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Analysis of salivary gland transcripts of the sand fly *Lutzomyia ayacuchensis*, a vector of Andean-type cutaneous leishmaniasis

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Abstract

The saliva of blood sucking insects contains potent pharmacologically active components that assist them in counteracting the host hemostatic and inflammatory systems during blood feeding. In addition, sand fly salivary proteins affect host immunity and have the potential to be a vaccine against *Leishmania* infection. In the present study, the salivary gland transcripts of *Lutzomyia ayacuchensis*, a vector of cutaneous leishmaniasis in Ecuadorian and Peruvian Andes, were analyzed by sequencing randomly selected clones of the salivary gland cDNA library of this sand fly. This resulted in the identification of the most abundant transcripts coding for secreted proteins. These proteins were homologous to the salivary molecules present in other sand flies including the RGD-containing peptide, PpSP15/SL1 family protein, yellow-related protein, putative apyrase, antigen 5-related protein, D7 family protein, and 27 kDa salivary protein. Of note, homologues of maxadilan, an active vasodilator abundantly present in saliva of *Lutzomyia longipalpis*, were not identified. This analysis is the first description of salivary proteins from a sand fly of the subgenus *Helcocyrtomyia* and from vector of cutaneous leishmaniasis in the New World. The present analysis will provide further insights into the evolution of salivary components in blood sucking arthropods.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found in the online version, at http://dx.doi.org/10.1016/j.meegid.2012.08.024

Keywords

Lutzomyia ayacuchensis; Salivary gland; Transcript; Bioinformatics; cDNA library

1. Introduction

Phlebotomine sand flies are hematophagous insects of the family Psychodidae in the order Diptera, and approximately 80 species can transmit Leishmania protozoa, the causative agents of leishmaniasis (Munstermann, 2005; Kato et al., 2010a). During blood-feeding, sand flies inject saliva containing various physiologically active substances including anticoagulants, vasodilators, and inhibitors of platelet aggregation to facilitate successful acquisition of a blood meal (Ribeiro, 1995; Ribeiro and Francischetti, 2003; Valenzuela, 2005). The infection of mammalian hosts with Leishmania protozoa occurs during the blood feeding of an infected female sand fly where the parasites are co-inoculated into the hosts together with sand fly saliva (Rohoušová and Volf, 2006; Oliveira et al., 2009). Sand fly saliva was shown to facilitate the transmission of *Leishmania* to mammalian hosts and to exacerbate the disease (Titus and Ribeiro, 1988; Theodos et al., 1991; Lima and Titus, 1996). Alternatively, cellular immune response to the salivary components protects the host from Leishmania infection (Belkaid et al., 1998, 2000; Kamhawi et al., 2000; Valenzuela et al., 2001; Gomes et al., 2008; Oliveira et al., 2008; Tavares et al., 2011). Humoral immunity to sand fly salivary components was shown in the case of Leishmania (Viannia) braziliensis to exacerbate infection (Oliveira et al., 2008). In addition, recent studies are showing that sand fly salivary proteins can be used as a specific marker of sand fly exposure and to use them to evaluate potential risks of *Leishmania* transmission (Teixeira et al., 2010; Vlkova et al., 2011). The profile of salivary components has been defined in 8 Old World Phlebotomus species; *Phlebotomus (P.) papatasi* [a vector of *Leishmania (Leishmania) major*] (Valenzuela et al., 2001), P. ariasi [a vector of L. (L.) infantum] (Oliveira et al., 2006), P. perniciosus [a vector of L. (L.) infantum] (Anderson et al., 2006), P. argentipes [a vector of L. (L.) donovani] (Anderson et al., 2006), P. duboscqi [a vector of L. (L.) major] (Kato et al., 2006), P. arabicus [a vector of L. (L.) tropica] (Hostomská et al., 2009), P. tobbi [a vector of L. (L.) infantum] (Rohoušová et al., 2012), and P. sergenti [a vector of L. (L.) tropica] (Rohoušová et al., 2012). On the other hand, the salivary components of the New World Lutzomyia species have been analyzed only in Lutzomyia (Lu.) longipalpis, a vector of L. (L.) infantum [a synonym of L. (L.) chagasi] causing visceral leishmaniasis (Charlab et al., 1999; Valenzuela et al., 2004). To date, no information is available on the repertoire of salivary transcripts from vectors of the New World cutaneous leishmaniasis. In the present study, to obtain further insight into the salivary components of *Lutzomyia* species, the salivary gland transcriptome analysis was performed on Lu. ayacuchensis, which is a proven vector of L. (L.) mexicana (Takaoka et al., 1990; Kato et al., 2005, 2008) and L. (V.) peruviana (Caceres et al., 2004; Kato et al., 2008), causative agents of cutaneous leishmaniasis in Andean areas of Ecuador and Peru, respectively. This is the first report of salivary proteins from a sand fly of the subgenus *Helcocyrtomyia* and from vector of cutaneous leishmaniasis in the New World.

2. Materials and methods

2.1. Lutzomyia ayacuchensis salivary glands

Sand flies were captured using protected human bait at Huigra ($2^{\circ}20'S$, $78^{\circ}58'W$; 1200 m above sea level) in the Department of Chimborazo, Ecuador, where Andean-type cutaneous leishmaniasis caused by *L*. (*L*.) *mexicana* is endemic. The sand flies were dissected and identified at the species level based mainly on the morphology of their spermathecae (Young

and Duncan, 1994). Salivary glands of *Lu. ayacuchensis* were dissected and transferred to tubes containing RNA*later* (Ambion, Austin, TX) and kept at -20 °C.

