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# The sialotranscriptome of *Amblyomma triste*, *Amblyomma parvum* and *Amblyomma cajennense* ticks, uncovered by 454-based RNA-seq

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## Abstract

**Background:** Tick salivary constituents antagonize inflammatory, immune and hemostatic host responses, favoring tick blood feeding and the establishment of tick-borne pathogens in hosts during hematophagy. *Amblyomma triste, A. cajennense* and *A. parvum* ticks are very important in veterinary and human health because they are vectors of the etiological agents for several diseases. Insights into the tick salivary components involved in blood feeding are essential to understanding vector-pathogen-host interactions, and transcriptional profiling of salivary glands is a powerful tool to do so. Here, we functionally annotated the sialotranscriptomes of these three *Amblyomma* species, which allowed comparisons between these and other hematophagous arthropod species.

**Methods:** mRNA from the salivary glands of *A. triste, A. cajennense* and *A. parvum* ticks fed on different host species were pyrosequenced on a 454-Roche platform to generate four *A. triste* (nymphs fed on guinea pigs and females fed on dogs) libraries, one *A. cajennense* (females fed on rabbits) library and one was *A. parvum* (females fed on dogs) library. Bioinformatic analyses used *in-house* programs with a customized pipeline employing standard assembly and alignment algorithms, protein databases and protein servers.

**Results:** Each library yielded an average of 100,000 reads, which were assembled to obtain contigs of coding sequences (CDSs). The sialotranscriptome analyses of *A. triste, A. cajennense* and *A. parvum* ticks produced 11,240, 4,604 and 3,796 CDSs, respectively. These CDSs were classified into over 100 distinct protein families with a wide range of putative functions involved in physiological and blood feeding processes and were catalogued in annotated, hyperlinked spreadsheets. We highlighted the putative transcripts encoding saliva components with critical roles during parasitism, such as anticoagulants, immunosuppressants and anti-inflammatory molecules. The salivary content underwent changes in the abundance and repertoire of many transcripts, which depended on the tick and host species.

**Conclusions:** The annotated sialotranscriptomes described herein richly expand the biological knowledge of these three *Amblyomma* species. These comprehensive databases will be useful for the characterization of salivary proteins and can be applied to control ticks and tick-borne diseases.

**Keywords:** Ticks, Amblyomma cajennense, Amblyomma parvum, Amblyomma triste, Salivary gland, Saliva, Transcriptome, RNA-seq

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### Background

Arthropods have developed several feeding habits during the course of evolution, and within this taxa, hematophagy evolved independently over millions of years, resulting in significant differences among hematophagous arthropods [1,2]. Most tick species are obligatory blood-feeding ectoparasites and transmit more infectious agents—bacteria, protozoa and viruses—to humans and animals than any other arthropod vectors [3].

Among the hematophagous ticks, there are 130 species of hard ticks of the genus *Amblyomma* (Acari: Ixodidae) [4], which has considerable medical and veterinary importance in the Americas and the Caribbean [5]. The *A. cajennense* tick is associated mainly with savannah biomes under natural conditions [6,7] and causes severe infestations in equines. Due to its low parasitic specificity, it also infests cattle, dogs, birds, capybaras [8-10] and humans [11] in the urban and peri-urban areas of Brazil, and it is the vector for *Rickettsia rickettsii*, the causative agent of spotted fever in South America [12]. Recent mitochondrial and nuclear DNA analyses suggest that *A. cajennense* is a complex of six species, such that *A. sculptum* is synonymous with *A. cajennense* ticks found in the coastal and central-western regions of Brazil [13,14].

The *A. parvum* tick may harbor *Rickettsia* species of yet unknown pathogenicity [15,16], and it is a potential vector for emerging pathogens, such as *Ehrlichia chaffeensis* (the infectious agent of human monocytotropic ehrlichiosis) [17] and *Coxiella burnetii*, which causes Q fever and chronic endocarditis in humans [18]. This tick species is found almost exclusively in dry areas [19]. It feeds on medium-size rodents, such as the yellow toothed cavy (*Galea musteloideds*) [20], but it is also considered a host generalist, as it feeds on domestic animals including cattle and goats [16].

In contrast to the above-mentioned *Amblyomma* species, the *A. triste* tick is mostly associated with marshes and environments prone to flooding [21-23]. Although the deer *Blastocerus dichotomous* is the main host for *A. triste* adults, domestic animals, wild carnivores and humans may also host *A. triste* (reviewed by Nava *et al.* [24]). Epidemio-ecological studies performed in Argentina, Brazil and Uruguay have reported specimens of *A. triste* infected with *Rickettsia parkeri* [25-28], which raises an important health issue due to suspected human cases of rickettsiosis caused by *R. parkeri* [29].

Successful blood meals and enhanced pathogen transmission result from morphological and biochemical adaptations of the tick salivary glands, which secrete bioactive molecules with profound impacts on host hemostasis and inflammation [30,31]. Additionally, salivary glands have osmoregulatory functions that enable ticks to survive in different environments for long periods away from the host [32]. Tick saliva has anti-clotting, anti-platelet, vasodilatory, anti-inflammatory and immunomodulatory components [33]. Thus, the components of tick saliva are relevant for global public health, as they may contain novel pharmacologically active compounds and provide possible targets for vaccines against ticks and the diseases they transmit [34].

