**Abstract**

The phlebotomine sand fly *Lutzomyia longipalpis* is the most important vector of American visceral leishmaniasis (AVL), the disseminated and most serious form of the disease in Central and South America. In the natural environment, most female *L. longipalpis* are thought to survive for less than 10 days and will feed on blood only once or twice during their lifetime. Successful transmission of parasites occurs when a *Leishmania*-infected female sand fly feeds on a new host. Knowledge of factors affecting sand fly longevity that lead to a reduction in lifespan could result in a decrease in parasite transmission. Catalase has been found to play a major role in survival and fecundity in many insect species. It is a strong antioxidant enzyme that breaks down toxic reactive oxygen species (ROS). Ovarian catalase was found to accumulate in the developing sand fly oocyte from 12 to 48 hours after blood feeding. Catalase expression in ovaries as well as oocyte numbers was found to decrease with age. This reduction was not found in flies when fed on the antioxidant ascorbic acid in the sugar meal, a condition that increased mortality and activation of the prophenoloxidase cascade. RNA interference was used to silence catalase gene expression in female *Lu. longipalpis*. Depletion of catalase led to a significant increase of mortality and a reduction in the number of developing oocytes produced after blood feeding. These results demonstrate the central role that catalase and ROS play in the longevity and fecundity of phlebotomine sand flies.

**Introduction**

The phlebotomine sand fly *Lutzomyia longipalpis*, Lutz and Neiva 1912 is the best studied and most important vector of American Visceral Leishmaniasis (AVL) [1,2]. It is widely found in Latin America, from the South of Mexico to the North of Argentina [3,4]. *Lu. longipalpis* is a permissive vector to *Leishmania* infections [5] and this together with its wide distribution in urban environments [6] makes this sand fly species an ideal model to study phlebotomine physiology to help develop future vector control methods for the disruption of *Leishmania* transmission.

Previous studies in other blood-feeding insect vectors have shown that reactive oxygen species (ROS) play an important role in both reproductive output [7] and survival [8,9,10,11,12]. Biological damage related to ROS production has also been implicated in the process of aging in dipterans like *Drosophila melanogaster*, previous work done on this species showed that oxidative stress increases with age, while antioxidant enzyme activity decreased over time [13,14,15].

ROS are regularly generated by mitochondrial electron transport [16]. Partially reduced and highly reactive metabolites of O2 such as superoxide anion (O2·−) and hydrogen peroxide (H2O2) are formed during cellular respiration. These partially reduced metabolites of O2 are often referred to as reactive oxygen species due to their higher reactivity in relation to molecular oxygen [17]. Excessive release of ROS damages lipids, proteins, and DNA [18] which leads to oxidative stress, loss of cell function, and programmed cell death [19]. ROS are also actively released as a response against bacterial and parasitic pathogens in different insect species [8,20,21]. To regulate oxidative stress, the eukaryotic cell produces different ROS-scavenging enzymes, such as superoxide dismutase (which reduces O2·− to H2O2), glutathione peroxidase and catalase (which reduces H2O2 to H2O) [17]. Although many studies have been published regarding mechanisms to resist oxidative stress in *Leishmania* parasites [22,23,24,25] there is little information on ROS-scavenging molecular mechanisms on sand flies, despite the fact that putative antioxidant enzymes have been found to be upregulated in two different species of phlebotomine sand flies upon *Leishmania* infection [26,27].

In order to investigate the biological role of ROS-scavenging in fecundity and survival, we analysed the expression of catalase in three different age groups of female sand flies. Fecundity and catalase expression decreased with age. Catalase was incriminated as an important component in the loss of fecundity using RNAi. Dietary supplementation with an exogenous ROS-scavenger was found to partially reverse the differences in fecundity and increase mortality with a concomitant activation of the phenoloxidase (PO)
cascade. These results show that both fecundity and survival are affected by endogenous and exogenous ROS-scavenging in female *Lutzomyia longipalpis*.

