Serological Markers of Sand Fly Exposure to Evaluate Insecticidal Nets against Visceral Leishmaniasis in India and Nepal: A Cluster-Randomized Trial

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Abstract

Background: Visceral leishmaniasis is the world’s second largest vector-borne parasitic killer and a neglected tropical disease, prevalent in poor communities. Long-lasting insecticidal nets (LNs) are a low cost proven vector intervention method for malaria control; however, their effectiveness against visceral leishmaniasis (VL) is unknown. This study quantified the effect of LNs on exposure to the sand fly vector of VL in India and Nepal during a two year community intervention trial.

Methods: As part of a paired-cluster randomized controlled clinical trial in VL-endemic regions of India and Nepal, we tested the effect of LNs on sand fly biting by measuring the antibody response of subjects to the saliva of Leishmania donovani vector Phlebotomus argentipes and the sympatric (non-vector) Phlebotomus papatasi. Fifteen to 20 individuals above 15 years of age from 26 VL endemic clusters were asked to provide a blood sample at baseline, 12 and 24 months post-intervention.

Results: A total of 305 individuals were included in the study, 68 participants provided two blood samples and 237 gave three samples. A random effect linear regression model showed that cluster-wide distribution of LNs reduced exposure to P. argentipes by 12% at 12 months (effect 0.88; 95% CI 0.83–0.94) and 9% at 24 months (effect 0.91; 95% CI 0.80–1.02) in the intervention group compared to control adjusting for baseline values and pair. Similar results were obtained for P. papatasi.

Conclusions: This trial provides evidence that LNs have a limited effect on sand fly exposure in VL endemic communities in India and Nepal and supports the use of sand fly saliva antibodies as a marker to evaluate vector control interventions.

Introduction

Visceral leishmaniasis (VL or kala azar) is a vector-borne parasitic disease with a fatal outcome if untreated. It is estimated that a large proportion of the annual 500,000 cases and 60,000 deaths occur in poor rural communities of the Indian subcontinent [1]. In these regions VL is exclusively caused by Leishmania donovani, transmitted by the bite of female Phlebotomus argentipes sand flies, an opportunistic blood feeding sand fly [2]. Phlebotomus papatasi, a man-biting sand fly sympatric with P. argentipes throughout the Indian subcontinent, does not transmit L. donovani, but is the Old World vector of zoonotic cutaneous leishmaniasis in much of Northern Africa and the Middle East [3]. Since there is no vaccine for VL, control measures depend on early case-detection, treatment and reduction in transmission through vector control measures. Current control of VL vectors in the Indian subcontinent is based on indoor residual spraying (IRS) of insecticides. Despite these efforts, the current strategy is failing to control VL in these regions [4]. Because L. donovani transmission is anthroponotic, and humans represent the only proven reservoir of infection, attention is being focused on the use of insecticide treated nets (ITNs), specifically, long-lasting insecticidal nets (LNs) to replace or compliment IRS.

Village-wide distribution of LNs have shown to significantly reduce indoor P. argentipes density by 25% [5], 44% [6] and 60% [7] in the Indian subcontinent. The variation observed in the effect of LN on P. argentipes density could be related to differences in experimental designs, vector behavior or insecticide susceptibility in Bangladesh, India and Nepal. Nevertheless the results of the first large-scale randomized controlled trial of the effectiveness LN to prevent VL in India and Nepal, indicate that LNs seem to have a small and not significant effect on the risk of L. donovani infection and clinical disease in VL endemic communities. During this trial...
Author Summary

Visceral leishmaniasis (VL), also known as kala azar, is one of the major public health concerns of the Indian subcontinent, caused by Leishmania donovani transmitted by the bite of the sand fly Phlebotomus argentipes. To date, Indoor Residual Spraying (IRS) campaigns have been unable to control the disease. This makes Long-lasting insecticidal nets (LNs) an attractive alternative or complement to IRS. Therefore, it is important to assess the extent that LNs reduce bites from P. argentipes. When female sand flies bite they require their saliva to efficiently bloodfeed. For humans and animals alike, the host’s immune response against components of sand fly saliva can be used as a marker of exposure to the vector. Here we describe how comprehensive coverage of LNs in trial communities over two years reduced antibody levels to the saliva of P. argentipes and P. papatasi (a man-biting sand fly that co-exists with P. argentipes but does not transmit VL) sand flies by 9–12% compared to communities without LNs. Our results demonstrate that the large-scale distribution of LNs did not confer significant additional protection against sand fly bites in VL-endemic regions of India and Nepal and questions the indoor transmission of L. donovani in these regions.

the risk for L. donovani infection, measured by means of Direct Agglutination Test (DAT), was reduced by 10% in clusters using LNs compared to controls [8]. Therefore, a tool to measure exposure to the VL vector will allow us to bridge a gap between the entomological and clinical results observed. The most direct way of doing this is by recording the numbers of bites individuals receive; however, since human landing catches are unethical for VL (VL is fatal with no effective prophylaxis) there are only a handful of studies reporting biting or landing rates of P. argentipes in VL foci [9–12]. An alternative method is required.