The characterization of gene expression in tissues (transcriptomes) is a robust strategy to uncover tick salivary gland components and functional mechanisms. Salivary gland transcriptomes, or sialotranscriptomes, of several tick species have been described through the analysis of Expressed Sequence Tags (ESTs) obtained by Sanger sequencing of cDNA libraries [35-41], which generally yield a few thousand ESTs. Next generation sequencing (NGS) technologies allow deeper and more accurate gene expression studies because in contrast to previous sequencing methods, millions of short sequences are produced yielding data at the gigabase rather than kilobase scale of traditional cDNA libraries based on cloning [42]. The NGS transcriptional profiling approach is called RNA-seq. To date, two tick sialotranscriptomes have been described by NGS technologies, resulting in a very detailed catalogue of salivary gland transcripts from A. maculatum [43], the gulf coast tick, and from Ixodes ricinus, the castor bean tick [44].

Here, we describe the sialotranscriptome of three species of Amblyomma ticks, A. triste, A. parvum and A. cajennense, using 454-based RNA-seq. These species present distinct biology in terms of preferred habitats and hosts. The repertoire of tick saliva may be complementary to its preferred host's homeostatic and immune strategies. Ticks can acquire host genes through horizontal gene transfer [45]; therefore, ticks with different feeding habits may have different repertoires of salivary proteins. A total of six RNA-seq libraries were generated from the salivary glands of nymph and female ticks. For the first time, the transcriptional profile of a tissue from A. parvum and A. triste ticks has been generated and analyzed. Furthermore, we examined the salivary gland transcripts of a third tick, A. cajennense, through the analysis of 67,677 reads, which clustered into 4,604 putative coding sequence (CDS) contigs. Comparison of the six annotated sialotranscriptomes revealed common and unique features in the ticks. The transcript catalogues described herein provide insights to the biology of these tick species, which are important ectoparasites, pathogen reservoirs and vectors for diseases.

## Methods

### Ticks, hosts and sample preparation

The following ticks and hosts were used in this study: *A. parvum*: semi-engorged females (54 ticks) fed on dogs; *A. cajennense:* semi-engorged females (45 ticks) fed on rabbits. One pool of salivary glands (SGs) was obtained

for both A. parvum and A. cajennense. As A. triste ticks were evaluated in four parasitological conditions, SGs samples generated four pools designed NGP1 (47 nymphs fed on guinea pigs during first infestation) and NGP2 (90 nymphs fed on guinea pigs during second infestation) to designate the semi-engorged nymph samples; FD1 (87 females fed on dogs during first infestation) and FD2 (47 females fed on dogs during second infestation) to designate the semi-engorged female samples. All tick infestations were performed under laboratory conditions as described previously [16,46,47]. Nymphs of A. triste and females of A. cajennense were removed from their hosts between the third and fourth day post-feeding, and A. triste and A. parvum females were removed between the fourth and fifth day post-feeding. Within the first hour of collection, ticks were washed and dissected to obtain the SGs under sterile conditions. The SG samples were organized in pools according to the tick species, which afterwards corresponded to the number of libraries sequenced. The pairs of SGs were gently washed in ice-cold phosphate buffered saline (PBS) and stored immediately in RNAlater solution (Ambion Inc., Austin, TX, USA) at 4°C for 24 hours and then at -70°C until RNA isolation.

### **RNA** isolation

Total RNA was isolated from the six pools of tick salivary glands with a protocol optimized in our laboratory. Briefly, after centrifugation, the SG samples were removed from RNAlater solution, and Trizol reagent (Invitrogen, Life Technologies, CA, USA) was added to the samples, which were then thoroughly blended with an overhead stirrer at high speed. Chloroform was gently mixed with the homogenate, and after centrifugation at maximum speed at 4°C, the resuspended aqueous phase was placed in a clean tube containing 96-100% ethanol and mixed. RNA was purified from the mixture with an SV Total RNA Isolation System Kit (Promega Corporation, Madison, WI, USA) according to the manufacturer's instructions. RNA was quantified using a NanoDrop 1000 Spectrophotometer (Thermo Scientific, Wilmington, DE, USA), and the quality was checked with a 2100 Bioanalyzer (Agilent Technologies Inc., Santa Clara, CA, USA). RNA was stored at -70°C.

### Library preparation and pyrosequencing

The raw sequencing data were generated on the 454-Roche platform by the High-Throughput Sequencing and Genotyping Unit, DNA services branch, Roy J. Carver Biotechnology Center, University of Illinois in Urbana-Champaign, IL, USA. Briefly, messenger RNA was isolated from 20  $\mu$ g of total RNA from each pool of SGs using an Oligotex mRNA kit (Qiagen, Valencia, CA, USA). The mRNA-enriched fraction was converted to a primary cDNA library with barcoded adaptors compatible with the 454-Roche system. The cDNA libraries were quantified using a Qubit fluorometer (Invitrogen, Life Technologies, CA, USA), and the average fragment sizes were determined using a DNA 7500 chip on the 2100 Bioanalyzer. The adaptor-ligated cDNA libraries were pooled in equimolar concentrations, diluted to  $1x10^6$  molecules/µl, subjected to emulsion-based clonal amplification and pyrosequenced on a full plate using the 454 Genome Sequencer FLX system (454 Life Sciences, Roche, Branford, CT, USA) according to the manufacturer's instructions. Signal processing and base calling were performed using the bundled 454 Data Analysis Software version 2.3.