2.2. Construction of salivary gland cDNA library

Lu. ayacychensis salivary gland mRNA was isolated from 38 pairs of salivary glands using the Micro FastTrack mRNA isolation kit (Invitrogen, San Diego, CA). The PCR-based cDNA library was prepared following the instructions for the SMART cDNA library construction kit (BD-Clontech, Palo Alto, CA) with some modifications (Valenzuela et al., 2004). The quality of the cDNA was checked by agarose gel electrophoresis and the absence of smaller fragments derived from degraded mRNA was confirmed. The obtained cDNA library was fractionated using a Chromaspin 1000 column (BD-Clontech) into small (approximately 400–800 bp), medium (approximately 800–1200 bp) and large (>1200 bp) transcripts based on their electrophoresis profiles on a 1.1% agarose gel. Pooled fractions were ligated into Lambda TriplEx2 vector (BD-Clontech) and packaged into lambda phage (Stratagene, La Jolla, CA).

2.3. Sequence analysis of cDNA library

Single isolated plaques were picked from the plate using sterile wooden sticks and placed into 50 μ l of water. Amplification of cDNA was performed in a volume of 15 μ l using a pair of primers, PT2F1 (5'-AAG TAC TCT AGC AAT TGT GAG C-3') and PT2R1 (5'-CTC TTC GCT ATT ACG CCA GCT G-3'), Premix Taq (Takara Bio, Shiga, Japan) and 3 μ l of template DNA. After an initial denaturation at 75 °C for 3 min and following 95 °C for 4 min, PCR amplification was performed with 35 cycles of denaturation (95 °C, 1 min), annealing (50 °C, 1 min) and polymerization (72 °C, 2 min). PCR products were cleaned using Multiscreen PCR cleaning plates (Millipore Corporation, Bedford, MA) and used as templates for cycle-sequencing kit (Applied Biosystems, Foster City, CA) with PT2F3 primer (5'-TCT CGG GAA GCG CGC CAT TGT-3'). Cycle-sequencing products were cleaned using sephadex and MultiScreen HV plates (Millipore Corporation), dried and stored at -20 °C. Sequencing was performed on an ABI 3730xl DNA sequencer (Applied Biosystems).

2.4. Bioinformatics

Expressed sequence tags (ESTs) were trimmed of primer and vector sequences and clustered. The ESTs were grouped based on nucleotide homology of 95% identity over 100 residues using the BLASTn algorithm (Altschul et al., 1997). The assembly of the ESTs into transcript contigs was done using the CAP3 algorithm, generating a consensus sequence (Huang, 1992). Contigs and singletons (contig containing only one sequence) were compared using BLASTx or BLASTn (Altschul et al., 1997) of the non-redundant (NR) protein database of the National Center of Biological Information (NCBI), the gene ontology database (GO) (Ashburner et al., 2000), and the Conserved Domains Database (CDD) that includes all Pfam (Bateman and Birney, 2000), SMART (Schultz et al., 1998) and COG protein domains in the NCBI (Marchler-Bauer et al., 2002). Additionally, contigs were compared using BLASTn (Altschul et al., 1997) to custom databases of mitochondrial (mitpla) and rRNA (rrna) nucleotide sequences. Identification of putative secreted proteins was conducted using SignalP server (Bendtsen et al., 2004).

2.5. T cell epitope prediction

For the T cell epitope prediction, ProPred MHC Class-II Binding Peptide Prediction Server (Singh and Raghava, 2001) was utilized. HLA class II-binding peptides were searched on the 51 different HLA-DR alleles, and the promiscuous epitopes were selected from the *Lu. ayacuchensis* salivary protein sequences tested that were predicted to bind at least 20 alleles.

2.6. Phylogenetic analysis

The sequences that had homologies with secreted proteins by BLASTx analyses were aligned with CLUSTAL W software (Thompson et al., 1994) and examined using the program MEGA (Molecular Evolutionary Genetics Analysis) version 4.0 (Tamura et al., 2007). Phylogenetic trees by the neighbor-joining method were constructed with the distance algorithms available in the MEGA package. Bootstrap values were determined on 1000 replicates of the data sets.

3. Results and discussion

3.1. Sequencing of Lutzomyia ayacuchensis salivary gland cDNA library

Lu. ayacuchensis salivary gland cDNA library was constructed, and sequencing was performed on randomly selected 1152 clones. As a result, 768 high-quality sequences were obtained. Three categories of expressed genes were derived from the manual annotation of the contigs: secreted, housekeeping and unknown. The putative secreted category comprised 45.7% of the clusters and 74.6% of the total sequences. The high ratio of transcripts encoding secreted proteins were also reported in other phlebotomine sand flies (Valenzuela et al., 2001; Oliveira et al., 2006; Anderson et al., 2006; Kato et al., 2006; Hostomská et al., 2009; Rohoušová et al., 2012). The housekeeping category had 35.0% of the clusters and 16.3% of the total sequences. Finally, the category of ''unknowns" comprised 19.3% of the cluster and 9.1% of the sequences.

3.2. Housekeeping genes

The clusters of sequences attributed to housekeeping genes (94 clusters with 125 sequences in total) were further divided into 18 subgroups, according to their possible function (Table 1). The two largest subgroups were associated with "Energy production and conversion" (24 sequences in 20 clusters) and "Translation, ribosomal structure and biogenesis" (23 sequences in 19 clusters). Six sequences in 6 clusters, which represent conserved proteins with unknown function, were classified as "unknown conserved". Other sequences were identified with homology to housekeeping genes and were associated with transport, metabolism, signal transduction, and cell structure, among other potential activities.