Sand flies rely on the vasodilatory and anti-haemostatic properties of their saliva to obtain blood for egg production and consequently salivate into the host’s skin with each bite. The relationship between the levels of antibodies to arthropod saliva, consequently salivate into the host skin with each bite. The properties of their saliva to obtain blood for egg production and measure human antibodies to successfully used to evaluate the efficacy of LNs against malaria transmission, epidemiology and risk of leishmaniasis and provides an opportunity to develop a versatile tool to understand the number of bites or amount of saliva injected [18]. This that the level of antibodies to salivary proteins are proportional to marker for vector infestation in domestic animals [15]. Sand fly saliva has been shown to be highly immunogenic for both humans and animals alike [16–17], and experimental studies have shown that the levels of antibodies to salivary proteins are proportional to the number of bites or amount of saliva injected [18]. This provides an opportunity to develop a versatile tool to understand the transmission, epidemiology and risk of leishmaniasis and evaluate vector intervention programs.

In Angola antibodies to the saliva of Anopheles gambiae was successfully used to evaluate the efficacy of LNs against malaria [19]. Recently, we developed a single saliva-based ELISA to measure human antibodies to P. argentipes and P. papatasi in VL-endemic areas [20]. An entomological survey of Indian and Nepalese households was used to assess the use of this ELISA as a tool to measure vector exposure. Indoor CDC light trap captures, used as proxy for sand fly exposure, were correlated to sand fly saliva antibodies in people. Similarly, in a small scale study of VL patients in Muzzafarpur, an endemic district of VL in India, we found that admission to hospital – thus protecting patients from sand fly bites for 30 days – resulted in a significant drop in antibodies to P. argentipes and P. papatasi saliva, which quickly rose again when treated patients returned to their villages and were re-exposed. To date, sand fly salivary antibodies have not been used to evaluate vector intervention programs at the community level. In the current study we screened sera from people given Deltamethrin-impregnated bednets, or not, to sleep under to assess their levels of anti-sand fly salivary antibodies over two years.

The objective of this study was to detect antibodies to P. argentipes and P. papatasi saliva to determine the effect of LNs on vector and non-vector sand fly exposure in VL-endemic villages of India and Nepal.

Materials and Methods

Study population

The blood samples included in this study are a subset of the samples collected in a large-scale, randomised controlled trial on the effectiveness of comprehensive LN distribution to prevent VL in the Indian subcontinent (KALANET, ClinicalTrials.gov CT-2005-015374). The study design is briefly described here. In May 2006, 26 VL endemic clusters with over 20,000 inhabitants were selected in India (n = 16) and Nepal (n = 10) based on their VL incidence from 2003 to 2005. The study clusters were matched by country, population size and pre-intervention VL incidence and randomly allocated to intervention or control groups, 13 clusters per arm. All households in the intervention group received Deltamethrin coated LNs (PermaNet 2.0) at baseline (November–December 2006). Enough LNs were distributed to ensure all households members slept under a net.

For the main trial outcome, we collected finger prick blood samples at baseline and 12 and 24 months post-intervention from all participants over 2 years of age. Incident L. donovani infections were determined by Direct Agglutination Test (DAT). Further details on the study design and on the effect of LNs on indoor sand fly density, L. donovani infection and VL are described elsewhere [5,8].

For this study, 15 to 20 individuals were selected in each study cluster in October 2006. The individuals were randomly selected among all the inhabitants in each cluster using the data collected in a demographic survey conducted in July 2006. Only individuals above 15 years of age were eligible these participants were asked to provide a larger amount of blood (3 ml) by vein puncture at baseline, 12 and 24 months post-intervention. The sera obtained by centrifugation were identified with the individual ID and kept at −20°C until the laboratory analyses were conducted. Information on the age, gender, VL history, DAT titre at baseline, malnutrition and Socio-Economic Status (SES) were available for all participants. The methods used to evaluate the malnutrition and SES are described in detail elsewhere [21].