### **Bioinformatic analysis**

The raw data were downloaded from the High-Throughput Sequencing and Genotyping Unit website, and the bioinformatics workflow for the pyrosequencing data analysis was performed as previously described [43]. The software was written and provided by JMCR in Visual Basic 6.0 (Microsoft, Redmond, Washington, USA). For functional annotation, the CDS contigs were queried against several databases with the blastx, blastn and rpsblast algorithms [48]. We used the followed databases: 1) the non-redundant protein database (NR) of the National Center for Biotechnology Information (NCBI); 2) Gene Ontology (GO) FASTA subset [49]; 3) Swissprot [50]; 4) tick salivary sequences described previously [33]; 5) custom databases from GenBank containing mitochondrial and rRNA nucleotide sequences; 6) CDD from NCBI [51] containing the SMART [52], KOG [53] and PFAM [54] motifs. The functionally annotated catalog for each sialotranscriptome was manually curated and plotted in a hyperlinked Excel spreadsheet. The three spreadsheets were designated Additional file 1 AF1 (A. triste), Additional file 2 AF2 (A. parvum) and Additional file 3 AF3 (A. cajennense) and are available for download at http:// exon.niaid.nih.gov/transcriptome/sm/AF1-Atris-web.xlsx, http://exon.niaid.nih.gov/transcriptome/sm/AF2-Aparvweb.xlsx and http://exon.niaid.nih.gov/transcriptome/sm/ AF3-Acaj-web.xlsx, respectively (the weblinks are best visualized with the following browsers: Google Chrome, Mozilla Firefox and Internet Explorer). The calculation for differential transcript abundance between the libraries of the three tick species and for the libraries of A. triste ticks under different parasitological conditions (Additional file 1 AF1, columns AH-AQ) was performed with a chisquare  $(X^2)$  test using the number of reads in each CDS from different libraries; results were considered statistically significant if P < 0.05. The raw sequencing files were manipulated with the ShortRead package in Bioconductor software [55].

Hierarchical clustering (HCL) to ascertain the gene expression pattern was performed with the standard

statistical algorithm described previously [56], using the percentage of reads for each sialotranscriptome, and represented by the  $\log_2$  transformation of reads + 1. HCL employed Euclidean distance for metric calculations and the average linkage method to indicate the cluster-tocluster distance in the hierarchical trees, which were displayed as heatmaps. These analyses were performed with MeV (Multi Experiment Viewer) software [57]. To compare the different datasets from A. cajennense ticks, one library of SGs (LIBEST\_019946 [58]) was downloaded from NCBI dbEST database [59,60] and a second library (LIBEST\_USP-RP) described herein (Additional file 4 AF4) were used for sequence similarity searches against the 454-Roche dataset with the blastn algorithm. The translated amino acid sequences of some ESTs from the first two datasets were obtained with NCBI ORF Finder [61].

Multiple sequence alignments were performed with the Clustal W algorithm [62] using Bioedit software [63]. Phylogenetic analyses were determined using the Neighbor-Joining (NJ) method in MEGA 5.0 software [64], and sequences from other tick species were downloaded from the NCBI database (listed in Additional file 5 AF5). The node support of each clade was evaluated using a bootstrap analysis (1000 replicates), and the evolutionary distances were derived using the Jones-Taylor-Thornton (JTT) method [65].

### Accession numbers

Sequence deposition for *A. triste* [GenBank: GBBM0 1000001 - GBBM01008098 (BioProject 241269)], for *A. parvum* [GenBank: GBBL01000001 - GBBL01002838 (BioProject 241271)] and *A. cajennense* [GenBank: GBBK01000001 - GBBK01005770 (BioProject 241272); JZ718403 – JZ718940].

### **Ethical approval**

The experiments were approved by Animal Experimentation Ethics Committee of the Federal University of Uberlândia, protocol #033/14, according to Ethical Principles on Animal Research of the National Council for the Control of Animal Experimentation (CONCEA).

## **Results and discussion**

## Overview of the annotated sialotranscriptomes of *A. triste, A. parvum* and *A. cajennense* ticks

We generated and analyzed the transcriptional profile of salivary glands (SG) from three neotropical hard ticks belonging to the *Amblyomma* genus, obtained from six 454-based RNA-seq libraries: one library from *A. par-vum* female ticks fed on dogs, one library from *A. cajennense* female ticks fed on rabbits and four libraries from *A. triste* ticks. The sialotranscriptome of the latter species addressed different parasitological conditions

including the developmental stage of the tick, the type of host that each stage prefers [23] and the status of host exposure to ticks and anti-tick immunity. This transcriptome was thus constructed with SGs from nymphs fed on naïve guinea pigs (NGP1), nymphs fed on guinea pigs undergoing a second tick infestation (NGP2), females fed on naïve dogs (FD1) and females fed on dogs undergoing a second infestation (FD2).

