HUMAN IMMUNE RESPONSE TO SAND FLY SALIVARY GLAND ANTIGENS: A USEFUL EPIDEMIOLOGICAL MARKER?

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Abstract. Antibody (IgG) responses to salivary gland homogenate and to a recombinant salivary protein from the sand fly Lutzomyia longipalpis were investigated using sera from children living in an endemic area of visceral leishmaniasis in Brazil. We classified children into four groups according to their responses to Leishmania antigen: (Group I) positive serology and positive delayed type hypersensitivity (DTH), (Group II) positive serology and negative DTH, (Group III) negative serology and positive DTH, and (Group IV) negative serology and negative DTH. A highly significant correlation was found between anti-salivary gland IgG levels and DTH responses. An L. longipalpis salivary recombinant protein used as an antigen in an enzyme-linked immunosorbent assay (ELISA) gave a significant but different result. A positive correlation was found between anti-Leishmania IgG and anti-recombinant protein IgG titers. The results indicate that sand fly salivary proteins may be of relevance to the study of the epidemiology of leishmaniasis.

INTRODUCTION

The leishmaniasis are a group of mostly zoonotic diseases caused by protozoan parasites of the genus Leishmania which are transmitted by phlebotomine sand fly vectors. Sand flies acquire the parasite, which lives as intracellular amastigotes in their vertebrate hosts, while taking a blood meal. The parasites develop from amastigotes to extracellular promastigotes in the sand fly gut, multiply, and transform to infective metacyclics that accumulate in the anterior portion of the fly’s digestive canal. Once infective protozoa invade their vector’s mouthparts, they can be transmitted to another vertebrate when the fly attempts to take another blood meal.1

Visceral American leishmaniasis (VL) caused by Leishmania chagasi is endemic in several areas in Brazil. Within these regions, most of the infections are subclinical with leishmanin-positive patients normal upon physical examination.2 Full-blown VL is characterized by unimpeded parasite multiplication in bone marrow, liver, and spleen, leading to severe weight loss, hepatosplenomegaly, anemia, leukopenia, and thrombocytopenia. Development of clinical disease is possibly related to nutritional status3 and young age4 among other factors. A strong delayed type hypersensitivity (DTH) to leishmanial antigen (leishmanin) is normally associated with resistance to manifestations of the disease.5,6 Additionally, laboratory experiments indicate that immunity to sand fly vector salivary antigens may also confer protection.5,6

While taking a blood meal, sand flies salivate on their hosts. The saliva of blood-sucking animals contains a repertoire of molecules that modulate their host’s hemostatic, inflammatory, and immune responses. The New World sand fly Lutzomyia longipalpis has the peptide maxadilan as its salivary vasodilator.8,9 Maxadilan also has immunosuppressive properties10,11 and the anti-clotting molecule of Lutzomyia has been purified and cloned.12 Additionally, several salivary gland cDNAs from Lutzomyia longipalpis have been recently cloned, including the salivary apprase and hyaluronidase.13 Salivary components may also play a role in helping the establishment of pathogens in their vertebrate hosts.6,8 Tick saliva has been implicated in the successful transmission of several tick-borne viruses14,15 and saliva of both Lutzomyia longipalpis and Phlebotomus papatasii has been implicated in enhancing transmission of Leishmania major parasites in mouse models of this disease.13,16 Conversely, host immunity to vector saliva may decrease infectivity of the transmitted pathogens.5,6

