triton X-100 for 10 min, and stained with 10 μg/ml DAPI for 1 h. The total number of macrophages and the total number of intracellular parasites per microscopic field (100 X N.A. 1.3 oil immersion objective, Nikon E200 epifluorescence microscope) were determined, and the results expressed as intracellular parasites per 100 macrophages. At least 300 host cells, in triplicate, were analyzed for each time point. The data were analyzed for statistical significance using an unpaired Student’s t test (p < 0.05 was considered significant).

In vivo virulence and parasite load estimation

A total of 1 x 10⁶ infective metacyclics purified from WT, SODA/Δsoda, or SODA/Δsoda + SODA stationary phase promastigote cultures and resuspended in a volume of 50 μl of PBS were used to inoculate 6-week-old female C57BL/6 mice (n = 5 per group) in the left hind footpad. Progression of footpad lesion development was monitored through weekly measurements with a caliper (Mitutoyo Corp., Japan), quantitating the difference between the left and right hind footpads. The parasite load was estimated in infected tissue collected from footpads of mice sacrificed 11 weeks post-infection using a limiting dilution assay (54).

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References

1. World Health Organization (2010) Control of the leishmaniases. World Health Organ. Tech. Rep. Ser., 2010, xii–xiii, 1–186
2. Bates, P. A. (2008) Leishmania sand fly interaction: progress and challenges. Curr. Opin. Microbiol. 11, 340–344
3. Rosenzweig, D., Smith, D., Oppedorpes, F., Stern, S., Olafson, R. W., and Zilberstein, D. (2008) Retooling Leishmania metabolism: from sand fly gut to human macrophage. FASEB J. 22, 590–602
4. Tsiganov, P., Gherardini, P. F., Helmer-Citterich, M., and Zilberstein, D. (2012) What has proteomics taught us about Leishmania? Parasitology 139, 1146–1157
5. Lahav, T., Sivam, D., Volpin, H., Ronen, M., Tsiganov, P., Green, A., Holland, N., Kuzyk, M., Borchers, C., Zilberstein, D., and Myler, P. J. (2011) Multiple levels of gene regulation mediate differentiation of the intracellular pathogen Leishmania. FASEB J. 25, 515–525
SODA-mediated redox signaling promotes Leishmania virulence