The A. parvum library yielded 243,567 pyrosequencing reads, and the A. cajennense yielded library 67,677 pyrosequencing reads. The four combined A. triste libraries yielded 1,195,924 pyrosequencing reads (data not shown). Only the pyrosequencing reads longer than 149 nucleotides were used to assemble primary contigs. The primary assembled contigs (minimum of 3-4 reads per contig) were analyzed to extract putative CDS, and the CDS contigs were used to annotate each sialotranscriptome. The CDS contigs, categorized according to their putative function, were plotted in hyperlinked Excel spreadsheets available as additional files for download: A. triste (Additional file 1: AF1), A. parvum (Additional file 2: AF2) and A. cajennense (Additional file 3: AF3). The annotation of the CDS contigs were performed automatically using a suite of customized programs written in Visual Basic and afterwards manually curated and refined. The functional annotation relied on significant similarities (determined by BLAST algorithms, considering only high identity scores and low E-values) between CDS contigs and sequences in public protein databases and customized databases of known and predicted tick proteins.

The *A. triste* sialotranscriptome was generated from four libraries and consisted of 7,532 CDS contigs (90,367 reads) for NGP1, 7,391 CDS contigs (85,871 reads) for NGP2, 9,311 CDS contigs (120,717 reads) for FD1 and 9,004 CDS contigs (145,801 reads) for FD2. Together, *A. triste* libraries produced a total of 11,240 CDS from 442,756 reads (Table 1 and Additional file 1: AF1). The *A. parvum* sialotranscriptome contained 3,796 CDS contigs assembled from 104,817 reads (Table 1 and Additional file 2: AF2). The *A. cajennense* sialotranscriptome contained 4,604 CDS contigs assembled from 67,677 reads (Table 1 and Additional file 3: AF3).

Transcripts in the three sialotranscriptomes were classified into seven main categories: a) *Secreted*, having a signal peptide and/or significant BLAST hits with secreted proteins; b) *Housekeeping*, constitutive genes required for the basal maintenance of tissues; c) *Unknown* and d) *Unknown Secreted*, which had non-significant BLAST hits or no matches against any database, being that the latter category (d) contained signal peptide sequences; e) *Transposable Elements*, mobile elements in genomes; f) *Viral*, transcripts with similarities to viral proteins and g) *Leishmania-related*, transcripts with

### Table 1 General distribution of reads in coding sequence contigs for the sialotranscriptomes of three Amblyomma ticks

	Amblyomma cajennense				Amblyomma parvum				Amblyomma triste*			
Category	CDS <sup>a</sup>	Reads <sup>a</sup>	Reads/CDS	Reads <sup>b</sup>	CDS <sup>a</sup>	Reads <sup>a</sup>	Reads/CDS	Reads <sup>b</sup>	CDS <sup>a</sup>	Reads <sup>a</sup>	Reads/CDS	Reads <sup>b</sup>
Secreted	1,015	16,480	16.2	24.4	493	22,796	46.2	21.7	1,861	76,193	40.9	17.2
Housekeeping	2,805	44,246	15.8	65.4	2,653	65,952	24.9	62.9	6,854	305,129	44.5	68.9
Unknown	338	3,245	9.6	4.8	272	10,739	39.5	10.2	1,071	26,373	24.6	6.0
Unknown secreted	424	3,509	8.3	5.2	359	5,141	14.3	4.9	1,345	33,418	24.8	7.5
Transposable Elements	16	38	2.4	0.1	19	189	9.9	0.2	97	1,405	14.5	0.3
Viral	6	159	26.5	0.2	-	-	-	-	10	193	19.3	0.04
Leishmania-related	-	-	-	-	-	-	-	-	2	45	22.5	0.01
Total	4,604	67,677			3,796	104,817			11,240	442,756		

a. total number; b. percentage; \*Combined data from the four libraries of *A. triste*: salivary glands from nymph ticks fed on guinea pig during first (NGP1) and second (NGP2) infestation; salivary glands from female ticks fed on dogs during first infestation (FD1) and second (FD2) infestation.

similarities to *Leishmania* proteins. These data are numerically summarized in Table 1.

On average, 20% of the reads in the three sialotranscriptomes were annotated as putative secreted proteins, whereas the majority of the reads (ranging from 62.9% to 68.9%) were annotated as housekeeping genes (Table 1). The *Unknown* and *Unknown Secreted* categories together presented an average of 13% of reads, and this relatively low proportion is a consequence of previous efforts to describe transcripts in salivary glands from several tick species [33].