A large body of work demonstrates the production by humans and other vertebrates of antibodies against the salivary gland components of blood-sucking insects and ticks.17–19 Although most of this work is aimed at preventing and treating undesirable allergic responses, or developing anti-arthropod vaccines, these responses may also be used as epidemiological markers of vector exposure, such as to ixodes scapularis, a vector of Lyme disease.20,21 In the case of leishmaniasis, there is a dual interest in understanding the nature of the human response to sand fly salivary antigens.1) Sand fly populations tend to be clustered (Miranda and others, unpublished data). Screening of human antibodies to sand fly saliva could be a useful indicator the spatial distribution of sand flies in a particular region, thus helping to direct vector and disease control efforts. 2) Immunity to sand fly salivary antigens may decrease expression of leishmaniasis or disease severity.6,9 Accordingly, the purpose of this work was to investigate whether sera from children living in an area endemic for visceral leishmaniasis were able to differentially recognize salivary gland antigens of the vector sand fly depending on the children’s exposure to Leishmania as determined by their cellular (measured by cutaneous DTH) and humoral IgG responses to Leishmania antigens. The results indicate that anti-sand fly antibody studies can be used within the context of the epidemiology of leishmaniasis.

MATERIALS AND METHODS

Sand flies. Sand flies were reared at the Walter Reed Army Medical Research Institute on a fermented mixture of...
rabbit chow and rabbit feces as described previously. Adult sand flies had with free access to a 10% solution of sucrose unless otherwise specified. Salivary glands from 3-to 10-day-old adult female flies were dissected and transferred to 10 or 20 μL Hepes 10 mM pH 7.0, NaCl 0.15 M in 1.5 mL polypropylene vials, usually in groups of 20 pairs of glands in 20 μL of Hepes saline, or individually in 10 μL of Hepes saline. Salivary glands were kept at −75°C until needed, when they were disrupted by sonication using a Branson Sonifier 450 homogenizer (Branson, Danbury, CT). Salivary homogenates were centrifuged at 10,000 g × 2 min; the supernatants were used for the experiments.

Preparation of recombinant antigen. The recombinant antigen used in this work refers to the salivary anti-clotting protein characterized in a previous publication. The GenBank accession number for the cDNA sequence is AF131932. A plasmid containing the Lutzomyia longipalpis anticoagulant sequence was used as a template to amplify only the sequence coding for the processed protein. Accordingly, we used CTA CAA GTT ACT GAG GAA CTT as a sense primer and CTT TTC TTG ACA TAC AAA ATG ATG as an antisense primer. Polymerase chain reaction (PCR) conditions were as follows: 1 cycle of 3 min at 94°C, 25 cycles of 1 min at 94°C, 30 sec at 45°C, 30 sec at 72°C, and a final cycle of 10 min at 72°C. Amplification of a single product was confirmed in a 1.5% agarose gel. The obtained PCR product was immediately ligated to pTrcHis2-TOPO TA expression vector (Invitrogen, San Diego, CA) following the manufacturer’s protocol.

Cell transformation. Cells were transformed with the ligation mixture, plated in LB-agar-ampicillin (100 μg/mL) plates and incubated overnight at 37°C. Isolated colonies were picked and transferred to 5 mL of LB-ampicillin (100 μg/mL). The cultures were grown at 37°C for 6 hours and stored at 4°C until PCR analysis. Five μL of the cultures were used as a template for a PCR reaction used to determine the presence of the anti-coagulant insert. We utilized the same sense and antisense primers described above and the following PCR conditions: 1 cycle of 5 min at 75°C, 1 cycle of 4 min at 94°C, 35 cycles of 30 sec at 94°C, 30 sec at 40°C, and 45 sec at 72°C. Amplification of a PCR product of about 420 base pairs (bp) suggested the presence of the insert. Colonies containing the insert were regrown and their plasmids extracted and sequenced in both directions to confirm the presence of the anti-coagulant and the orientation of the sequence.