**Results**

**Age-related decrease of fecundity**

To evaluate the effect of ageing in fecundity, females of different age groups were blood fed and dissected to examine difference in developing oocyte numbers. Female *Lu. longipalpis* from the older age group showed a decrease in the number of developing oocytes dissected five days after blood feeding in comparison to younger sand flies (fig. 1). Female *Lu. longipalpis* that were bloodfed 3 and 6 days post-emergence (PE) showed no significant difference in oocyte numbers. However, sand flies bloodfed at 9 days PE showed a significant decrease in oocyte numbers after dissecting 5 days after blood feeding (fig. 1; \( P < 0.005 \), ANOVA).

**ROS-scavenging reverses age related loss of fecundity**

To evaluate the role of ROS scavenging in age-related decrease of fecundity, 9 day old female *Lu. longipalpis* were fed a sucrose meal supplemented with a ROS-scavenger upon emergence until end of the experiment. Sand flies were offered a 70% sucrose solution supplemented with 20 mM ascorbic acid and subsequently blood-fed on day 9 PE. 20 mM ascorbic acid was chosen after evaluating mortality of sand flies when offered 100, 50, 20, 10 and 5 mM ascorbic acid in 70% sucrose (data not shown). The number of developing oocytes dissected 5 days after blood feeding was significantly higher (fig. 2; \( P < 0.0001 \), t-test) in sand flies that received a sugar meal supplemented with 20 mM ascorbic acid in comparison to control sand flies fed on 70% sucrose solution. This suggests that exogenous ROS-scavenging can reverse age-related loss of fecundity in sand flies blood fed 9 days PE.

**Catalase activity is reduced in developing oocytes of older flies and ROS scavengers reverse catalase depletion**

Flies were assayed at 24 h and 48 h to find out if catalase accumulated in developing oocytes. Ovaries of *Lu. longipalpis* dissected 6 days PE contained higher catalase enzymatic activity at 48 h compared to 24 h after blood feeding (fig. 3A; \( P < 0.0001 \), t-test). Moreover, mRNA expression of catalase increased with oocyte development from 12 to 48 hours after blood feeding (fig. 3B).

To further understand the role of endogenous ROS-scavenging and ageing, flies from different age groups were assayed for catalase *LlonKat1* expression. Flies from different age groups (3, 6 and 9 days PE) showed a decrease in expression in ovaries dissected at 48 hours after blood feeding (fig. 3C; \( P = 0.001 \), ANOVA). Interestingly, when 9 day old sand flies were fed with a 20 mM ascorbic acid supplemented sugar meal, catalase *LlonKat1* mRNA expression was significantly higher compared to flies of the same age fed on sucrose only (fig. 3C; \( P < 0.002 \), t-test). The results show that a) catalase accumulates in the developing oocyte as shown by increase in enzymatic activity and relative expression, b) catalase expression is age-dependent and is lower in older flies and c) the dietary supplementation with an exogenous ROS-scavenger increases catalase expression in older flies.

*Lutzomyia longipalpis* catalase sequence was already described [26,27] and was retrieved from the GenBank (ABV60342.1). It codes for a protein (named *LlonKat1* in this study) with molecular
mass of 57682 Da and isoelectric point of 8.28, without a signal peptide and mitochondrial or peroxisomal targeting sequences of types 1 and 2. LlonKat1 has high identity (ranging from 46–73%) to catalase sequences from other insects, crustaceans, yeast and mammals and lower identity to the bacterial catalase from *Pseudomonas syringae* (fig. 4). LlonKat1 sequence contains the conserved residues His73 and Asn147 (catalytic), Ser113, Val115, Phe152, Phe160, Leu298, Met349, Arg353, Tyr357 (heme binding/coordination) and His193, Arg202, Ile301, Gln304 (putative NADPH binding pocket) (fig. 4).