3.3. Putative secreted proteins

The transcripts coding for secreted proteins were further analyzed using the BLASTx program for comparison to the NCBI non-redundant protein database. Table 2 lists clusters for the most abundant transcripts coding for the putative secreted salivary gland proteins from Lu. ayacuchensis. The table was arranged from the most abundant to the least abundant transcripts found in the cDNA library. The nomenclature for the transcripts on the cDNA library is the following: Lay = $\underline{L}u$. ayacuchensis, $S = \underline{s}alivary$ glands, and the number (i.e. 45) denotes the contig number on the cDNA library where a contig is a cluster of identical transcripts. Many of the isolated transcripts code for proteins previously identified from the saliva of sand flies, Lu. longipalpis and P. perniciosus, including RGD-containing peptide, 14 kDa salivary protein, antigen 5-related protein, yellow-related protein and SL1-like protein (Table 2). Table 3 shows the classification of transcripts coding for putative secreted proteins in Lu. ayacuchensis salivary glands. Of the 573 transcripts associated with putative secreted proteins, the most abundant transcript (18.7%) had homology with that of RGDcontaining peptide from salivary glands of Lu. longipalpis. The following abundant transcripts were SL1-like protein (14.3%) and 14 kDa salivary protein (12.4%), both of which belong to a family of PpSP15/SL1 originally identified as 15 kDa protein from P. papatasi saliva. Additional transcripts coding for secreted proteins include homologs to yellow-related protein (11.1%), putative apyrase (7.9%), D7-related salivary protein (7.3%),

27 kDa salivary protein (6.6%), antigen 5-related protein (6.5%), 16.4 kDa salivary protein (5.6%), and others (Table 3).

3.3.1. RGD-containing peptide—The most abundant sequences found in the *Lu. ayacuchensis* salivary gland transcriptome contain C-terminal RGD (Arg-Gly-Asp) sequence; a motif that has been shown to bind integrins such as $\alpha\nu\beta3$ and $\alpha\nu\beta5$ (Hynes, 1992). Ten transcripts of RGD-containing peptide (LayS38–LayS47) were identified in the cDNA library of *Lu. ayacuchensis* salivary glands. These *Lu. ayacuchensis* RGD-containing molecules have a predicted molecular mass of 5.3 kDa in the mature form and a calculated isoelectric point of 3.44 (Table 2). The only homologous protein has been identified in the salivary gland transcriptome of *Lu. longipalpis* (LuloRGD), and speculated to act as an inhibitor of platelet aggregation via its RGD motif; although, the function has not yet been characterized (Charlab et al., 1999). Alignment of the RGD-containing peptide showed a high level of homology among contigs, and an RGD-motif in the C-terminal was strictly conserved in all molecules (Fig. S1).

3.3.2. PpSP15/SL1 family—The PpSP15/SL1 family of proteins has been reported only in sand flies and the biological function during blood-feeding remains unknown. Interestingly, immunization of mice with PpSP15 protected against Leishmania major infection co-injected with *P. papatasi* saliva, suggesting that this family of protein is a vaccine candidate for Leishmania infection (Valenzuela et al., 2001). Two groups of molecules were identified from the Lu. ayacuchensis salivary gland cDNA library, SL-1-like proteins and 14 kDa salivary proteins (PpSP14-like protein), as members of the PpSP15/SL1 family. Twenty-five contigs (LayS48-LayS72) showed the highest homology with SL1 protein from Lu. longipalpis, and six contigs (LayS32–LayS37) coding for a 14 kDa salivary protein were more homologous to the 14.6 kDa salivary protein from P. perniciosus (PpeSP09) (Table 2). Alignment of PpSP15/SL1 family showed a high level of homology among SL-1-like proteins and among 14 kDa salivary proteins (Figs. S2 and S3). When both groups of proteins from Lu. ayacuchensis were aligned with PpSP15/SL1 family proteins from other sand flies, only six cysteine residues and three other amino acids (P, F, and A) were conserved in the amino acid sequence of mature proteins (Fig. 1A), reflecting the divergence among PpSP15/SL1 family proteins in sand flies (Anderson et al., 2006). Since the proteins of this family are immunogenic and potential vaccines for Leishmania infection (Valenzuela et al., 2001), T cell epitopes were analyzed using ProPred MHC Class-II Binding Peptide Prediction Server. We identified two potential T cell epitopes, YRITKKHIE and IIHYYRCVV, in Lu. ayacuchensis SL-1-lile proteins (Fig. S2), and two potential epitopes, IKNKAVDGS and FKYESAINSY, in 14 kDa salivary proteins (Fig. S3). A phylogenetic analysis of sand fly PpSP15/SL1 family proteins revealed that SL-1-like proteins and 14 kDa salivary proteins from Lu. ayacuchensis constructed separate clades (Fig. 1B).

3.3.3. Yellow-related protein—Yellow-related proteins are abundantly expressed in salivary glands of sand flies (Valenzuela et al., 2001; Oliveira et al., 2006; Anderson et al., 2006; Kato et al., 2006; Hostomská et al., 2009; Rohoušová et al., 2012). The proteins of this family are immunogenic and host antibody responses to this protein can be a potential marker for sand fly exposure (Teixeira et al., 2010; Vlkova et al., 2011). Interestingly, prior DNA vaccination of mice with PpSP44 coding for a 44 kDa yellow-related protein from *P. papatasi* saliva primed strong humoral immunity and exacerbated subsequent *L. major* infection in the presence of sand fly saliva (Oliveira et al., 2008). In contrast, a yellow-related protein from *Lu. longipalpis* saliva, LJM11, conferred protective cellular immunity in mice against *L. major* infection plus sand fly saliva. Other *Lu. longipalpis* yellow-related proteins, LJM111 and LJM17, were not protective, suggesting that structural features are a