13. Chandel, N. S. (2015) Evolution of mitochondria as signaling organelles. Cell Metab. 22, 204–206
16. Mittra, B., Cortez, M., Haydock, A., Ramasamy, G., Myler, P. J., and Andrews, N. W. (2013) A trypanosomatid iron transporter that regulates mitochondrial function is required for Leishmania amazonensis virulence. PLoS Pathog. 12, e1005340
18. Taylor, M. C., and Kelly, J. M. (2010) Iron metabolism in trypanosomatids. Mol. Biochem. Parasitol. 154, 62–69
20. Wilson, M. E., Andersen, K. A., and Britigan, B. E. (1994) Response of Leishmania chagasi promastigotes to oxidant stress. Infect. Immun. 62, 5133–5141
21. Alzate, J. F., Arias, A. A., Moreno-Mateos, D., Alvarez-Barrientos, A., and Jiménez-Ruiz, A. (2007) Mitochondrial superoxide mediates heat-induced apoptotic-like death in Leishmania infantum. Mol. Biochem. Parasitol. 152, 192–202
22. Riemann, A., Schneider, B., Ihling, A., Nowak, M., Sauvant, C., Thews, O., and Gekle, M. (1994) The role of pH and temperature in the development of Leishmania parasites. Annu. Rev. Microbiol. 48, 449–470
23. Getachew, F., and Gedamu, L. (2007) Leishmania donovani iron superoxide dismutase A is targeted to the mitochondria by its N-terminal positively charged amino acids. Mol. Biochem. Parasitol. 154, 62–69
24. Boitz, J. M., Strasser, R., Yates, P. A., Jardim, A., and Ullman, B. (2013) Adenylosuccinate synthetase and adenylosuccinate lyase deficiencies trigger growth and infectivity deficits in Leishmania donovani. J. Biol. Chem. 288, 8977–8990
25. da Silva, M. F., Zampieri, R. A., Muxel, S. M., Beverley, S. M., and Foeter-Winter, L. M. (2012) Leishmania amazonensis arginase compartmentalization in the glycosome is important for parasite infectivity. PLoS ONE 7, e34022
26. Pal, S., Dolai, S., Yadav, R. K., and Adak, S. (2010) Ascorbate peroxidase from Leishmania major controls the virulence of infective stage of promastigotes by regulating oxidative stress. PLoS ONE 5, e11271
27. van Weelden, S. W., Fast, B., Vogt, A., van der Meer, P., Saas, J., van Hellemont, J. J., Tielen, A. G., and Boshart, M. (2003) Procyclic Trypanosoma brucei do not use Krebs cycle activity for energy generation. J. Biol. Chem. 278, 12854–12863
28. Bringuè, F., Rivière, L., and Coustou, V. (2006) Energy metabolism of trypanosomatids: adaptation to available carbon sources. Mol. Biochem. Parasitol. 149, 1–9
29. Bates, P. A. (2007) Transmission of Leishmania metacyclic promastigotes by phlebotomine sand flies. Int. J. Parasitol. 37, 1097–1106
30. Mehta, A., and Shaha, C. (2004) Apoptotic death in Leishmania donovani promastigotes in response to respiratory chain inhibition: complex II inhibition results in increased pentamidine cytotoxicity. J. Biol. Chem. 279, 11798–11813
31. Dikalov, S. (2011) Cross-talk between mitochondria and NADPH oxido-dases. Free Radic. Biol. Med. 51, 1289–1301
32. Nazarewicz, R. R., Dikalova, A., Bikineyeva, A., Ivanov, S., Kirilyuk, I. A., Grigor’ev, I. A., and Dikalov, S. I. (2013) Does scavenging of mitochondrial superoxide attenuate cancer prosurvival signaling pathways? Antioxid. Redox Signal. 19, 344–349
33. Murphy, M. P., and Smith, R. A. (2007) Targeting antioxidants to mitochondria by conjugation to lipophilic cations. Annu. Rev. Pharmacol. Toxicol. 47, 629–656
34. Gaur, U., Shovalter, M., Hickerson, S., Dalvi, R., Turco, S. J., Wilson, M. E., and Beverley, S. M. (2009) Leishmania donovani lacking the Golgi GDP-man transporter LP2G exhibit attenuated virulence in mammalian hosts. Exp. Parasitol. 122, 182–191
35. Owusu-Ansah, E., and Banerjee, U. (2009) Reactive oxygen species prime mitochondrial dismutase A is targeted to the mitochondria by its N-terminal positively charged amino acids. Mol. Biochem. Parasitol. 175, R453–R462
36. Huynh, C., Sacks, D. L., and Andrews, N. W. (2006) A Leishmania amazonensis ZIP family iron transporter is essential for parasite replication within macrophage phagolysosomes. J. Exp. Med. 203, 2363–2375
37. Flannery, A. R., Huynh, C., Mittra, B., Mortara, R. A., and Andrews, N. W. (2011) LFR1 ferric iron reductase of Leishmania amazonensis is essential for the generation of infective parasite forms. J. Biol. Chem. 286, 23266–23279
38. Miguel, D. C., Flannery, A. R., Mittra, B., and Andrews, N. W. (2013) Heme superoxide dismutase A is targeted to the mitochondria by its N-terminal positively charged amino acids. Mol. Biochem. Parasitol. 154, 62–69
50. Oberholzer, M., Lopez, M. A., Ralston, K. S., and Hill, K. L. (2009) Approaches for functional analysis of flagellar proteins in African trypanosomes. Methods Cell Biol. **93**, 21–57

51. Redmond, S., Vadivelu, J., and Field, M. C. (2003) RNAit: an automated web-based tool for the selection of RNAi targets in *Trypanosoma brucei*. Mol. Biochem. Parasitol. **128**, 115–118

52. Fulwiler, A. L., Soysa, D. R., Ullman, B., and Yates, P. A. (2011) A rapid, efficient and economical method for generating leishmanial gene targeting constructs. Mol. Biochem. Parasitol. **175**, 209–212

53. Pinto-da-Silva, L. H., Fampa, P., Soares, D. C., Oliveira, S. M., Souto-Padron, T., and Saraiva, E. M. (2005) The 3A1-La monoclonal antibody reveals key features of *Leishmania (L.) amazonensis* metacyclic promastigotes and inhibits procyclics attachment to the sand fly midgut. Int. J. Parasitol. **35**, 757–764

54. Tabbara, K. S., Peters, N. C., Afrin, F., Mendez, S., Bertholet, S., Belkaid, Y., and Sacks, D. L. (2005) Conditions influencing the efficacy of vaccination with live organisms against *Leishmania major* infection. Infect. Immun. **73**, 4714–4722

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53. Pinto-da-Silva, L. H., Fampa, P., Soares, D. C., Oliveira, S. M., Souto-Padron, T., and Saraiva, E. M. (2005) The 3A1-La monoclonal antibody reveals key features of *Leishmania (L.) amazonensis* metacyclic promastigotes and inhibits procyclics attachment to the sand fly midgut. Int. J. Parasitol. **35**, 757–764

54. Tabbara, K. S., Peters, N. C., Afrin, F., Mendez, S., Bertholet, S., Belkaid, Y., and Sacks, D. L. (2005) Conditions influencing the efficacy of vaccination with live organisms against *Leishmania major* infection. Infect. Immun. **73**, 4714–4722

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