Saliva preparation

Salivary gland lysate (SGL) of colonised P. argentipes and P. papatasi sand flies (Charles University, Prague, Czech Republic) was obtained as described previously [16–17,20]. Salivary glands were dissected from female flies maintained on sucrose solution ad libitum at five days old post-emergence. SGL was lyophilized and reconstituted in its original volume of distilled water for 1 hr at room temperature (25°C) before use.

Laboratory analyses

Pre-adsorption of sera against P. papatasi saliva significantly improves the specificity of the P. argentipes ELISA by reducing the levels of cross-reaction [20]. This is achieved by reducing the amount
of antibodies which commonly recognise salivary antigens of both these sand flies. To do this, 30 ng \( P. \ papatasi \) SGL in bicarbonate buffer (pH 9.6) was coated in each well of microtiter plates (maxisorp, Nunc) at 4°C overnight. After washing 4 times (PBS-0.05% Tween 20 (PBS-T) Fluka, Sigma), plates were blocked with 5% bovine serum albumin in PBS-T for 2 hr at 37°C. After washing, 1:50 diluted human sera in PBS-T were added and incubated overnight at 4°C (the \( P. \ papatasi \) pre-adsorption step). Simultaneously another plate was coated with SGL of \( P. \ argentipes \) (50 ng/well) at 4°C overnight. The next day after washing and blocking of the \( P. \ argentipes \) plate, sera were transferred from the \( P. \ papatasi \) plate and incubated at 37°C for 2 hr. From this point both plates were processed in parallel. Plates were incubated with biotinylated goat anti-human IgG (1:1000 in PBS-T, Sigma) for 1 hr at 25°C, washed and incubated with streptavidin-conjugated alkaline phosphatase (1:100 dilution in PBS-T, Sigma) overnight at 25°C. To develop the reaction substrate (paranitrophenylphosphate, 1 mg/ml, Sigma) was added and the optical density (OD) measured at 405 nm using a Spectramax 190 ELISA plate reader after 20 minutes incubation in the dark. To minimise day to day variation in ELISA performance three sera from the same individual collected over the entire trial (baseline; 12 and 24 months follow-up samples) were processed in the same plate.

Cut offs for positive \( P. \ argentipes \) and \( P. \ papatasi \) ELISA were determined as the average OD values plus two standard deviations of 9 Indian non-endemic controls (NEC) from urban, non-VL areas of Western Uttar Pradesh [20].

Statistical analyses

Individual and geometric mean ELISA OD per immunological survey: baseline (Nov-Dec 2006), 12 and 24 months follow-up; and intervention group (LN and control clusters) were plotted and tabulated. A random effect linear regression model was used to estimate the effect of LN on the log transformed ELISA OD at 12 and 24 months. The following model was applied:

\[
Y_{3ijk} = \gamma_j + \beta_i Y_{1ijk} + U_j + \epsilon_{ijk}
\]

Where the outcome \( Y_{3ijk} \) is the log-transformed OD at 24 months for person \( k \) in cluster \( j \) and treatment arm \( i \). And \( \gamma_j \) is a fixed pair effect to take the matching into account, \( \beta_i \) is the intervention effect, \( \gamma \) is the effect of the log-transformed baseline value \( Y_{1ijk} \), \( U_j \) is a random cluster effect assumed normally distributed with mean 0 and variance \( \sigma_u^2 \) (the between cluster variation within matched pairs) and \( \epsilon_{ijk} \) the individual measurement error also assumed normal with mean 0 and variance \( \sigma_e^2 \) (variation between individuals within same cluster). The main parameter of interest is \( \beta_i \) which measures the mean difference in log OD at 24 months for two persons from the same pair, with the same baseline OD, one from the intervention cluster and the other from the control cluster. The fit of the model was checked by residual plots. An analogous model was used to study the log-transformed ELISA OD result at 12 months.

In separate analyses those individuals with anti-\( P. \ argentipes \) or anti-\( P. \ papatasi \) OD values below the cut offs or no records at baseline were removed to increase the sensitivity of the data [22]. The data were analysed in Stata 11 (StataCorp LP, College Station, TX, USA).