Expression of recombinant anticoagulant. A 500 mL cell culture containing the anticoagulant expression vector was grown in LB-ampicillin (100 μg/mL) at 37°C, shaking at 300 rpm, until it reached 1 unit O.D. (600 nm). Isopropyl thiogalactoside (1 mM) was added to the culture and incubated for 6 hours at 37°C. Expression of the recombinant protein was confirmed by SDS-PAGE (4–20% gel). The control sample was a cell culture with the pTrcHis2-TOPO vector alone. Cells were harvested by spinning for 20 min at 1,500 × g (4°C). The pellet was resuspended in Tris-HCl pH 7.4, 20 mM EDTA, and lysozyme (8 mg/mL) was added to the culture. The culture was left at room temperature for one hour with intermittent shaking. Sodium chloride (0.5 M final concentration) and triton X-100 (2.5% final) were added and the sample was incubated at room temperature for 30 min with shaking. The culture was then spun at 10,000 × g for 50 min (4°C). The pellet was resuspended with Tris-EDTA-1% triton solution and the mixture sonicated three times for 30 sec then cooled for 1 min on ice. The cells were centrifuged again and the entire procedure repeated five times with washes of the pellet using Tris-EDTA buffer. The pellet was then stored at −75°C until use. The pellet was solubilized in GuHCl 6 M, Tris-HCl pH 8.0, EDTA 2 mM and sonicated three times for 30 sec with 1 minute cooling on ice. After sonication, the solution was incubated for two hours at room temperature and then centrifuged at 25,000 × g × 30 min at 4°C. The supernatant was recovered, DTE was added for a final concentration of 65 mM, and it was incubated at room temperature for 2 hours. The solution was then added to a refolding buffer (Tris-HCl 0.1 M pH 8.0, L-arginine-HCl 0.5 M, GSSG 0.9 mM, and EDTA 2 mM), and the mixture was incubated for 36 hours at 4°C, then dialyzed against Tris-HCl 20 mM pH 7.4. After dialysis, the recombinant protein was purified by reverse phase HPLC using a 60 min gradient of 10–60% acetonitrile with trifluoroacetic acid 0.1%, at 2 mL/min using a 1 cm × 25 cm octadecyl-silica column model 218TP510 (Vydac, Hesperia, CA). Absorbance at 280 nm was monitored. Fractions of the main peak were pooled, and analyzed by reducing SDS-PAGE, mass spectrometry, and Edman degradation. The pooled fractions were homogenous and yielded the expected amino-terminal sequence ALLQVTVKELSDGKKIFISK, with the two first amino acids being from the vector.

Anti-insect antigen serology by ELISA. ELISA plates were coated with 5 pairs of salivary glands/mL (approximately 5μg protein/mL), or with 1 μg recombinant protein/mL in carbonate buffer (NaHCO₃, 0.45 M, Na₂CO₃, 0.02 M, pH 9.6) overnight at 4°C. After three washes with PBS-0.05% Tween, the plates were blocked for 1 hour at 37°C with PBS-0.1% Tween plus 0.05% BSA. Sera were diluted 1:100 with PBS-0.05% Tween and incubated overnight at 4°C. After further washings, the wells were incubated with alkaline-phosphatase-conjugated anti-human IgG (Sigma, St. Louis, MO) at a 1:1,000 dilution for 45 minutes at 37°C. Following another washing cycle, the color was developed for 30 minutes with a chromogenic solution of p-nitrophenylphosphate in sodium carbonate buffer pH 9.6 with 1mg/mL of MgCl₂.