determinant for the host immunity against yellow-related proteins (Xu et al., 2011). Regarding the biological function, yellow-related proteins, LJM11, LJM111 and LJM17, were shown to act as high affinity binders of proinflammatory biogenic amines such as serotonin, catecholamines and histamine, suggesting that the proteins play a role for the reduction of inflammation during sand fly blood-feeding (Xu et al., 2011). In the Lu. ayacuchensis cDNA library, five contigs (LayS22-LayS24, LayS117 and LayS118) are putative yellow-related proteins, of which LayS117 was truncated in the 5' region. When these proteins were aligned with salivary yellow-related proteins from other sand fly species, four cysteine residues were conserved in the amino acid sequence of mature proteins (Fig. 2A and S4). In addition, an amino acid motif, T-x(52,63)-Y-Q-x(85,90)-[FY]-x(44,46)-Fx(54)-[IVL]-x(45,46)-[FY]-x-[TS]-D-x(13)-[NT]-x-[QHFL], recently identified in the ligand binding pocket of yellow-related proteins (Xu et al., 2011) were conserved in those from Lu. ayacuchensis (Fig. 2A), suggesting that these proteins work as anti-inflammatory agents in Lu. ayacuchensis saliva by binding biogenic amines. Two potential Nglycosylation sites were present in yellow-related proteins from Lu. ayacuchensis, but the positions of LayS118 were different from those of others (LayS22-LayS24) (Fig. 2A). T cell epitopes were analyzed on the immunogenic proteins of this family, and we identified seven potential epitopes in LayS22-LayS24 (Fig. S4A) and 5 potential epitopes in LayS118 (Fig. S4B). A phylogenetic analysis of sand fly yellow-related proteins revealed that LayS22, LayS23 and LayS24 were closer related to LJM11 whereas LayS118 were closer related to LJM17 from Lu. longipalpis saliva (Fig. 2B). The result suggested that effect on the host immunity of LayS118 is different from those of LayS22, LayS23 and LayS24.

3.3.4. Apyrase—Apyrases are nucleoside triphosphate-diphosphohydrolases present in a variety of organisms. In the saliva of blood sucking arthropods, apyrases function to hydrolyze ADP in a Ca²⁺-dependent manner and inhibit ADP-induced platelet aggregation to facilitate blood feeding (Ribeiro and Francischetti, 2003; Faudry et al., 2004). Apyrases identified in sand fly saliva belong to the *Cimex* apyrase family (Valenzuela et al., 2001; Hamasaki et al., 2009). Thirteen contigs (LayS8–LayS14 and LayS16–LayS21) were homologous to putative apyrase from *Lu. longipalpis* (LuloAPY). A multiple sequence alignment analysis together with LuloAPY confirmed the close relationship among putative apyrases of *Lu. ayacuchensis* (Fig. S5).

3.3.5. D7 family—D7 family proteins are abundantly expressed in salivary glands of blood-feeding Diptera such as mosquitoes, sand flies, black flies, and biting midges (James et al., 1991; Valenzuela et al., 2002; Campbell et al., 2005; Andersen et al., 2009). To date, the function of sand fly D7 proteins remains to be elucidated; however, several D7 family proteins have been functionally characterized in mosquitoes. Hamadarin from *Anopheles stephensi* saliva was identified as a blood coagulation inhibitor affecting the activation of the plasma contact system (Isawa et al., 2002). Salivary D7 proteins from *Anopheles gambiae* and *Aedes aegypti* were characterized as binders of biogenic amines such as serotonin, histamine and norepinephrine (Calvo et al., 2006). Nine contigs (LayS95–LayS103) encoding a 26.6 kDa protein were homologous to the 27 kDa D7-related salivary protein from *P. argentipes* (PagSP10) and *P. perniciosus* (PpeSP04b). In a multiple sequence alignment analysis of D7 family proteins from sand flies, 10 cysteine residues were strictly conserved in the amino acid sequence of mature proteins (Fig. 3A and Fig. S6). Alignment and phylogenetic analysis together with PpeSP04b showed that D7 family proteins of *Lu. ayacuchensis* had close relationships (Fig. 3B).

3.3.6. 27 kDa Salivary protein—The protein family is also known as PpSP32-like family. Eleven contigs (LayS83–LayS93) encoding a 27 kDa protein were homologous to the 29.2 kDa salivary protein from *Lu. longipalpis* (Table 2). These transcripts are also

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al., 2001; Oliveira et al., 2006; Anderson et al., 2006; Kato et al., 2006; Hostomská et al., 2009; Rohoušová et al., 2012). To date, no homologues of these proteins have been reported in organisms other than sand flies and the biological function remains to be clarified. Although no conserved domains were found in the amino acid sequences, these proteins contain glycine-rich sequences in the middle region (amino acids 66–179 in LayS91) (Fig. S7).

3.3.7. Antigen 5-related protein—This family of proteins belongs to the cysteine-rich secretory proteins (CRISPs) and is related to venom allergens in social wasps and ants (Lu et al., 1993; Hoffman, 1993; King and Spangfort, 2000). Transcripts coding for the members of this protein family have been identified in the salivary glands of blood sucking insects such as mosquitoes (Francischetti et al., 2002; Arcà et al., 2007), sand flies (Valenzuela et al., 2001; Oliveira et al., 2006; Anderson et al., 2006; Kato et al., 2006; Hostomská et al., 2009; Rohoušová et al., 2012) and triatomine bugs (Ribeiro et al., 2004; Santos et al., 2007; Assumpção et al., 2008; Kato et al., 2010b). Although the antigen 5 family of protein has been widely identified in blood-sucking arthropods, the function in their saliva has yet to be determined. Nine contigs (LayS73–LayS81) from *Lu. ayacuchensis* salivary glands were included in this family. A multiple sequence alignment analysis of antigen 5-related proteins with that of *Lu. longipalpis* revealed the close relationship among the proteins of *Lu. ayacuchensis* and strict conservation of 12 cysteine residues in mature forms (Fig. S8).

