Short Report: Lack of Protection of Pre-Immunization with Saliva of Long-Term Colonized *Phlebotomus papatasi* against Experimental Challenge with *Leishmania major* and Saliva of Wild-Caught *P. papatasi*

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**Abstract.** Immunity to saliva of *Phlebotomus papatasi* protects against *Leishmania major* infection as determined by co-inoculation of parasites with salivary gland homogenates (SGHs) of this vector. These results were obtained with long-term colonized female *P. papatasi*. We investigated the effect of pre-immunization with SGH of long-term colonized *P. papatasi* against *L. major* infection co-inoculated with SGH of wild-caught *P. papatasi*. Our results showed that pre-exposure to SGH of long-term, colonized *P. papatasi* do not confer protection against infection with *L. major* co-inoculated with wild-caught *P. papatasi*. These preliminary results strongly suggest that the effectiveness of a vector saliva-based vaccine derived from colonized sand fly populations may be affected by inconsistent immune response after natural exposure.

Several studies have shown that pre-immunization of mice with needle-injected saliva or pre-exposure to uninfected bites of *Phlebotomus papatasi* provided protection against infection with *Leishmania major*, the etiologic agent of zoonotic cutaneous leishmaniasis (ZCL).1–3 These studies were performed using long-term laboratory colonized sand flies mainly because of the difficulty of working with wild-caught flies. A puzzling fact is that people in leishmaniasis-endemic areas succumb of ZCL despite the high frequency of uninfected bites compared with infected ones.4 We showed in a previous study that pre-immunization of mice with salivary gland homogenate (SGH) of wild-caught *P. papatasi* does not confer protection against *L. major* co-inoculated with the same type of SGH compared with a significant protection obtained when pre-immunization and challenge were performed with SGH of long-term colonized flies.5 It was reported that colonized and wild-caught *Lutzomyia longipalpis* differ in the composition and the amount of salivary proteins and these differences may account for the lower effect observed on the modulation of experimental *Leishmania* infection by wild-caught SGH.6,7 Thus, these studies provide good evidence that the outcome of *Leishmania* infection differs significantly between colonized and wild-caught salivary gland proteins. Our aim in this study was to assess the protective effect of pre-immunization with SGH of long-term colonized *P. papatasi* on experimental *L. major* challenge co-inoculated with SGH of wild-caught *P. papatasi*.

*Phlebotomus papatasi* (Tunisian strain that originated in the Governorate of Sidi Bouzid) have been reared at the *Vector Ecology Laboratory of the Institut Pasteur* de Tunis since 2003.3 This colony is maintained without being supplemented periodically with wild-caught *P. papatasi*. Generation F39 was used in this study. Wild sand flies were collected using CDC light traps from animal shelters located in the village of Felta (Governorate of Sidi Bouzid), a focus highly endemic for ZCL.9 Preparation of SGH from laboratory-reared and wild-caught *P. papatasi* was performed as described.3 A highly virulent strain of *L. major*, MHOM/TN/95/GLC94, isolated from a patient in Tunisia was used in this study.5 We used female BALB/c mice (6–8 weeks of age) bred in the animal facility of the Institut Pasteur de Tunis.

Mice were immunized intradermally in the right ear with the equivalent of two pairs of salivary glands in 10 µL of phosphate-buffered saline (PBS). Two groups of 10 mice each were pre-immunized with SGH of long-term colonized female *P. papatasi* (F39), one a week for two weeks. In the fourth week, the first [CSGH (F39)-L.m+CSGH (F39)] and the second [CSGH (F39)-L.m+WSGH] groups were challenged with 106 stationary phase *L. major* promastigotes in 50 µL of PBS co-inoculated subcutaneously in the right hind footpads with SGH of long-term, colonized and SGH of wild-caught, female *P. papatasi*, respectively. Three control groups of 10 mice each were used in this study. The third [PBS-L.m+CSGH (F39)] and the fourth group [PBS-L.m+WSGH] (control groups) were injected with PBS instead of SGH and challenged with 106 stationary phase *L. major* promastigotes in 50 µL of PBS co-inoculated subcutaneously in the right hind footpads with SGH of long-term colonized, and SGH of wild-caught female *P. papatasi*, respectively. The fifth group [PBS-L.m+PBS] (control group) was injected with PBS instead of SGH and challenged with only 105 stationary phase *L. major* promastigotes in 50 µL of PBS inoculated subcutaneously in the right hind footpads. All experiments were replicated three times. The footpad swelling at the site of inoculation was monitored at weekly intervals by using a vernier caliper. Lesion size was defined as the increase in the footpad thickness after subtracting the size of the contralateral uninfected footpad.