Preparation of Western blots. For Western blots, we used 16% Tris-Glycine, 1.0 mm thick gels, running with sodium dodecyl sulfate (Novex, San Diego, CA). The gel type, recommended for 2D gels, has one small side well for prestained molecular weight markers (myosin, bovine serum albumin, glutamic dehydrogenase, alcohol dehydrogenase, carbonic anhydrase, myoglobin, lysozyme, aprotinin, and insulin, chain B, obtained from Novex) and a large single well, to which were added the contents of 40 pairs of salivary glands (approximately 40 μg of protein) treated with 8% SDS and 4% 2-mercaptoethanol in Tris-HCl buffer 0.5 M pH 6.8, 10% glycerol, and 1% bromophenol blue dye, and heated for 5 min at 100°C. The gel was developed with Tris-Glycine buffer according to the manufacturer’s instructions and transferred to nitrocellulose using a Blot-Module for the XCell II Mini-Cell (Novex). After transfer, free sites were blocked with PBS-0.05% Tween plus 0.5% non-fat dried milk for two hours at 37°C. The strips were washed with
PBS-0.05% Tween and incubated with sera (diluted 1:100 in PBS-0.05% Tween and 0.5% nonfat dried milk) overnight at 4°C. Following further washings, strips were incubated with rabbit alkaline-phosphatase-conjugated anti-human IgG for 1 hour at 37°C. Blots were then washed and developed with nitro blue tetrazolium and 5-bromo-4-chloro-3-indolyl phosphate.

**Anti-Leishmania serology by ELISA.** Leishmania were cultivated in Schneider’s medium, and the organisms disrupted by freeze-thaw cycles and then centrifuged at 40,000 g for 20 minutes. Protein content of the supernatant was determined by Lowry’s method. Samples were stored at −20°C until used. The test was performed as previously described, and has been documented to yield specific (98%) and sensitive (99%) responses in serological tests. The spectrophotometric absorbance values (OD) of 1/100 dilution of normal sera samples (mean ± 3 SD) were used as the cutoff for a positive response.

**Delayed-type hypersensitivity test.** Delayed-type hypersensitivity (DTH) tests were performed as previously described. Briefly, cutaneous DTH was assessed by intradermally inoculating patients with L. chagasi protein. Induration was measured after 48 hours. A positive test was defined as induration greater than 5 mm at maximum width.

**Study population.** Sera used in the present study were obtained from an epidemiological survey of visceral leishmaniasis in an endemic region of São Luiz, capital of Maranhão State. During the prospective study, anti-Leishmania DTH and serology were performed twice a year in children whose age was less than 7 years during 1996–1997. Sera from individuals with visceral leishmaniasis or those who were positive either by serology or DTH in the first survey were not used in the present report. A visceral leishmaniasis case was defined by the following criteria: fever, weight loss, hepatosplenomegaly, and anemia. Positivity in the anti-leishmanial tests reported here indicates a recent conversion determined by a sensitive and specific ELISA and/or by DTH. Thus, none of the individuals in the data set had disease, and all had had negative responses to leishmanial antigen during the preceding six-month period. Accordingly, anti-Leishmania-reacting children were presumably exposed recently to the pathogen. Assuming that recent seroconversion represents infection and a positive DTH is a marker of protection against leishmaniasis in subclinical cases, classified children into four groups: (Group I) positive serology and positive DTH, (Group II) positive serology and negative DTH, (Group III) negative serology and positive DTH, and (Group IV) negative serology and negative DTH. Informed consent was obtained from parents or legal guardians of minors. The project was approved by an appropriate institutional review board from the Oswaldo Cruz Foundation.

**Statistical analysis.** The data obtained in this work was not normally distributed, even after several attempts at transformation. Accordingly, the Kruskal-Wallis Analysis of Variance on Ranks (KW-ANOVA) was used to evaluate significance among group medians. The Kolgorovov-Smirnov test was used to test for a normally distributed population. Dunn’s procedure was used to perform pairwise multiple comparisons. Correlation tests were done with the Spearman rank order test. A P-value < 0.05 was used to establish the significance level. A P < 0.01 is indicated in the text to be highly significant. Sigmastat version 2.0 (Jandel Scientific, San Raphael, CA) was used to perform the statistical tests.

**RESULTS**

To investigate whether a human population from an endemic area of visceral leishmaniasis had IgG antibodies against salivary gland antigens of the vector sand fly Lutzomyia longipalpis, we measured by ELISA the antibody level in sera of 60 children, which were stratified into four groups according to their reaction to both the DTH reaction (D+ or D−) and antibody level (S+ or S−) (Figure 1). The KW-ANOVA test indicated that the group medians were significantly different (P = 0.001). Dunn’s pairwise multiple comparison procedure indicated that the significant (P < 0.05) comparisons were between the groups S+D+ and S+D− and between S+D+ and S−D−.