Because Amblyomma ticks are vectors of several microorganisms that are important public health threats, we examined the sialotranscriptomes for CDS related to microbes. The A. triste and A. cajennense sialotranscriptomes contained 10 and 6 CDSs, respectively, that matched viral-like proteins (Additional file 1: AF1 and Additional files 3: AF3). Forty-five reads were annotated into two CDS coding for Leishmania-like proteins in the A. triste sialotranscriptome (Additional file 1: AF1). Interestingly, several reports describe the presence of Leishmania parasites in ticks [66]. Additionally, we recovered a CDS from A. parvum (Ap-19721) that was primarily annotated as ferredoxin protein and that best matched Rickettsia sibirica (Additional file 2: AF2), an etiologic agent of North Asian tick typhus. Rickettsial strains phylogenetically related to R. parkeri, R. africae, and R. sibirica have been found in several Amblyomma ticks in Brazil [67]. Collectively, these findings support the role of Amblyomma ticks as pathogen vectors.

In total, eighty-seven protein families (602,347 reads) were shared within all *Amblyomma* libraries (Additional file 1: AF1, Additional file 2: AF2 and Additional file 3: AF3). Some protein families that were absent in a given sialotranscriptome may be expressed in different developmental stages or even are supplied by members of other families that are functionally redundant.

## Comparative analysis of the *A. triste*, *A. parvum* and *A. cajennense* sialotranscriptomes

The *Secreted* and *Housekeeping* categories contained several functional subcategories, and most of these were common between two or three sialotranscriptomes. To understand how the transcriptional profile of these different functional subcategories changes across the three *Amblyomma* species, we performed a hierarchical clustering using the abundance of transcripts (relative percentage of reads) for each sialotranscriptome. For the *A. triste* sialotranscriptome, we deconstructed it according to the experimental conditions (two developmental stages on once- and twice-infested hosts).

The expression data are displayed as heatmaps where the black to red colors indicate low and high expression levels, respectively (Figure 1). The abundances of the 33 subcategories of housekeeping genes together with the Unknown category genes were clustered in two main groups according to the expression level (Figure 1a, transcript clusters numbered 1 and 2). For instance, weakly expressed transcripts encoding calpain-like proteins, sulfatases, O-methyltransferases and catalases are in the first cluster, whereas the second cluster is composed of more abundant transcripts such as those encoding protein synthesis machinery, energy metabolism, transcription machinery, cytoskeletal and signal transduction proteins.

Regarding the clustering of *Amblyomma* species using the expression patterns of housekeeping genes (Figure 1a, library clustering from right to left), the libraries from *A. parvum* and *A. triste* formed the main cluster, where the two minor clusters were arranged according to the host and life stage of *A. triste* ticks, nymphs fed on guinea pigs (NGP) or females fed on dogs (FD) (Figure 1a, library clustering from right to left). Notably, libraries from different tick species using the same host for blood meals were clustered closer (i.e.,



Figure 1 Hierarchical clustering of putative protein families from the sialotranscriptome of Amblyomma ticks. The heatmaps display the expression level of the protein families (rows) for each sialotranscriptome (columns). a) Families classified as housekeeping proteins. b) Families classified as secreted proteins. The color scale represents the  $log_2$  transformation of % of reads + 1. The transition of colors from blue to red represents an increase in the % of reads. Black color indicates the % of reads <0.01; red color indicates the % of reads  $\geq$  4.5. The length of branches in hierarchical trees indicates the degree of similarity between objects, either tick libraries (columns) or protein families (rows). Side scales represent the node height (clusterto-cluster distance). NGP1: nymphs fed on tick-naïve guinea pigs; NGP2: nymphs fed on guinea pigs undergoing a second infestation; FD1: females fed on tick-naïve dogs (first infestation) and FD2: females fed on dogs undergoing a second infestation. FR: females fed on rabbits. The clustering method used Euclidian distance with average linkage

more similar), as observed for A. parvum FD and A. triste FD1 and FD2 libraries (Figure 1a). Additionally, the branch length of the sub-cluster formed with A. triste FD libraries indicated that the expression pattern was slightly more similar for female ticks than nymph ticks. Although the patterns in the four A. triste libraries were similar, the FD libraries were enriched for transcription machinery, protein export machineries and transport and storage, while the NGP libraries had a higher abundance of extracellular matrix/adhesion-related transcripts. The A. cajennense library was externally positioned and distant from the other female tick libraries. In addition to the species difference, this distance (i.e., less similarity with the others) might have been influenced by the blood meal source for this tick, which was fed on rabbits (FR). Almost all A. cajennense transcript families had lower expression compared with the other libraries. These transcription profiles suggest that the amount and content of proteins produced by salivary glands depends on the tick species and life stage as well as the host background.

Although the *Secreted* category contained about onefifth of the transcripts (Table 1), many diverse protein families were observed, and approximately double the number of housekeeping protein families were present, considering all subcategories of putative secreted proteins (Additional file 1: AF1, Additional file 2: AF2 and Additional file 3: AF3). Despite the lower numbers of transcripts than the housekeeping genes, this large repertoire with relatively small quantities of secreted proteins may have driven the performance of ticks during blood meals. Thus, tick saliva relies on the specificity of this secreted protein repertoire, which in turn may vary between tick species, which live in different habitats and have a range of preferred hosts.