Catalase gene RNAi mediated depletion leads to a decrease in sand fly fecundity

The gene sequence of *Lu. longipalpis* catalase was obtained from a cDNA library constructed from sand fly whole bodies (NSFM-142e04.q1k [27]) and aligned with a previous described catalase obtained from *Lu. longipalpis* midguts (GenBank Accession number: EU124624.1), showing a high level of identity (99%) and similarity (99%) (fig. S1). As any other sequence from *Lu. longipalpis* was identified as putative catalase, either in the whole body or midgut cDNA libraries, this gene is probably a single copy gene in *Lu. longipalpis*. To confirm the role of endogenous ROS-scavenging in fecundity catalase was depleted using RNAi. Flies injected with 144 ng of dsRNA for catalase (dsCAT) showed a dramatic decrease in oocyte number dissected 48 hours after blood feeding (fig. 5A; \( P < 0.005 \), ANOVA) compared to sand flies injected with a non-related dsRNA (dsGFP) and uninjected sand flies. A change in appearance of ovaries was observed during dissections with matured ovaries, in sand flies injected with dsRNA for catalase, appearing underdeveloped in comparison to both mock-injected and uninjected controls (fig. 5B). A dsRNA-mediated significant reduction in catalase expression in whole flies was observed by RT-PCR (fig. 5C), with no effects on the catalase expression in the midgut (data not shown). These results confirm that endogenous ROS-scavenging in developing oocytes plays a major role in female *Lu. longipalpis* fecundity.

Effect of ROS-scavenging in the survival of sand flies

To evaluate the role of exogenous ROS scavenging in survival, female *Lu. longipalpis* were fed with an antioxidant-supplemented sugar meal upon emergence. Mortality was recorded from day 1 PE up to day 7 PE. Survival curves depict an increase in mortality due
to exogenous ROS-scavenging by an exogenous antioxidant (fig. 6A). In order to assess whether the higher mortality rate was related to an effect on sand fly immune homeostasis, phenoloxidase (PO) activity was measured in control and antioxidant-supplement ed females. Spontaneous PO is defined as the activity measured upon reaction with 3,4 dihydroxy-DL-phenylalanine (DOPA), and corresponds to the enzyme that is already activated in physiological conditions and total activity was the activity observed after in vitro activation of the enzyme, by preincubating the sample with bovine trypsin. Sand flies fed on ascorbic acid-supplemented sucrose showed a significant increase in spontaneous PO (fig. 6B; P \(, 0.05 \), t-test) but no difference in total PO activity.

To further investigate if ROS-scavenging was implicated in increased mortality, catalase LlonKat1 was depleted via RNAi injection in female \( \text{Lu. longipalpis} \) and mortality was recorded from day 1 PE up to day 7 PE. Mortality rates was higher in knocked down (dsCAT) sand flies (fig. 7), compared to flies injected with a non-related dsRNA (dsGFP) and non-injected flies. These results show that ROS-scavenging by either endogenous or exogenous antioxidants play an important role in female \( \text{Lu. longipalpis} \) survival.

**Discussion**

The present study suggests for the first time that catalase-mediated ROS scavenging has a significant impact on female \( \text{Lu. longipalpis} \) fecundity and survival. Female \( \text{Lu. longipalpis} \) from different age groups showed differences in developing oocytes numbers, with the oldest (9 days PE) presenting the lowest number of oocytes (fig. 1). The age-related loss of fecundity could be reversed with dietary supplementation of a potent exogenous ROS-scavenger (fig. 2). This underlines the importance of catalase in the reproductive success of blood sucking dipterans. Evidence from other dipterans show that aging results in increase of oxidative stress and loss of enzymatic antioxidant efficiency \[13,14,28\]. Moreover, inactivation or silencing of catalase in \( \text{Drosophila melanogaster} \) \[29\], \( \text{Musca domestica} \) \[30\], \( \text{Rhodnius prolixus} \) \[31\] and \( \text{Anopheles gambiae} \) \[9\] led to increased mortality due to increase in ROS levels. It is likely that accumulation of ROS in older flies could account for the decrease of female sand fly fecundity due to an increase in oxidative stress, loss of antioxidant enzymatic efficiency or both. In \( \text{An gambiae} \), fecundity of female...
mosquitoes declined with age, with reduction of number of eggs oviposited and number of larvae hatched per female [7]. We did not measure differences in fecundity in terms of larval develop-

Figure 5. RNAi-mediated depletion of catalase LlonKat1 in female Lu. Longipalpis and its effect on fecundity. (A) Average number of developing oocytes dissected 48 hours days after blood meal ± SEM (combined samples derived from 2 independent experiments). Asterisk indicates statistical significance at \( P<0.005 \) (ANOVA). (B) Relative development of female Lu. longipalpis ovaries observed upon catalase gene knockdown by RNAi, in comparison to mock-injected and uninjected control sand flies. Bar = 1 mm. (C) Relative expression of catalase LlongKat1 mRNA in whole fly homogenates from dsRNA-injected catalase knock-down sand flies. Bar charts represent mean ± SEM of combined samples from at least 2 independent experiments. Asterisk indicates statistical difference at \( P<0.05 \) (ANOVA).