Ethical issues

Written informed consent was obtained from each participant or their guardian for those under 18 years old. Ethical approval was obtained from the Institutional Review Boards (IRB) of the B. P. Koirala Institute of Health Sciences, Nepal; the Institute of Medical Sciences Banaras Hindu University, India and the Institute of Tropical Medicine, Antwerp, Belgium.
Discussion

The results of this study show that *P. argentipes* exposure was reduced by 9 to 12% in people living in villages where LNs were used compared to controls. This reduction is in the same order of magnitude of the effect of LN on *L. donovani* infection observed in the same study clusters in India and Nepal [8]. Even if the use of LN reduced the *P. argentipes* indoor density in the study clusters [5] and seemed to provide some degree of personal protection [8,23], a significant number of subjects living in intervention clusters had high levels of antibodies against *P. argentipes* after 24 months of LN use (43.5% were ELISA positive). These results could be explained if LN failed to reduce the sand fly abundance as shown in a previous study in the area [24] or by the incorrect use of LN. However, as over 90% of the participants in the intervention clusters use the LN regularly (i.e. over 80% of the nights), they seem to support the theory that a substantial fraction of *L. donovani* transmission occurs outside the house where LNs would not prevent sand fly-human contact [8]. This goes against the traditional narrative that *P. argentipes* predominantly bite at night, and inside houses [9–12]. This hypothesis cannot be proved with this study design but it is supported by the trial results as a whole: i.e. similar *L. donovani* infection (5.4% vs. 5.5%) and VL (0.38% vs. 0.40%) rates were reported in both intervention and control clusters [8]. Moreover, *P. argentipes* are known to breed outside households [25], significant numbers of *P. argentipes* captured around households [26–27] and about 15–20% of those collected in cattle sheds had fed on humans [28–29]. The latter results could be related to the movement of blood fed females but they also suggest that *P. argentipes* are somehow exophagic.

The effect of LNs on *P. papatasi* exposure was similar with a 9–11% reduction in exposure. *Phlebotomus papatasi* is a sand fly sympatric with *P. argentipes* throughout the Indian subcontinent, is also endophilic and highly anthropophilic [30–31]. Although it is man-biting, *Phlebotomus papatasi* does not vector *L. donovani*. However, it would appear that LNs protect against the bite of *P. argentipes* and *P. papatasi* equally.

### Table 1. Characteristics of individuals excluded and lost to follow-up.

<table>
<thead>
<tr>
<th></th>
<th>Lost to follow-up</th>
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<tbody>
<tr>
<td></td>
<td>Control</td>
</tr>
<tr>
<td>Total Individuals</td>
<td>62</td>
</tr>
<tr>
<td>Mean age (SD)</td>
<td>27.5 (7.7)</td>
</tr>
<tr>
<td>No. males (%)</td>
<td>39 (63%)</td>
</tr>
<tr>
<td>Percentage of DAT positive at baseline</td>
<td>17% (10/57)</td>
</tr>
<tr>
<td>No. Individuals with past history of VL (%)</td>
<td>6 (9.7%)</td>
</tr>
<tr>
<td>Mean SES indicator² (SD)</td>
<td>2.1 (1.6)</td>
</tr>
<tr>
<td>Percentage of individuals with Moderate or Severe Malnutrition³ (n/N)</td>
<td>5.2% (3/57)</td>
</tr>
<tr>
<td>No. Individuals living in houses with at least one VL case in past 24 months (%)</td>
<td>6 (9.7%)</td>
</tr>
</tbody>
</table>

¹Direct Agglutination Test (DAT) titre ≥1:1600. ²Socio-Economic Status indicator calculated as detailed in Singh et al [21]. ³Nutrition status calculated as detailed in Singh et al [21].

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By removing the non-responders we improve the specificity of the saliva ELISA. The adjusted data revealed that the difference in geometric mean of ELISA ODs for *P. argenteipes* shows a greater drop that that observed for *P. papatasi*. This may be due to differences in bloodfeeding or resting behaviour, as hypothesised above. Currently, *P. argenteipes* is considered more endophilic than endoendemic, often found digesting their bovine bloodmeals within households which have live stock nearby, or are commonly housed in the same building [25–27–28]. In contrast, *P. papatasi* is considered less opportunistic and more endophilic [30–31]. If this is the case one would expect a larger drop in ELISA ODs against *P. papatasi* as they would come into contact with the LN, attracted to the sleeping occupant. Recently, Dinesh and colleagues showed that *P. argenteipes* from the same areas of India and Nepal as our study were very sensitive to deltamethrin but *P. papatasi* was not compared [32]. Therefore, there remains the possibility of different susceptibilities to this insecticide between different sand fly populations. An alternative hypothesis is that the LNs repelled *P. papatasi,* although no such properties have been reported in the literature.