The hierarchical clustering of the putative secreted transcripts resulted in distribution pattern similar to housekeeping protein families, such that *A. cajennense* 

was placed as external member and the *A. triste* FD libraries were grouped with the *A. parvum* FD library instead of the *A. triste* NGP libraries (Figure 1b, library clustering). In addition, the branches were longer between the *A. triste* libraries in the hierarchical tree for secreted transcripts, reflecting less similarity. In general, the expression profile of transcripts encoding putative secreted proteins was less well defined across species than the housekeeping genes. The transcription of secreted proteins may be more prone to parasitological changes (e.g., nymphs versus females) as well as the different host environments (e.g., dogs or guinea pigs, once- or twice-infested).

Compared with the housekeeping transcripts, the expression level of genes encoding putative secreted proteins was lower. Most transcripts encoding secreted proteins were expressed at a low level (Figure 1b, transcript cluster 1), whereas a few families showed medium to high expression levels (Figure 1b, transcript cluster 2). Additionally, HCL analysis showed that some cluster 1 transcripts encoding secreted proteins differed significantly between the *A. cajennense* and the *A. parvum* and *A. triste* libraries (Table 2).

Among the families with low expression (Figure 1b, protein cluster 1), we highlight the metalloproteases, Salp-15, antimicrobial peptides and evasins. The metalloproteases were represented by reprolysin, neprilisin-type, and other proteins that have metal-dependent proteolytic activity to gelatin, fibrinogen and fibronectin, which could help regulate host inflammatory and immune responses [68,69].

Salp-15 protein was first described in *I. scapularis* saliva as an immunomodulatory protein that inhibits  $CD4^+$  T cell differentiation through its interaction with the CD4 co-receptor and favors the transmission of diverse *Borrelia* species to their hosts [70,71]. Salp-15 was only found in FD libraries from *A. triste*. As *A. triste* is a vector of human rickettsiosis [23], the putative Salp-15 might interfere with the host immune response, supporting both the hematophagous ectoparasite as well as the pathogenic bacteria carried by the tick. Hence, it will be interesting to further examine *A. triste* as a pathogen vector.

The superfamily of antimicrobial peptides is formed by CDSs encoding defensins, microplusins, lysozymes and antigen 5 families. Ticks produce antimicrobial peptides as part of their innate immunity; these peptides control and combat invading microorganisms [72-74]. These families were expressed in the sialotranscriptomes of all three *Amblyomma* species, indicating their importance for the protection of the ticks against potentially harmful microorganisms during hematophagy. Notably, these transcripts were differentially expressed, with higher expression in *A. cajennense* followed by the *A. triste* and *A. parvum* female libraries (Table 2).

All libraries contained CDSs encoding evasin, a salivary protein previously identified in *R. sanguineus* ticks [75]. Evasins hinder the host immune response by binding chemokines and impairing cell migration [75]. These transcripts were more highly expressed in the *A. cajennense* and female libraries compared with the *A. triste* nymph libraries (Figure 1b and Table 2).

Aside from unknown secreted proteins, the most abundant transcripts encoding secreted proteins were members of the GRP category (Figure 1b, transcript cluster 2). Significant differences in GRP expression were observed between female libraries of the three species of ticks; A. cajennense females (FR) contained significantly less GRP transcripts than A. parvum females and A. triste fed on dogs (Table 2). A. triste NGP libraries also had an abundance of GRP expression (Figure 1b). We previously showed that the amount and types of GRP are related to the biology of the tick species, namely, whether it is monoxenous or heteroxenous and whether it has a short or long hypostome [76]. In this study, all ticks were heteroxenous Longirostrata ticks, and A. triste and A parvum females were fed on dogs. One possible explanation is the different ecosystems where these ticks live. A triste is found on hosts living in marshes, where humidity is high, and A. parvum is found in the dry savannah. In addition to their roles in tick attachment to host skin, GRPs are also hygroscopic proteins. The ticks in our experiments were reared in controlled laboratory conditions, but basal expression of GRP may have been maintained each species.

The Kunitz domain family is subclassified according to the number of KU domains in each sequence (i.e., Monolaris, Bilaris, Trilaris, Tetralaris, Pentalaris and Hexalaris). Members of this family are abundant in the sialotranscriptomes of hematophagous arthropods, such as flies and ticks [33,77,78], and typically, they are protease inhibitors that impair various processes including blood coagulation and promote vasodilatation [79]. Therefore, this family of proteins facilitates hematophagy and plays an important role during tick engorgement. The Kunitz domain family was represented by significantly fewer transcripts in the A. triste FD transcriptome (Table 2). In all developmental stages, hosts and their immune statuses, this family was expressed at a lower level in A. triste than in A. cajennense and A. parvum (Figure 1b and Table 2). This finding merits further investigation to determine if other mediators of antihemostasis may substitute for this family in the sialome of A. triste. Another hypothesis is that temporal differences affect the coagulation and fibrinolytic systems of different mammalian hosts [80]: whilst A. triste and A. parvum females were both fed on dogs, the genetic composition of these species of ticks may be complementary to the hemostatic mechanisms of their preferred natural hosts, ungulates and rodents, respectively.