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Figure 6. Effect of dietary supplementation of ascorbic acid on mortality of sugar fed Lu. Longipalpis. (A) Female sand flies were offered a 70% sucrose solution supplemented with 20 mM ascorbic acid or a non-supplemented sucrose solution. Experimental flies (sucrose + 20 mM ascorbic acid) exhibited a significantly lower survival rate compared to control flies, \( (p<0.001, \) Kaplan-Meier, Log Rank \( \chi^2 \) test). (B) Spontaneous and total phenoloxidase (PO) activity in Lu. longipalpis females after 7 days of feeding with 70% sucrose solution or 70% sucrose solution supplemented with 20 mM ascorbic acid. Spontaneous PO activity in ascorbic acid supplemented flies was significantly higher than control flies \( (P<0.05, T\)-test). Results are mean ± SEM from 2 independent experiments with 10 sand flies per experiment.

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ment but it is likely that the age-related differences in fertility would have resulted in less viable larvae being produced from older flies, as they would be presumably exposed for a longer periods to oxidative damage.

Catalase enzymatic activity as well as catalase LlonKat1 mRNA relative expression increased in the ovaries of older female sand flies (6 days PE) after the blood feeding (figs. 3a and b). Protein expression and accumulation increased upon blood feeding in maturing ovaries of mosquitoes due to nutrient allocation for egg production [32,33]. It has been shown in different insect species that antioxidant activity increases in the ovaries to protect the embryo from oxidative damage [34,35]. It is conceivable that such
accumulation of catalase in sand fly ovaries also provides the means to protect developing eggs from oxidative damage. Additional support for this hypothesis was given by the dramatic decrease in developing oocyte numbers upon successful catalase gene depletion by RNAi in female sand flies (fig. 5A and b).

Interestingly, oral delivery of ascorbic acid seemed to stimulate catalase LlonKat1 mRNA expression in older flies to levels similar to that of younger flies (fig. 3C). It has been shown that age-related accumulation of ROS/oxidative stress leads to loss of efficiency in cellular processes [13,14,15,28,36], therefore it is possible that ROS-scavenging by an exogenous antioxidant slowed or lowered such deleterious effects in either catalase LlonKat1 mRNA or in other molecules involved in its upregulation. On the other hand, it has been shown that ascorbate is a potent inhibitor of catalase [37], the inhibition being independent of substrate concentration and pH and strongly influenced by temperature. Furthermore, catalase incubation with ascorbate leads to degradative changes to the catalase molecule [38]. In our experiments, the increase in catalase gene expression might reflect a compensation response to replenish normal catalase levels in the sand fly body after catalase was degraded, by an unknown mechanism, during ascorbic acid supplementation with the sugar meal.

Catalase LlonKat1 does not have a signal peptide or targeting subunits with molecular mass ranging from 43–75 kDa [40], which is consistent with LlonKat1 monomer predicted molecular mass (57.7 kDa). All conserved catalytic and heme binding residues are present in LlonKat1 sequence, suggesting a full catalytic activity, and the presence of residues Leu298, Met349 indicate that His70 is above the ring III of the hemeprotein molecule (His-III orientation), as seen in other clade 3 catalases [41].