Because the positive reaction described above could derive from nonspecific antibody responses, we measured the reactivity of the sera against salivary homogenates of the Old World sand fly, Phlebotomus papatasi (Figure 2). Although the optical densities of the ELISA tests were greater than when using L. longipalpis salivary homogenates, no significant differences were found among the medians of the four groups as tested by KW-ANOVA (P = 0.233).

Because large epidemiological investigations using salivary gland antigens are prohibitive due to the need for dissection of thousands of pairs of salivary glands, we tested the use of a recombinant salivary protein as an antigen for ELISA studies with 15 of the same sera in each group described above (Figure 3). The KW-ANOVA test indicated...
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FIGURE 2. Human serum IgG response against salivary gland homogenate of the sand fly Phlebotomus papatasi. Fifteen sera in each class were used, except for the S+D− group, where 14 sera only were used. Other conditions as in Figure 1.

FIGURE 3. Human serum IgG response against recombinant salivary anti-clotting protein from the sand fly Lutzomyia longipalpis. Fifteen sera in each class are represented, except for the S+/H11002 D+/H11002 group where 13 sera were used. Other conditions as in Figure 1.

FIGURE 4. Western blots indicating human IgG serum reactivity against Lutzomyia longipalpis salivary gland antigens separated by SDS-polyacrylamide gel electrophoresis. 1 = positive serum 2 = negative serum. The results are representative of 4 positive and 4 negative sera, with the positive sera being the four most-reactive sera by enzyme-linked immuno sorbent assay against L. longipalpis salivary homogenate, and the negative sera being the four least-reactive sera.

the differences among the medians to be highly significant \((P < 0.001)\). A pairwise multiple comparison test indicated that there were significant differences \((P < 0.05)\) between groups S+/D− and S−D− and groups S+/D− and S−D+.

To test whether the serologic response to the recombinant antigen correlated with that measured against whole salivary antigen as well as with the other two variables tested, namely, DTH induration and antibody to leishmanial antigen, we performed a Spearman rank order correlation test on these four variables by pooling the results reported in Figures 1 and 3. The results (Table 1) indicate that the response to the recombinant antigen correlates significantly with serological antibody response to leishmanial antigen \((P < 0.001)\), and that anti-saliva antibody response correlates \((P = 0.002)\) with DTH response to Leishmania. Unexpectedly, no correlation was found between responses to the recombinant antigen and whole salivary homogenate.

We evaluated by Western blots the complexity of the antigenic mixture in salivary gland homogenates of L. longipalpis using a selected sample of sera that tested either positive or negative by ELISA against L. longipalpis salivary homogenates. Sera with the 4 highest (thus designated positive) and 4 lowest (designated negative) OD readings in ELISA were tested in the blots. Results indicate that there are 6 main antigens, of which 3 are commonly recognized by sera from individuals with low response to Lutzomyia antigen. Antigens with molecular weights of approximately 6, 12, 36, and 96 kDa were recognized only by sera from individuals with serum IgG antibodies to Lutzomyia antigen (Figure 4).

DISCUSSION

Results presented in this paper indicate that children positive to leishmanin as assayed by a skin test react positively to salivary gland antigen of the sand fly L. longipalpis (Figure 1, Table 1). This serum IgG antisalivary homogenate response correlated positively with DTH intensity (Table 1). The complexity of the antigenic mixture comprised in the salivary gland homogenate is indicated by Western blots,
where up to 6 bands were observed in some positive sera. These results have a two-fold implication: 1) individuals exposed to Leishmania recognize salivary gland antigens of the vector sand fly, a finding reported here for the first time and 2) to the extent that response to the vector salivary gland antigen is protective in the development of disease (a possibility substantiated in laboratory models of leishmaniasis), the possibility exists to track this cohort of children to investigate their disease outcome according to their anti-salivary gland titer. This last implication, if proven negative (high anti-saliva titer negatively correlated with disease outcome), can give support for development of an anti-vector-based vaccine for leishmaniasis.