Using a linear mixed-effects model for longitudinal data10 but allowing for nested random effects, and where the within-group errors are permitted to be correlated and/or have unequal variances, we tested for difference in trends (group effect) and time-group interaction between curves illustrating the variation of the lesion size through time for each group of mice immunized and challenged differently as described above. In addition, for specific time point analysis (post-challenge week) the Wilcoxon test11 and Student t-test were used to test for median and mean difference of lesion size between groups. Holm’s correction for multiple testing12 of the reported *P* values was used when appropriate. All statistical analyses
were performed with R software for statistical computing (version 2.7).

After challenge with \( L. \text{major} \) co-inoculated with SGH of long-term, colonized, female \( P. \text{papatasi} (F_{39}) \), mice pre-immunized with the same type of SGH developed footpad lesions that were significantly smaller in size and grew more slowly than in the control groups \( (P < 0.0001) \) (Figure 1). In contrast, mice pre-immunized with SGH of long-term colonized \( P. \text{papatasi} \) and challenged with \( L. \text{major} \) co-inoculated with SGH of wild-caught \( P. \text{papatasi} \) developed lesions that grew as rapidly and as large in size as the control groups \( (P = 0.7389) \) (Figure 1). Lesion size differed significantly in mice pre-immunized with SGH of long-term colonized \( P. \text{papatasi} \) and then challenged with \( L. \text{major} \) co-inoculated with the same type of SGH compared with lesion size observed in the group of mice pre-immunized with SGH of long-term colonized \( P. \text{papatasi} \) and then challenged with \( L. \text{major} \) co-inoculated with SGH of wild-caught female \( P. \text{papatasi} \) \( (P < 0.0001) \). Overall, no significant difference was observed in lesion size among three control groups \( (P = 0.18) \) (Figure 1).

Our results confirmed the finding of previous studies showing that pre-immunization with SGH of long-term laboratory-colonized female \( P. \text{papatasi} \) induced significant protection against \( L. \text{major} \) co-inoculated with the same type of SGH.\(^{1,3}\) However, all aforementioned studies were performed with long-term laboratory colonized sand flies. Because persons in leishmaniasis-endemic areas are exposed to bites of wild populations of \( P. \text{papatasi} \), we investigated the effects of sand fly saliva as close as possible to natural transmission. Our results showed that pre-immunization of mice with SGH of long-term, laboratory-colonized, female \( P. \text{papatasi} \) did not protect mice against \( L. \text{major} \) co-inoculated with SGH of wild-caught female \( P. \text{papatasi} \).

Previous studies using small numbers of parasites (500–1,000 promastigotes) reported that lesion size was exacerbated in the presence of SGH compared with parasites alone.\(^{1,3}\) In our study, SGH of either colonized or wild-caught \( P. \text{papatasi} \) did not exacerbate \( L. \text{major} \) infection compared with lesion sizes observed when parasites were injected alone. This finding is probably caused by the high number of parasites (10\(^6\) promastigotes) used in our study.

Among nine identified salivary gland proteins of long-term, colonized, female \( P. \text{papatasi} \), one with an apparent molecular weight of 15 kD \( (\text{SP-15}) \) provided significant protection of mice when challenged with \( L. \text{major} \).\(^{13}\) Wild-caught \( P. \text{papatasi} \) exhibited higher genetic variation in \( \text{SP-15} \) than compared with colonized flies of the same species.\(^{13}\) Many variants of SP15 were found in natural field populations of \( P. \text{papatasi} \).\(^{13}\) It was hypothesized that the development of a vaccine based on SP-15 will not be affected by an inconsistent immune response because of genetic variation in natural populations of \( P. \text{papatasi} \).\(^{13}\) However, the protective ability of one variant of SP15 against exposure to another variant needs to be studied.