ROS-scavenging by dietary supplementation of ascorbic acid (fig. 6a) led to a reduction in sand fly survival. When antioxidants were provided to a susceptible strain of *Anopheles gambiae* to *Plasmodium* infection, a similar but more drastic effect was observed with female mosquitoes [7]. Magwere et al. [42] observed that antioxidant supplementation did not extend the lifespan of wild type *Drosophila*. Similarly, Bayne et al. [43] showed that overexpression of MnSOD and catalase, despite protecting *Drosophila* from oxidative stress, were detrimental for lifespan and physical fitness of the insects. Kang et al. [44] observed a reduction in the lifespan of *Anopheles stephensi* when the mosquitoes were blooded with the antioxidant MnTBAP in comparison with the buffer control. It has been hypothesized that a minimal level of ROS might be required to maintain the balance of the gut microbiota and that a baseline level of ROS activity might be crucial for basic midgut physiology. Previous studies done with other dipteran species had showed that ROS release constitutes a first line of defence against pathogens in the midgut [45]. Experiments in *D. melanogaster* have demonstrated the existence of a midgut-specific active ROS releasing system against orally delivered bacteria [8]. In the present study, higher activities of spontaneous PO were recorded and this might be due to an increase in microbial infection associated with sand flies that fed on an ascorbic acid-supplemented sugar meal. In insects, PPO activation is often related to bacterial or fungal infections [45]. Since only the soluble form of PO was measured (see Material and Methods), it is more likely that the activity was related to the immune response rather than to the melanisation of the adult cuticle or egg shell. PO has already been described in gut tissues or adhered hemocytes in other dipterans [46]. It is possible that mortality in our experimental group fed with sucrose supplemented with ascorbic acid may be due to a decrease in ROS production inside the midgut and that ROS activity, similar to the events in certain strains of mosquitoes, may play a role in sand fly immunity towards opportunistic microbes or be involved in important cellular signalling pathways [47,48].

There is evidence of other antioxidant enzymes with catalase-like functions found in the sand fly midgut, such as peroxiredoxins [26,27]. These are a family of thioredoxin-dependent peroxidases, found in several insect species [49,50,51,52], that function as ROS-scavengers as well as other cellular processes. However their efficiency in converting *H₂O₂* was found to be significantly lower compared to catalase [53]. We are currently investigating the role of these antioxidant enzymes in ROS detoxification and fecundity and survival of female sand flies, as well as studies to confirm if proliferation of bacteria due to ROS reduction could account for the differences in sand fly survival.

Recent studies on transgenic *Anopheles stephensi* (the leading malaria vector in India and parts of Asia and the Middle East) overexpressing the protein kinase AKT gene increased the insulin signalling in the mosquito midgut, significantly reducing mosquito lifespan and inhibiting *P. falciparum* development [54]. The role of genes involved in stress responses in *Plasmodium* survival within the mosquito midgut was investigated by Jaramillo-Gutierrez et al. [55]. RNAi gene knockdown of the OXR1 gene (oxidation resistance gene) in *Anopheles gambiae* showed that this gene regulates the basal levels of catalase and glutathione peroxidase expression and that OXR1 gene knockdown decreased *Plasmodium berghei* oocyst formation. The findings of a *Lu. longipalpis* OXR1 gene homologue (unpublished) will shed some light into the regulation of ROS production within the sand fly gut and will also help to understand how ROS production impacts *Leishmania* development in the sand fly midgut.

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*Figure 7. Survival in female *Lu. Longipalpis* after RNAi-mediated depletion of catalase LlonKat1. Experimental group (dsCAT) exhibits a significantly lower survival rate compared to both dsGFP and pricked control groups, (P<0.0001, Kaplan-Meier, Log Rank χ² test). Results represent mean ± SEM of 3 independent biological replicates.*

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Current vector control strategies rely on spraying of residual insecticides to control vector population. Insect transgenesis and paratransgenesis are novel strategies that aim at reducing insect vectorial capacity by using genetic manipulation of disease vectors, rendering them incapable or less efficient to transmit a given pathogen [56] or even reducing the longevity and fecundity of a given insect vector. This study shows that catalase is a key gene in determining survival and fecundity of phlebotomine sand flies and future developments may warrant this gene being included as a potential target to reduce female sand fly fitness and reproductive capacity in the field.

Materials and Methods

Insects

All experiments were carried out using insectary-reared *Lu. longipalpis* from a colony first started with individuals caught in Jacobina, Brazil. Insects were kept under standard laboratory conditions [57]. Insects were fed with 70% w/v sucrose solution in cotton wool (unless stated differently in experiments), kept under a photoperiod of 8 hours light/16 hours darkness, temperature of 27°C (±2) and a relative humidity of >80% inside the rearing cages. Rabbit blood feeding was via a Hemotek membrane feeder (Discovery Workshops, UK) at 37°C. All procedures involving animals were performed in accordance with UK Government (Home Office) and EC regulations.