Protein families	Number of reads											
	A. cajennense	vs		A. parvum	A. cajennense	VS		A. triste <sup>#</sup>	A. parvum	VS		A. triste <sup>#</sup>
	Obs.	Exp.	Obs.	Exp.	Obs	Exp.	Obs.	Exp.	Obs.	Exp.	Obs.	Exp.
Metalloproteases	1,439*	946.1	906	1,399	1,439*	835.5	828	1,431	906*	775.1	828	959.9
8.9 kDa Family	1,400*	1,268	1,832	1,964	1,400*	712	582	1,270	1,832*	1,122	582	1,292
ML Domain	0	14.9	38*	23.1	0	3.6	10	6.4	38*	22.3	10	25.7
Cystatin	62*	24.3	0	37.7	62*	33	30	59	0	13.9	30*	16.1
23 kDa Family	8	3.1	0	4.9	8	18,3	43*	32.7	0	20	43*	23
TIL domain	256	414.7	801*	642.3	256	243.6	422	434.4	801*	568.4	422	654.6
Basic tail Superfamily	941*	427.7	149	662.3	941*	481.7	400	859.3	149	255.1	400*	293.9
Carboxypeptidases	0	14.1	36*	21.9	0	5	14*	9	36*	23.2	14	26.8
Cystein Proteases	132	194.5	349*	286.5	132*	81.5	89	139.5	349*	195.5	89	242.5
Serine Proteases	424*	199.3	69	293.6	424*	188.9	88	323.1	69	70	88	86.9
Secreted Nucleases	784*	352.5	87	518.5	784*	354	175	604.9	87	116.9	175*	145
Antimicrobial Peptides	202*	116.2	94	179.8	202*	156	226	271.9	94	145.5	226*	174.5
Evasins	459*	284.1	265	439.9	459*	198.3	93	353.7	265*	166.4	93	191.6
Thyropin	12	18	34	28	12	21.9	49*	39.1	34	38.6	49	44.4
DAP-36 immunosuppressant family	58	82.8	153*	128.2	58*	44.9	67	80.1	153*	102.2	67	117.8
Kazal domain containing peptides	9	20	42*	31	9	10.8	21	19.2	42*	29.3	21	33.7
Phosphatidylethanolamine-binding protein	2	11.8	28*	18.2	2	10.8	28*	19.2	28	26	28	30
Salp-15 family	0	0	0	0	0	6.5	18*	11.5	0	8.4	18*	9.6
Serpin	133*	89.5	95	138.5	133*	60.4	35	107.6	95*	60.4	35	69.6
Serum Amyloid A Proteins	17	127.5	308*	197.5	17	12.2	17	21,8	308*	151	17	174
Hirudin	533*	211.1	5	326.9	533*	194.3	8	346.7	5	6	8	7
Prokineticin domain-containing peptides	0	18.4	47*	28.6	0	1.8	5	3.2	47*	24.2	5	27.8
Carboxypeptidase inhibitor	121*	49	4	76	121*	53.5	28	95.5	4	14.9	28*	17.1
Ferritin	103	245.2	522*	379.8	103	113.9	214	203.1	522*	342.1	214	393.9
Glycine-rich proteins	1,026	2,888	6,336*	4,474	1,026	1,288	2,512*	2,250	6,336*	4,021	2,512	4,827
Mucins	957	1,218	2,147*	1,886	957	1,923	4,396*	3,430	2,147	3,041	4,396*	3,502
Histamine binding proteins	1,311*	1,170	1,671	1,812	1,311	1,626	3,216*	2,901	1,671	2,271	3,216*	2,616
Kunitz Domains	4,033*	3,122	3,748	4,659	4,033*	1,916	1,190	3,307	3,748*	2,210	1,190	2,728
Lipocalins	1,585	1,745	2,864*	2,703	1,585*	1,103	1,486	1,968	2,864*	2,022	1,486	2,328

Table 2 Transcripts coding secreted proteins differentially represented between A. cajennense, A. parvum and A. triste ticks

Calculation was performed with chi-square test ( $\chi^2$ ) using number of reads assigned for protein families, according functional annotation (Additional file 1: AF1, Additional file 2: AF2 and Additional file 3: AF3); Obs:: Observed; Exp:: Expected. The higher number of reads observed between the comparisons are highlighted in bold if statically significant according the  $\chi^2$  test result (\* *P* < 0.05). #: The comparisons were performed only with the library of *A. triste* female ticks fed on dog during the first infestation (FD1).

Serine/threonine-rich proteins containing potential glycosylation sites are termed mucins. Mucins are mainly present in mucosal tissues and function as mechanical barriers to damage from pathogens and toxins [81]. Transcripts encoding members of this protein family have been recovered in several tick sialotranscriptomes [35,39,43]. We found that these transcripts were highly abundant in all *A. triste* libraries (Figure 1b). Within the female libraries of *Amblyomma* ticks, these transcripts were significantly higher in the FD libraries from *A. triste* and *A. parvum*  (Table 2). Although this evidence suggests that the host may influence the expression of mucins, for instance, *A. cajennense* FR library had fewer of these transcripts than other libraries (Figure 1b and Table 2), it is possible that other protein families may overcome the protection provided by them, such as antimicrobial peptides that were significantly more transcribed in *A. cajennense* (Figure 1b and Table 2).