3.3.8. 16.4 kDa Salivary protein—Thirteen contigs (LayS120–LayS132) coded for 16.4 kDa protein containing a C-type lectin/C-type lectin-like domain (Table 2). This putative domain may function as a Ca²⁺-dependent carbohydrate-binding pocket involved in extracellular matrix organization, pathogen recognition and cell-to-cell interactions (Weis et al., 1998). Homologous proteins with molecular weight of 16.2–16.5 kDa have been identified from *Lu. longipalpis* saliva (Charlab et al., 1999; Valenzuela et al., 2004), and they were similar to the previously described anticoagulant from *Lu. longipalpis* saliva (Charlab et al., 1999; Valenzuela et al., 2004), and they were similar to the previously described anticoagulant from *Lu. longipalpis* saliva (Charlab et al., 1999). A recombinant protein of mature LayS127 protein was expressed in *Escherichia coli*, and its activity on coagulation pathway was assessed; however, the protein inhibited neither intrinsic nor extrinsic coagulation pathways (data not shown). At present, the biological function of these proteins from *Lu. ayacuchensis* 16.4 kDa proteins together with *Lu. longipalpis* 16.5 kDa protein, three cysteine residues were strictly conserved in the amino acid sequence of mature proteins (Fig. S9).

3.3.9. 11.5 kDa Salivary protein—Four contigs encoding 10.7–11.9 kDa proteins (LayS4–LayS7) were identified in the cDNA library of *Lu. ayacuchensis* salivary glands. The different size of these proteins was mainly resulted from the number of "SSDGSSG"repeat sequences with unknown function (Fig. S10). The only homologous protein has been identified in salivary glands of *Lu. longipalpis* (9 kDa salivary protein) with unknown function. The amino acid sequences of these proteins are rich in serine from the middle to the C-terminal regions (Fig. S10).

3.3.10. Other putative secreted proteins—Three contigs (LayS25–LayS27) coding for a 33.6 kDa protein had homology with a 34 kDa salivary protein from *P. ariasi*. Three contigs (LayS141–LayS143) coded for a 9.3 kDa protein. Three singletons (LayS109–LayS111) coding for a 15.9 kDa protein had homology with a 15.5 kDa salivary protein from *P. argentipes* with unknown function. Two contigs (LayS2 and LayS3) coded for a 25.9 kDa protein homologous to allergen from *Culex quinquefasciatus* belonging to the

sperm-coating protein (SCP) superfamily. Two singletons (LayS167 and LayS168) and one singleton (LayS153) coded for homologous proteins with a 32.4 kDa salivary protein and a 71 kDa salivary protein from *Lu. longipalpis*, respectively, although the sequence of LayS168 was truncated in the 5' region. A singleton (LayS147) coded for homologous protein with the 43.7 kDa salivary protein from *Lu. longipalpis*, a putative endonuclease, but the sequence was truncated in the 5' region. Singletons, LayS106 and LayS215, coded for homologous protein with the 27 kDa salivary protein from *P. ariasi* and trypsin inhibitor like cysteine rich domain containing protein from *Drosophila*, respectively.

4. Conclusions

The present study identified abundant salivary gland transcripts of *Lu. ayacuchensis*, a proven vector of *L. (L.) mexicana* (Takaoka et al., 1990; Kato et al., 2005, 2008) and *L. (V.) peruviana* (Caceres et al., 2004; Kato et al., 2008) causative agents of cutaneous leishmaniasis in Andean areas of Ecuador and Peru, respectively. The relatively divergent transcripts resulting in the generation of a variety of contigs noted for each component probably reflects population of the sand flies used in this study; that is, our cDNA library was constructed from field-captured *Lu. ayacuchensis* while most studies prepared the cDNA libraries from colonized insects. To date, the salivary components of the New World *Lutzomyia* species have been analyzed only in *Lu. longipalpis* (Charlab et al., 1999; Valenzuela et al., 2004), a subgenus *Lutzomyia* species, transmitting *L. (L.) infantum*, a causative agent of visceral leishmaniasis (Young and Duncan, 1994). On the other hand, *Lu. ayacuchensis* is a vector species of cutaneous leishmaniasis and belongs to the subgenus *Helcocyrtomyia* (Young and Duncan, 1994). Therefore, the present analysis is the first description of salivary proteins of a sand fly in the subgenus *Helcocyrtomyia* and a vector of cutaneous leishmaniasis in the New World.