Fecundity assays

Female *Lu. longipalpis* were allowed to mate under regular rearing conditions and fed with rabbit blood at three, six and nine days post-emergence (DPE). A batch of >500 flies was released into a large (20 m²) rearing cage and groups of ~100 individuals were transferred to medium sized cages (5 m²) at 3, 6 and 9 DPE and blood-fed as above. Fifteen fully-engorged females were then transferred to a new medium rearing cage. Insects were dissected five days later to count developing oocytes.

ROS-scavenger feeding

Fecundity assays were carried out as described above with female *Lu. longipalpis* fed on a 70% sucrose solution supplemented with 20 mM ascorbic acid and blood-fed at 9 DPE. Supplemented sucrose-meal was freshly changed daily and continued after blood-feeding. A 9 DPE control group was reared under the same rearing conditions and fed with rabbit blood at three, six and nine days post-emergence (DPE). A batch of >500 flies was released into a large (20 m²) rearing cage and groups of ~100 individuals were transferred to medium sized cages (5 m²) at 3, 6 and 9 DPE and blood-fed as above. Fifteen fully-engorged females were then transferred to a new medium rearing cage. Insects were dissected five days later to count developing oocytes.

Ovarian Catalase Activity

Ovaries were collected from 5 female sand flies at 24 and 48 hrs post blood feeding (PBF). Samples were homogenised in 50 µl of 0.15 M NaCl solution, kept on ice and transferred to a −80°C freezer until needed. Before assays, samples were centrifuged at 5000 RPM for 2 minutes and 1 µl of the supernatant was diluted in 24.9 µl of 0.15 M NaCl solution. Catalase activity was determined using Amplex Red Catalase Assay Kit (Invitrogen Ltd) following the manufacturer’s protocol. Enzyme-specific activities were expressed as units/mg of protein. One unit of catalase activity was defined as 1 µM of H$_2$O$_2$ consumed per minute. All assays were carried out in triplicate. Fluorescence was measured using a Varioskan fluorescence spectrometer (Thermo Electron) with an excitation wavelength of 560 nm and an emission wavelength of 590 nm. Ovarian catalase activity was normalised using the total amount of protein in the whole body (minus dissected ovaries) using the BIORAD® Protein assay reagent following the manufacturer’s protocol and using bovine serum protein as standard. Endpoint absorbance was measured at 595 nm in a 96 well plate with a microplate reader (VersaMax Microplate Reader, Molecular Devices Inc.).

Table 1. Oligonucleotides for dsRNA synthesis and Reverse Transcriptase PCR.

| Oligonucleotide     | 5'-3' sequence                  | Size (bp) |
|---------------------|---------------------------------|-----------|
| dsCAT484 Forward    | TAATACGACTCACTATAGGGAAGTGCAGCTCTTTGCC | 524       |
| dsCAT484 reverse    | TAATACGACTCACTATAGGGAAGTGCAGCTCTTTGCC | 524       |
| dsGFP Forward       | TAATACGACTCACTATAGGGAAGTGCAGCTCTTTGCC | 693       |
| dsGFP Reverse       | TAATACGACTCACTATAGGGAAGTGCAGCTCTTTGCC | 693       |
| RT CAT484 Forward   | TGTTGCAGGAGCTCTTTGCC             | 484       |
| RT CAT484 Reverse   | AGGTTGACAGCTCTTTGCC              | 484       |
| RT Ribo605 Forward  | TCTCATGGAAGTTTCTTGCC            | 850       |
| RT Ribo60 Reverse   | GGCGTTGACACCTTTGACT             | 850       |

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Ovarian Catalase Expression

Six DPE sand flies were blood-fed and ovaries from 10 sand flies (two pools of 5 flies) were dissected at 12, 24 and 48 hours PBF, homogenised in 50 µl of TRI Reagent® (Ambion, Austin, TX) and kept at −80°C until needed. RNA was extracted following the manufacturer’s protocol. Total RNA was quantified using a NanoDrop® (NanoDrop Technologies, Wilmington, USA) and normalised to 10 ng/µl. RT-PCR was carried out with SuperScript® III One-Step RT-PCR System with Platinum® Taq DNA Polymerase Kit (Invitrogen, San Diego, CA) performing 19 cycles and following the manufacturer’s protocol (primers listed on table 1). Relative expression of catalase was normalised using a housekeeping gene (AM088777, 60S ribosomal protein L3). RT-PCR products were analysed by 1.5% agarose/ethidium bromide gel electrophoresis and reduction in catalase expression was determined by densitometric measurement of bands using the softwares GeneSnap/GeneTools (Sygen, UK).