The results of this study support the use of the sand fly saliva ELISA as a sensitive tool to evaluate vector control intervention. Similar methods have been used to assess the exposure to *Anopheles gambiae* in natural conditions in Senegal [33] and to evaluate the efficacy of ITNs in malaria vector control in Angola [19]. The latter study reported a significant decrease in the antibody response to *An. gambiae* after the introduction of ITNs. However, in contrast to our study, the magnitude of the effect was not assessed, a "before and after intervention" design was used [so there were no concurrent controls] and only 109 samples were analysed [19].

The baseline sand fly saliva antibody values were different between intervention and control groups; people in intervention clusters seemed to have a higher sand fly exposure before the LNs were distributed (Figure 1). This contrasts with the baseline data from the trial which showed that intervention and control clusters had similar indoor *P. argenteipes* density [3] and similar population characteristics [8]. This difference may be due to random error as the number of samples per cluster was small (6 to 17 subjects/cluster) and there were some differences between groups at baseline; i.e. more past VL cases.

### Table 2. Study population characteristics.

<table>
<thead>
<tr>
<th></th>
<th>All available samples</th>
<th>Restricted <em>P. argenteipes</em></th>
<th>Restricted <em>P. papatasi</em></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control</td>
<td>Intervention</td>
<td>Control</td>
</tr>
<tr>
<td>Total Individuals (range per cluster)</td>
<td>155 (7–17)</td>
<td>150 (6–17)</td>
<td>72 (2–14)</td>
</tr>
<tr>
<td>Mean age (SD)</td>
<td>28.2 (7.6)</td>
<td>28.3 (7.5)</td>
<td>27.4 (8.5)</td>
</tr>
<tr>
<td>No. males (%)</td>
<td>48 (31.0)</td>
<td>63 (420)</td>
<td>25 (34.7)</td>
</tr>
<tr>
<td>No. DAT positive at baseline (%)</td>
<td>24 (15.5)</td>
<td>27 (18.0)</td>
<td>14 (19.4)</td>
</tr>
<tr>
<td>No. individuals with past history of VL (%)</td>
<td>8 (5.2)</td>
<td>13 (8.7)</td>
<td>3 (4.2)</td>
</tr>
<tr>
<td>Mean SES indicator (SD)</td>
<td>2.2 (1.4)</td>
<td>1.8 (1.4)</td>
<td>2.3 (1.4)</td>
</tr>
<tr>
<td>No. individuals with Moderate or Severe Malnutrition (%)</td>
<td>12 (7.7)</td>
<td>18 (12.0)</td>
<td>1 (1.4)</td>
</tr>
<tr>
<td>No. Individuals living in houses with at least one VL case in past 24 months (%)</td>
<td>11 (7.1)</td>
<td>14 (9.3)</td>
<td>6 (8.3)</td>
</tr>
</tbody>
</table>

**Notes:**
1. Excluding records with no ELISA results or OD for *P. argenteipes* below 0.9 at baseline (n = 163).
2. Excluding records with no ELISA results or OD for *P. papatasi* below 1.8 at baseline (n = 73).
3. Direct Agglutination Test (DAT) titre ≥:1:1600.
4. Socio-Economic Status indicator calculated as detailed in Singh et al [21].
5. Nutrition status calculated as detailed in Singh et al [21].
6. Geometric mean (GM) and inter quartile range (IQR) of ELISA Optical Density (OD) per immunological survey (baseline, 12 and 24 months follow-up) and intervention group (LN and control clusters) for *Phlebotomus argenteipes* and *P. papatasi*. Number of samples positive per survey using 0.9 and 1.8 ELISA OD as cut off values for *P. argenteipes* and *P. papatasi* respectively. Estimates of the intervention effect at 12 and 24 months adjusting for pair and baseline ELISA OD value. Results were obtained using all samples available (n = 305).
in the intervention group (Table 1). Differential dropout between the study groups may have also caused the differences observed at baseline. However, even if there were more individuals lost to follow-up in control group than in the intervention group, the individuals excluded from both groups had similar characteristics (Table 1). To take into account the differences at baseline, the statistical model used to evaluate the impact of LN on sand fly exposure was adjusted for baseline values. Similarly, when the analyses were restricted to positive ELISA results at baseline to increase the sensitivity of the test [22], the baseline values were equilibrated between groups and the effect of LN on *P. argentipes* exposure remained unaltered (14% reduction).