In the present analysis, transcripts homologous to maxadilan, an active vasodilator abundantly present in saliva of Lu. longipalpis, were not identified. Maxadilan was originally isolated from saliva of Lu. longipalpis (Lerner et al., 1991), and its homologue was reported from Lu. neivai (Aires et al., 2005). Conversely, maxadilan-related proteins have never been identified in Old World *Phlebotomus* species (Valenzuela et al., 2001; Oliveira et al., 2006; Anderson et al., 2006; Kato et al., 2006; Hostomská et al., 2009). Phlebotomus sand flies except for P. duboscqi lack adenosine deaminase (ADA) that hydrolyze adenosine and adenosine monophosphate (AMP) (Oliveira et al., 2006; Anderson et al., 2006; Kato et al., 2006; Hostomská et al., 2009), but contain large amounts of adenosine and AMP, very active vasodilators, in their saliva (Ribeiro et al., 1999; Katz et al., 2000; Ribeiro and Modi, 2001). On the other hand, Lu. longipalpis has ADA (Charlab et al., 1999, 2000; Valenzuela et al., 2004) and lacks adenosine and AMP in the saliva (Ribeiro et al., 1989). Therefore, with an exception, salivary vasodilators of sand flies are considered to be adenosine and AMP in *Phlebotomus* species and maxadilan in *Lutzomyia* species. Unexpectedly, neither ADA nor maxadilan were identified in Lu. ayacuchensis, which is similar to *Phlebotomus* species, suggesting that salivary vasodilators of this species may be adenosine and AMP. Further analysis of small molecules in the saliva of Lu. ayacuchensis will elucidate this issue.

In conclusion, the most abundant proteins of *Lu. ayacuchensis* saliva were identified in this study. These results will provide further insights into the evolution of salivary components in blood sucking arthropods. In addition, the cDNAs and future recombinant proteins prepared from these transcripts will result in the discovery of vaccine candidates and markers of sand fly exposure as well as novel pharmacologically active compounds.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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		PduM07 (ABI15937)	
		100 PpSP12 (AAL11045)	

Fig. 1.

(A) Sequence alignment of PpSP15/SL1 family proteins from *Lutzomyia* (*Lu.*) ayacuchensis (LayS36, LayS37, LayS58, LayS59, LayS66, LayS68, LayS71, and LayS72) together with those from *Phlebotomus* (*P.*) arabicus (Pab), *P. argentipes* (Pag), *P. ariasi* (Par), *P. duboscqi* (Pdu), *P. papatasi* (Pp), *P. perniciosus* (Ppe), and *Lu. longipalpis* (Lulo). Black-shaded amino acids represent identical amino acids and gray-shaded amino acids represent conserved amino acids denote conserved cysteine residues, and an arrowhead indicates the predicted signal peptide cleavage site. (B) Phylogenetic tree analysis of amino acid

- PduM50 (ABI20148)

- PpSP14 (AAL11046)

1001 LayS36 (AK416777) LayS37 (AK416778) PagSP13 (ABA12144) PpeSP09 (ABA43057)

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sequences of PpSP15/SL1 family proteins from sand flies. Accession numbers are in parentheses and node values indicate branch support.

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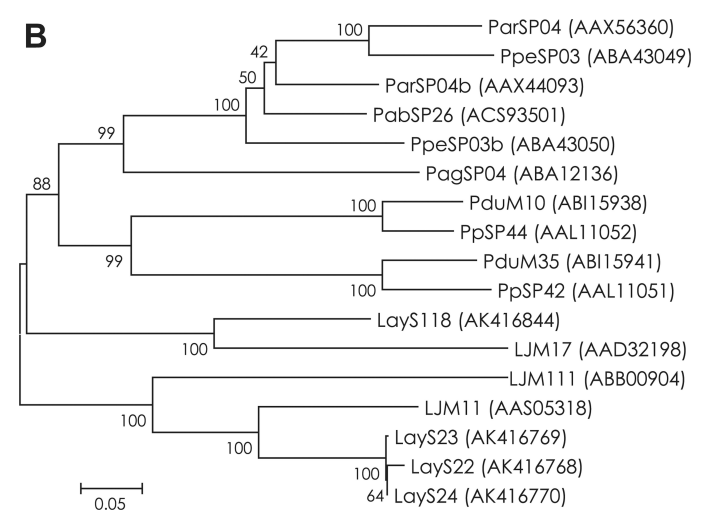


Fig. 2.

(A) Sequence alignment of yellow-related proteins from *Lu. ayacuchensis* (LayS22, LayS23, LayS24, and LayS118) together with those from *P. arabicus* (Pab), *P. argentipes* (Pag), *P. ariasi* (Par), *P. duboscqi* (Pdu), *P. papatasi* (Pp), *P. perniciosus* (Ppe), and *Lu. longipalpis* (LJM). Black-shaded amino acids represent identical amino acids and gray-shaded amino acids represent conserved amino acids. Dashes indicate gaps introduced for maximal alignment. Asterisks at the top of the amino acids denote conserved cysteine residues, and an arrowhead indicates the predicted signal peptide cleavage site. Closed circles show conserved amino acids contained in the ligand binding pocket of yellow-related protein family. Potential N-glycosylation sites of *Lu. ayacuchensis* yellow-related proteins are boxed. (B) Phylogenetic tree analysis of amino acid sequences of yellow-related proteins from sand flies. Accession numbers are in parentheses and node values indicate branch support.

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Fig. 3.

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(A) Sequence alignment of D7 family proteins from Lu. ayacuchensis (LayS101–LayS103) together with those from P. arabicus (Pab), P. argentipes (Pag), P. ariasi (Par), P. duboscqi (Pdu), P. papatasi (Pp), P. perniciosus (Ppe), and Lu. longipalpis (Lulo). Black-shaded amino acids represent identical amino acids and gray-shaded amino acids represent conserved amino acids. Dashes indicate gaps introduced for maximal alignment. Asterisks at the top of the amino acids denote conserved cysteine residues, and an arrowhead indicates the predicted signal peptide cleavage site. (B) Phylogenetic tree analysis of amino acid sequences of D7 family proteins from sand flies. Accession numbers are in parentheses and node values indicate branch support.