Age-related expression of ovarian catalase

To measure catalase LlongKat1 mRNA expression levels in different age groups, 3, 6 and 9 DPE sand flies were blood fed and ovaries from 10 sand flies (two pools of 5 flies) were dissected at 48 hours PBF. Additionally, to evaluate the effect of feeding a ROS-scavenger in age-related expression of ovarian catalase, a
group of 9 days old sand flies was fed with ascorbic acid-supplemented sucrose solution as described above, blood fed and dissected at 48 hours. RNA was extracted and checked for catalase relative expression as above.

**RNAi-mediated catalase knockdown**

Sense and anti-sense catalase-specific primers flanked by the T7 promoter site (Table 1) PCR amplified a 404 bp product from a plasmid obtained from a whole body Lu. longipalpis normalised cDNA library [27] that was used as template for double-stranded RNA synthesis dsRNA. Transcription reactions and column purification were carried out using the Megascript RNAi Kit (Ambion®) following the manufacturer’s protocol. dsRNA purity was assessed by 1.5% agarose/ethidium bromide gel electrophoresis and dsRNA was quantitated using a Nanodrop ND-1000 Spectrophotometer (LabTech, UK). dsRNA was eluted with nuclease-free water at 65°C, concentrated to 4.5 μg/μL with a ChriSt® RVC 2–25 rotational vacuum concentrator and stored at −80°C until needed. Enhanced Green Fluorescent protein (eGFP) dsRNA was produced from a 653 bp amplicon of the pEGFP-N1 expression plasmid (Clontech) and used as a ‘mock’ injected control. RNAi was achieved by dsRNA injections as previously described [58]. After injections, sand flies were transferred to cages and kept with access to 70% sucrose solution ad libitum. Developing oocytes were dissected and counted 48 hours after blood feeding. Non-injected flies of the same age and kept under the same conditions were used as second control. Three pools of three whole sand flies were collected from each group to evaluate knockdown by RT-PCR.

**Survival assays**

To assess sand fly survival mediated by ROS-scavenging related to catalase activity, RNAi-mediated catalase knock down was carried out in a group of 50 sand flies. Flies were injected with dsRNA for catalase (dsCAT) as described above. To exclude wound-related mortality, all dead flies at 24 hrs post-injections were removed and were not included in the experiment. Dead sand flies were counted and removed from the cage daily from day 2 to 7 after injection. Flies injected with dsRNA for GFP (dsGFP) and a needle-pricked group were used as controls. To assess exogenous ROS-scavenging related survival, 50 female Lu. longipalpis were collected upon emergence and sugar fed on a 2% sucrose solution supplemented with 20 mM ascorbic acid. Dead sand flies were counted and removed from the cage every day until day seven. A group of sand flies fed with 70% sucrose was used as a control.

**Phenoloxidase assays**

Phenoloxidase activity was determined by measuring the production of dopachrome from 3,4 dihydroxy-DL-phenylalanine (DOPA) [59,60]. Briefly, single flies were homogenized in 60 μL of PBS and centrifuged at 25,000 g for 5 min at 4°C to recover the soluble fraction. 20 μL of supernatant was mixed with 10 μL of PBS (spontaneous PO) or trypsin solution (for total PO activity; 1 mg/mL in PBS, FLUKA cat. no. 93614), incubated for 20 min at 37°C followed by the addition of 20 μL of a saturated solution of DOPA (4 mg/mL in PBS) and absorbance (490 nm) measured by kinetic assay for 1 h at 5 minute intervals in a microplate reader at 30°C.

The PO activity was measured to ensure that activity was proportional to protein concentration and incubation time. Independent experiments showed that the PO activity was stable in the conditions above. Controls with no enzyme or no substrate were included. One unit of enzyme (U) is defined as the amount that produces 0.001 unit of absorbance/min.