The fact that pre-immunization of mice with SGH of long-term, laboratory-colonized, female \( P. \text{papatasi} \) did not protect mice against \( L. \text{major} \) co-inoculated with SGH of wild-caught female \( P. \text{papatasi} \) strongly suggests that the effectiveness of a sand fly saliva–based vaccine will be affected by the antigenic diversity of sand fly salivary proteins that results from the genetic variation in natural populations of \( P. \text{papatasi} \). To clarify this hypothesis, studies concerning salivary proteins of wild populations of \( P. \text{papatasi} \) are needed. Similarly, among natural field populations of \( L. \text{longipalpis} \), extensive amino acid sequence variation (up to 23\%) was observed in maxadilan peptides.\(^{14}\) Natural selection may favor the polymorphism observed in maxadilan peptides to escape host immune evasion.

**Figure 1.** Course of lesion development in vaccinated and control BALB/c mice after challenge with \( 10^6 \) Leishmania major metacyclic promastigotes. Results are expressed as increases in infected footpad thickness (in millimeters) and are means + SD per group. Data are representative of three experiments combined. CSGH \( (F_{39}) \)-L.m+CSGH \( (F_{39}) \) = mice pre-immunized with salivary gland homogenate \( (\text{SGH}) \) of long-term laboratory-colonized female \( P. \text{papatasi} \) \( (F_{39}) \) and challenged with \( L. \text{major} \) co-inoculated with the same type of SGH; CSGH \( (F_{39}) \)-L.m+WSGH = mice pre-immunized with SGH of long-term colonized female \( P. \text{papatasi} \) \( (F_{39}) \) and challenged with \( L. \text{major} \) co-inoculated with SGH of wild-caught female \( P. \text{papatasi} \); PBS-L.m+WSGH \( (\text{control group}) \) = mice pre-immunized with PBS only and challenged with \( L. \text{major} \) co-inoculated with PBS. Symbols represent the mean + SD curve for each group. Asterisks indicate a significant statistical difference in lesion size (by \( t \)-test or Wilcoxon test, adjusted \( P < 0.05 \)) for specific time point \( (\text{week}) \) between the CSGH-L.m+CSGH \( (F_{39}) \) group and the remaining groups (three control groups and the CSGH-L.m+WSGH group).
responses. This hypothesis is corroborated by the fact that although maxadilan exacerbates infection with *L. major*, vaccination against one variant of maxadilan protected mice against *L. major* infection. Therefore, the protective effect observed with one variant of maxadilan may differ when exposed to another variant. Although SP-15 is protective against *L. major*, immunization with another salivary gland protein (SP-44) from the same colony of *P. papatasi* induced disease exacerbation. Thus, proper selection of a vector-based vaccine candidate is of major importance.

Laboratory colonies of insects are often accepted as being representative of field populations from which they have been derived, but this assumption may be challenged because colonies frequently incorporate only a fraction of the genetic variability present in the original populations. It has been reported that laboratory colonization of sand flies reduces natural genetic variability, and might foster selection for certain traits that are normally suppressed in field populations. The loss of genetic variation as a result of colonization might figure prominently in the protection observed in mice pre-immunized with SGH of long-term colonized *P. papatasi* and challenged with *L. major* co-inoculated with the same type of SGH. Antigenic variation of salivary gland proteins of field populations of *P. papatasi* similar to that observed in *Lu. longipalpis* might explain the absence of protection observed in mice pre-immunized with SGH of long-term colonized *P. papatasi* and challenged with *L. major* co-inoculated with SGH of wild-caught *P. papatasi*. Thus, our preliminary results strongly suggest that the development of a vaccine based on salivary gland proteins needs to include consideration of variability in natural populations of *P. papatasi*.

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