PpeSP10 (ABA43058)

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Table 1

Functional classification of housekeeping genes expressed in Lutzomyia ayacuchensis salivary glands

Type of transcripts	Cluster	%	Sequence	%
Energy production and conversion	20	21.3	24	19.2
Translation, ribosomal structure and biogenesis	19	20.2	23	18.4
General function prediction only	9	9.6	10	8.0
Signal transduction mechanisms	8	8.5	8	6.4
Posttranslational modification, protein turnover, chaperones	7	7.4	14	11.2
Lipid transport and metabolism	5	5.3	5	4.0
Amino acid transport and metabolism	4	4.3	4	3.2
Chromatin structure and dynamics	2	2.1	2	1.6
Inorganic ion transport and metabolism	2	2.1	2	1.6
Intracellular trafficking, secretion, and vesicular transport	2	2.1	3	2.4
RNA processing and modification	2	2.1	2	1.6
Secondary metabolites biosynthesis, transport and catabolism	2	2.1	2	1.6
Transcription	2	2.1	14	11.2
Carbohydrate transport and metabolism	1	1.1	3	2.4
Cell cycle control, cell division, chromosome partitioning	1	1.1	1	0.8
Cell wall/membrane/envelope biogenesis	1	1.1	1	0.8
Defense mechanisms	1	1.1	1	0.8
Unknown conserved	6	6.4	6	4.8
Total	94	100.0	125	100.0

Table 2

Most abundant salivary gland transcripts from Lutzomyia ayacuchensis

Cluster	Sequence name	No. of	Mature	pI	Best match to N	Best match to NR protein database			Accession No.
		seq in cluster	MW (KUa)		Accession No.	Organism	${\cal E}$ value	% Identity	
LayS45	RGD-containing peptide	06	5.3	3.44	AAD32196	Lu. longipalpis	3E-07	47	AK416785
LayS37	14 kDa Salivary protein	58	14.1	8.21	ABA43057	P. perniciosus	6E-39	50	AK416778
LayS81	Antigen 5-related protein	24	28.5	9.08	AAD32191	Lu. longipalpis	1E-135	80	AK416820
LayS24	Yellow-related protein	23	43.5	9.37	AAS05318	Lu. longipalpis	0	78	AK416770
LayS59	SL1-like protein	19	13.8	8.22	AAD32197	Lu. longipalpis	1E-51	64	AK416798
LayS91	27 kDa Salivary protein	17	27.2	10.14	AAS16906	Lu. longipalpis	2E-92	59	AK416828
LayS117*	Yellow-related protein	16			AAD32198	Lu. longipalpis	4E-77	66	AB744655
LayS118	Yellow-related protein	14	43.8	8.05	AAD32198	Lu. longipalpis	1E-164	67	AK416844
LayS127	16.4 kDa Salivary protein	11	16.4	8.05	ABB00903	Lu. longipalpis	1E-29	39	AK416852
LayS21	Putative apyrase	11	35.7	9.19	AAD33513	Lu. longipalpis	1E-123	65	AK416767
LayS142	9.3 kDa Protein	10	9.3	4.22	XP 001238639	Eimeria tenella	0.012	44	AK416859
LayS103	D7-related salivary protein	6	26.6	8.81	ABA43052	P. perniciosus	7E-86	59	AK416840
LayS36	14 kDa Salivary protein	6	14.1	8.21	ABA43057	P. perniciosus	2E-39	50	AK416777
LayS6	11.5 kDa Salivary protein	8	11.5	6.67	AAS16919	Lu. longipalpis	2E-07	33	AK416753
LayS102	D7-related salivary protein	8	26.6	8.81	ABA43052	P. perniciosus	7E-86	59	AK416839
LayS19	Putative apyrase	8	35.8	60.6	AAD33513	Lu. longipalpis	1E-127	65	AK416765
LayS58	SL1-like protein	8	13.8	8.22	AAD32197	Lu. longipalpis	1E-51	64	AK416797
LayS68	SL1-like protein	8	13.9	8.61	AAD32197	Lu. longipalpis	7E-52	64	AK416807
LayS90	27 kDa Salivary protein	8	26.6	10.08	AAS16906	Lu. longipalpis	1E-90	58	AK416827
LayS101	D7-related salivary protein	7	26.6	8.81	ABA43052	P. perniciosus	2E-86	59	AK416838
LayS66	SL1-like protein	9	13.9	8.61	AAD32197	Lu. longipalpis	1E-51	64	AK416805
LayS20	Putative apyrase	9	35.5	9.00	AAD33513	Lu. longipalpis	1E-128	65	AK416766
LayS22	Yellow-related protein	9	43.4	9.39	AAS05318	Lu. longipalpis	1E-125	75	AK416768
LayS17	Putative apyrase	5	35.5	9.11	AAD33513	Lu. longipalpis	1E-128	65	AK416763
LayS18	Putative apyrase	5	36.0	9.46	AAD33513	Lu. longipalpis	1E-128	65	AK416764
LayS23	Yellow-related protein	5	43.5	9.37	AAS05318	Lu. longipalpis	0	78	AK416769
LayS26	33.6 kDa Salivary protein	5	33.7	8.95	AAX55751	P. ariasi	6E-91	53	AK416772