**Sequence analysis**

The coding sequence of LlonKat1 was analyzed using the algorithms pl/Mw tool [61], signal IP [62], PRT1 Predictor [63], PeroxiP [64], TargetP [62] based at the EXPASY Proteomics Server (http://expasy.org/). Selected amino acid sequences of catalases were aligned with catalase LlonKat1 using the ClustalW Multiple Alignment tool in BioEdit Sequence Alignment Editor (http://www.mbio.ncsu.edu/BioEdit/BioEdit.html). Alignment was generated using Boxshade (http://www.ch.embnet.org/software/BOX_form.html).

**Statistical analysis**

Comparisons between means of two independent groups were carried out using a pair-wise t-test. Multiple comparisons were done by one-way ANOVA. Survival curves were analyzed with the Kaplan-Meier Log Rank test. Significance was considered when P<0.05. All data were analysed with the use of the SPSS Data Editor software (version 17.0, SPSS Inc).

**Supporting Information**

**Figure S1 Structure-based alignment of the aminoacid sequence of Lutzomyia longipalpis catalase, translated from a whole body (GeneDB NSFM-142e04.q1k [27]) and a midgut-specific (GenBank Accession number: EU124624.1 [26]) cDNA library.** Sequences show a 99% identity and a 99% similarity.

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**Author Contributions**

Conceived and designed the experiments: HDA MRVS FAG RJD. Performed the experiments: HDA RM FAG. Analyzed the data: HDA RM FAG. Contributed reagents/materials/analysis tools: RJD. Wrote the paper: HDA MRVS FAG RJD.

**References**

1. Lainson R, Rangel E (2005) Lutzomyia longipalpis and the eco-epidemiology of American visceral leishmaniasis, with particular reference to Brazil. A review. Memórias do Instituto Oswaldo Cruz 100: 811-827.
2. Soares RPP, Turco SJ (2003) Lutzomyia longipalpis [Diptera: Psychodidae: Phlebotominae]: a review. Anais da Academia Brasileira de Ciências 75: 301-330.
3. Grimaldi G, Tesh RB (1993) Leishmaniases of the New World: current concepts and implications for future research. Clinical Microbiology Reviews 6: 230-250.
4. Romero GAS, Boelaert M (2010) Control of Visceral Leishmaniasis in Latin America: A Systematic Review. PLoS Negl Trop Dis 4: e594.
5. Myskova J, Svobodova M, Beverley S, Volf P (2007) A lipophosphoglycan-independent development of Leishmania in permissive sand flies. Microbes and infection 9: 317-324.
6. Costa C (2008) Characterization and speculations on the urbanization of visceral leishmaniasis in Brazil. Cadernos de Saúde Pública 24: 2959-2963.
7. Djelong RJ, Müller LM, Molina-Cruz A, Gupta L, Kumar S, et al. (2007) Reactive oxygen species detoxification by catalase is a major determinant of fecundity in the mosquito Anopheles gambiae. Proceedings of the National Academy of Sciences 104: 2121.
14. Sohal R, Arnold L, Orr W (1990) Effect of age on superoxide dismutase.

10. Graca-Souza AV, Maya-Monteiro C, Paiva-Silva GO, Braz GR, Paes MC, et al. (2006) Adaptations against heme toxicity in blood-feeding arthropods. Insect Biochemistry and Molecular Biology 36: 392–395.

19. Wheeler D (1996) The role of nourishment in oogenesis. Annual Review of Entomology 41: 407–431.

52. Kim I, Lee K, Hwang J, Ahn M, Li J, et al. (2005) Molecular cloning and characterization of a peroxiredoxin gene from the mole cricket, Gryllotalpa ilu. longipalpis. Comparative Biochemistry and Physiology Part B: Biochemistry and Molecular Biology 140: 579–587.

7. Mandal G, Wyllie S, Singh N, Sundar S, Fairlamb AH, et al. (2007) Increased levels of thiols protect antimony unresponsive Leishmania donovani field isolates against oxidative stress. Antioxidants & Redox Signaling 9: 1527–1548.

5. Modr G Care and maintenance of phlebotomine sandfly colonies. Bulletin of the Entomological Research 41: 407–431.