Antibody-based assays to measure vector exposure represents an advance from traditional methods of vector sampling since light traps are not effective in catching bloodfed *P. argentipes* [34] and are unable to measure the human-sand fly contact outside households. It is important that such assays are specific to the vector, sensitive to the number of bites received and responsive to changes in exposure over time [35,20]. We have previously shown that our saliva-ELISA correlates with indoor *P. argentipes* densities, and pre-absorption of sera against *P. papatasi* saliva reduced cross-reaction with this non-VL vector, which may lead to false positive results [20]. In the future, recombinant peptides screened from cDNA libraries constructed from *P. argentipes* (and *P. papatasi*) salivary glands will insure against this problem. Despite the drawbacks of using whole saliva as ELISA antigen (labour intensive, costly and time consuming) experimental studies have shown that not all saliva-positive human sera recognize the same protein bands [36,17,22]. In this respect, whole saliva has an advantage as it represents all peptides.

In conclusion we demonstrate that the current *P. argentipes* saliva antibody test is a useful tool for the evaluation of vector intervention programmes in human populations from VL-endemic areas. It would appear that LNs have a limited effect on sand fly exposure.

### Table 4. Average anti-saliva antibody response – baseline adjusted.

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>Intervention</th>
<th>Intervention effect</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. Samples</td>
<td>GM ELISA OD (IQR)</td>
<td>No. Positive (%)</td>
</tr>
<tr>
<td><strong>P. argentipes</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>72</td>
<td>1.52 (1.15; 2.14)</td>
<td>72 (100)</td>
</tr>
<tr>
<td>12 months</td>
<td>59</td>
<td>1.35 (0.97; 1.82)</td>
<td>48 (81.4)</td>
</tr>
<tr>
<td>24 months</td>
<td>60</td>
<td>1.33 (0.99; 1.75)</td>
<td>49 (81.7)</td>
</tr>
<tr>
<td><strong>P. papatasi</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>26</td>
<td>2.30 (2.01; 2.50)</td>
<td>26 (100)</td>
</tr>
<tr>
<td>12 months</td>
<td>22</td>
<td>2.22 (1.92; 2.75)</td>
<td>18 (81.8)</td>
</tr>
<tr>
<td>24 months</td>
<td>21</td>
<td>2.23 (1.95; 2.62)</td>
<td>17 (81.0)</td>
</tr>
</tbody>
</table>

Geometric mean (GM) and inter quartile range (IQR) of ELISA Optical Density (OD) per immunological survey (baseline, 12 and 24 months follow-up) and intervention group (LN and control clusters) for *Phlebotomus argentipes* and *P. papatasi*. Number of samples per survey using 0.9 and 1.8 ELISA OD as cut off values for *P. argentipes* and *P. papatasi* respectively. Estimates of the intervention effect at 12 and 24 months adjusting for pair and baseline ELISA OD value. Results were obtained excluding records with no ELISA results or OD below the cut off values at baseline.

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exposure and combined interventions that address the peri-and intradomestic environment seem the way forward. VL control will require strengthening vector control methods. Rapid case detection and treatment alone may be insufficient to control L. donovani transmission if asymptomatic infected individuals play a role in VL epidemiology as a recent mathematical model suggests [37]. More research on the behaviour of P. argenteipes in relation to L. donovani transmission would be prudent to refine future intervention strategies for VL.

References


Figure 3. Effect of LNs on sand fly exposure, adjusted for non-endemic controls. Individual ELISA Optical Density (OD) per immunological survey (baseline, 12 and 24 months follow-up) for intervention (long-lasting insecticidal net, LN – black triangles) and control clusters (grey circles), for Phlebotomus argentipes (Panel A) and P. papatasi (Panel B). Individuals with no ELISA results or below the average non-endemic control OD+2×S.D. cut-off values (0.9 for P. argenteipes and 1.8 for P. papatasi) at baseline were excluded. The geometric means ELISA OD are represented as a solid line + S.D. and 1.8 for P. papatasi) at baseline were excluded. The geometric means ELISA OD are represented as a solid line + S.D. and dotted line for control groups. The Mann Whitney t-test was used to compare 12 and 24 month follow-up samples compared to their corresponding baseline values, asterisks denote statistical significance (*, P<0.05; **, P<0.005; *, P<0.005; ns, not significant P>0.05).

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Supporting Information

Checklist S1 CONSORT checklist. (DOCX)

Author Contributions

Conceived and designed the experiments: MR AP MB SS SR. Performed the experiments: KG AP SR BO MS JC PV MR. Analyzed the data: EWA AP MR. Contributed reagents/materials/analysis tools: VV PV. Wrote the paper: MR AP EWA MB.