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Cluster	Sequence name	No. of	Mature	pI	Best match to N.	Best match to NR protein database			Accession No.
		seq in cluster	Mw (kDa)		Accession No.	Organism	E value	% Identity	
LayS44	RGD-containing peptide	5	5.3	3.47	AAD32196	Lu. longipalpis	8E-07	46	AK416784
LayS71	SL1-like protein	5	13.8	8.22	AAD32197	Lu. longipalpis	2E-50	64	AK416810
LayS72	SL 1-like protein	5	13.8	8.43	AAD32197	Lu. longipalpis	3E-51	64	AK416811
LayS100	D7-related salivary protein	4	26.6	8.61	ABA43052	P. perniciosus	4E-86	59	AK416837
LayS131	16.4 kDa Salivary protein	4	16.4	8.32	ABB00903	Lu. longipalpis	7E-26	39	AK416856
LayS3	Allergen-related protein	4	26.0	5.68	XP 001865175	Culex quinquefasciatus	8E-31	35	AK416750
LayS64	SL 1-like protein	4	13.9	8.61	AAD32197	Lu. longipalpis	5E-48	59	AK416803
LayS97	D7-related salivary protein	4	26.6	8.81	ABA43052	P. perniciosus	2E-86	59	AK416834
LayS128	16.4 kDa Salivary protein	3	16.4	8.05	ABB00903	Lu. longipalpis	1E-29	39	AK416853
LayS129	16.4 kDa Salivary protein	3	16.4	7.69	ABB00903	Lu. longipalpis	2E-29	40	AK416854
LayS16	Putative apyrase	3	36.6	9.34	AAD33513	Lu. longipalpis	1E-125	64	AK416762
LayS27	33.6 kDa Salivary protein	3	33.7	8.69	AAX55751	P. ariasi	1E-90	52	AK416773
LayS46	RGD-containing peptide	3	5.3	3.44	AAD32196	Lu. longipalpis	1E-07	49	AK416786
LayS47	RGD-containing peptide	3	5.3	3.44	AAD32196	Lu. longipalpis	2E-07	47	AK416787
LayS70	SL 1-like protein	3	13.9	7.98	AAD32197	Lu. longipalpis	5E-51	64	AK416809
LayS77	Antigen 5-related protein	3	28.6	9.21	AAD32191	Lu. longipalpis	1E-135	80	AK416816
LayS89	27 kDa Salivary protein	3	26.6	10.08	AAS16906	Lu. longipalpis	5E-90	58	AK416826
LayS98	D7-related salivary protein	3	26.6	8.81	ABA43052	P. perniciosus	3E-86	59	AK416835
LayS130	16.4 kDa Salivary protein	2	16.4	8.05	ABB00903	Lu. longipalpis	2E-29	40	AK416855
LayS132	16.4 kDa Salivary protein	2	16.4	8.05	ABB00903	Lu. longipalpis	2E-29	40	AK416857
LayS2	Allergen-related protein	2	26.0	5.46	XP 001865175	Culex quinquefasciatus	5E-30	34	AK416749
LayS60	SL 1-like protein	2	13.9	7.98	AAD32197	Lu. longipalpis	7E-51	64	AK416799
LayS61	SL 1-like protein	2	13.8	8.22	AAD32197	Lu. longipalpis	1E-51	65	AK416800
LayS62	SL 1-like protein	2	13.8	8.52	AAD32197	Lu. longipalpis	6E-51	64	AK416801
LayS63	SL 1-like protein	2	13.8	8.23	AAD32197	Lu. longipalpis	1E-51	64	AK416802
LayS65	SL 1-like protein	2	13.8	7.98	AAD32197	Lu. longipalpis	1E-51	64	AK416804
LayS67	SL 1-like protein	2	13.8	8.35	AAD32197	Lu. longipalpis	1E-50	64	AK416806
LayS69	SL 1-like protein	2	13.9	8.22	AAD32197	Lu. longipalpis	4E-51	64	AK416808
LayS7	11.5 kDa Salivary protein	2	10.8	8.36	AAS16919	Lu. longipalpis	2E-07	34	AK416754
LayS78	Antigen 5-related protein	2	28.4	9.16	AAD32191	Lu. longipalpis	1E-133	80	AK416817

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E value % Identity 80 80 58 58 59 58 59

AK416818

1E-135 1E-135

Lu. longipalpis

AK416819 AK416829 AK416830

AK416832 AK416833

5E-86 1E-85

9E-91 1E-91

Lu. longipalpis Lu. longipalpis Lu. longipalpis AK416836

6E-86

P. perniciosus P. perniciosus

P. perniciosus

Accession No.

Best match to NR protein database

 \mathbf{pI}

Mature Mw (kDa)

No. of seq in cluster

Sequence name

Cluster

Accession No. Organism

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ABA43052	8.81	26.6	2	D7-related salivary protein 2	LayS99
ABA43052	8.81	26.6	2	D7-related salivary protein	LayS96
ABA43052	8.81	26.6	2	D7-related salivary protein	LayS95
AAS16906	10.14	27.2	2	27 kDa Salivary protein	LayS93
AAS16906	10.14	27.3	2	27 kDa Salivary protein	LayS92
AAD32191	9.01	28.5	2	Antigen 5-related protein	LayS80
AAD32191	9.08	28.4	2	Antigen 5-related protein	LayS79

* Truncated in the 5' region.

Table 3

Classification of transcripts coding for putative secreted proteins in Lutzomyia ayacuchensis salivary glands

	Cluster	Sequence	% Sequence
RGD-containing peptide	10	107	18.7
SL1-like protein	25	82	14.3
14 kDa Salivary protein	6	71	12.4
Yellow-related protein	5	64	11.1
Putative apyrase	13	45	7.9
D7-related salivary protein	9	42	7.3
27 kDa Salivary protein	11	38	6.6
Antigen 5-related protein	9	37	6.5
16.4 kDa Salivary protein	13	32	5.6
9.3 kDa Protein	3	16	2.8
11.5 kDa Salivary protein	4	12	2.1
33.6 kDa Salivary protein	3	9	1.6
Allergen-related protein	2	6	1.0
Others	9	12	2.1
Total	122	573	100.0