Results indicate that individuals with low ELISA reactivity to Lutzomyia salivary antigen recognize 3 antigens bands by Western blots, but that up to 6 bands are recognized by sera from individuals having high antibody titers in the same ELISA test. The positive bands found in ELISA-negative individuals could be either the result of low exposure to the sand fly or due to cross-reactivity of some Lutzomyia antigens with other arthropod antigens. Indeed, antibody responses to the tick vector of Lyme’s disease correlated with Borrelia burgdorferi antibody titer and also with ELISA titer against salivary gland homogenates of the non-vector tick Dermacentor variabilis, or even the mosquito Aedes aegypti. Use of a recombinant salivary protein from the tick Ixodes scapularis is now being used as a marker of exposure to the vector of Lyme disease, where antibody responses to the recombinant salivary calreticulin homologue correlate with the degree of engorgement of the tick on human. Serum IgG responses against Phlebotomus papatasi did not differ among the 4 groups tested (Figure 2). Although this result supports the conclusion that the response to L. longipalpis antigens is not an artifact due to nonspecific polyclonal IgG activation, it does not exclude the possibility of cross reactivity between antigens of L. longipalpis and P. papatasi, or, for that matter, any other arthropod antigen. Indeed, the greater optical densities shown in the P. papatasi ELISA could represent cross-reactivity of P. papatasi salivary antigens with some widespread antigen recognized by all four groups of sera.

Thorough evaluation of sand fly salivary components to the epidemiology or control of leishmaniasis will require large amounts of antigen, which is not feasible due to technical difficulties in rearing and dissecting sand flies. Until recently, only one salivary gland cDNA was cloned from L. longipalpis salivary glands, coding for the relatively small (6.5 kDa) vasodilator maxadilan. More recently, 11 new cDNA clones were identified, 9 of which encode secretory material as indicated by their leader sequences or isolation and amino-terminal sequence of their coded proteins. One such salivary gland cDNA, coding for the salivary anti-clotting protein was expressed in E. coli. Large amounts of product were obtained in inclusion bodies, which were further purified by HPLC to serve as an antigen in the present study. Reactivity to recombinant anti-clotting protein correlated positively with serum anti-Leishmania titers. However, it did not correlate with DTH response or with the response against total salivary homogenate, indicating that the response to individual salivary components may not be representative of that to the total homogenate.

Immune reaction to salivary homogenate is complex, as indicated by the Western blot data (Figure 4). Each antigen in the homogenate may elicit a different pattern of response in different individuals, and may have a different temporal pattern of induced reactivity according to the timing and intensity of exposure. Ultimately, production and testing of several recombinant antigens may yield a mix that will substitute for the whole salivary homogenate. However, individual antigens may have unique value, as indicated by the recombinant anti-clotting protein. Serum anti-recombinant protein IgG levels correlated positively with serum IgG levels against leishmanial antigen, indicating that this recombinant antigen may be a good marker of vector exposure. Ultimately, before either saliva or recombinant antigens can be used for widespread ELISA screening, two controlled, basic studies should be accomplished: 1) the frequency with which exposed humans develop antibodies to those antigens will have to be determined; and 2) the specificity and sensitivity of an ELISA to detect those antibodies will have to be determined.

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**Table 1**

<table>
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<th>Delayed type hypersensitivity</th>
<th>α-saliva IgG</th>
<th>α-Leishmania IgG</th>
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<td>α-TAC IgG (n = 59)</td>
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r = correlation coefficient. 
P = probability.
REFERENCES