Transcripts for histamine-binding proteins (HBPs) were more abundant in *A. triste* females fed on dogs

(FD1 and FD2), and the expression of other lipocalins was significantly higher in *A. parvum* compared with *A. cajennense* and *A. triste* females (Figure 1b and Table 2). Histamine is an important mediator of inflammation, and its concentration varies depending on the host skin. Tick histamine-binding proteins have a high affinity for histamine [82] and they suppress inflammation caused by tick bites and saliva. HBPs was previously characterized in the saliva of the *R. appendiculatus* tick [83], and transcripts encoding HBP and lipocalin are also present in several other tick sialo-transcriptomes [36,39,43].

Additionally, several CDS contigs displayed a high level of identity with protein sequences previously described in the sialotranscriptomes of other species in the Amblyomma genus, namely, A. maculatum [43] and A. variegatum [41], which employed the same bioinformatics strategy for the sequence assembly. Most of the characterized salivary proteins for the Amblyomma genus-for example, Amblin [84], Variegin [85] and Serpins [86,87]-have anti-hemostatic functions. One CDS similar to Amblin was present in the A. triste sialotranscriptome (At-143156), and we found sequences in the sialotranscriptome of A. triste (At-34965, At-35820 and At-30336) and A. cajennense (Ac-23315) annotated as serpins with at least 82% of amino acid identity with the serpins/lospins from A. americanum [86,87]. CDSs similar to the well-characterized thrombin inhibitor Variegin could not be found in our analyses, because any small protein sequences (less than 49 amino acids) were excluded during bioinformatics steps; nevertheless, this does not mean that transcripts for Variegin or any other small proteins were not expressed. CDSs similar to proteins that presents non-anti-hemostatic functions, -such as ferritins, iron carriers found in several ticks [88], and Hebraein [72], an antimicrobial described for A. hebraeum— were also presented in A. parvum (Ap-3247, 98% identical to A. americanum ferritin) and A. cajennense (Ac-5954, 98% identical to A. americanum ferritin and Ac-6284/Ac-6285, 86% identical to Hebraein). Protein sequence similarities with other Amblyomma species can be explored in the three annotated sialotranscriptomes (see BLAST results for Acari protein database in the Additional file 1: AF1, Additional file 2: AF2 and Additional file 3: AF3).

## Amblyomma triste: changes in the sialotranscriptome depend on host immunity and the life stage of the tick

The sialotranscriptomes of *A. triste* ticks in four different biological conditions (NGP1, NGP2, FD1 and FD2) were compared to determine which transcripts were shared and unique in these four libraries (Figure 2). Most transcripts (68.7%) were shared by all four libraries. In the libraries from ticks feeding on hosts infested once or twice and

thus presenting different levels of immunity, the total number of unique transcripts was decreased in the nymphs (from 1% in NGP1 to 0.2% in NGP2) and adults (from 1.2% in FD1 to 0.8% in FD2). Only 0.3% of the transcripts are uniquely shared between the libraries representing the first (FD1 and NGP1) or second (FD2 and NGP2) tick infestation, regardless of the host background or the tick life stage. A small increase in uniquely shared reads was observed when ticks in the same life stage were feeding on the same host, for example, overlaps between FD1 and FD2 (11.5%) or NGP1 and NGP2 (4.2%) (Figure 2a).

Among the numerous families annotated within the *Secreted* category, we emphasize the abundance of 12 major families that were significantly differentially expressed between the libraries (Figure 2b and Additional file 1: AF1). The CDSs for Thyropin, Serpin, Ferritin and Cuticle-like protein were more abundant in nymphs. Tick serpins are serine protease inhibitors that play key roles in the regulation of blood coagulation and inflammation through the inhibition of cathepsin G, chymase and vertebrate elastase [89,90]. They were highly expressed in NGP1, with an average 10-fold decrease in NGP2 (Figure 2b). This decrease might be caused by the acquired resistance of guinea pig hosts during the second infestation, which affected the gene expression in NGP2 by unknown mechanisms.

The optimal development of eggs and offspring viability requires the ingestion of a large volume of blood by female ticks. Due to the large amount of blood consumed, hematophagous arthropods must process a high content of iron in hemoglobin, which in excess becomes toxic. In this context, tick ferritins neutralize iron toxicity through the metabolism and storage of iron [91,92], thus allowing successful parasitism. Ferritin transcripts were significantly more abundant in the sialotranscriptome of the nymphal stage (Additional file 1: AF1, columns AH-AQ), possibly in preparation for the female stage of development in which rapid engorgement with blood and iron accumulation occurs.

Cuticle proteins are also an essential factor for parasite success. In addition to forming part of the external tegument of the tick exoskeleton, the cuticle lines internal tissues, such as salivary gland ducts, trachea and the digestive tract [93]. Thus, cuticle is important for molting, and it supports physiological changes in internal tissues during parasite development and hematophagy.

In the *A. triste* sialotranscriptome, we observed that transcripts encoding secreted ferritins and cuticle-like proteins had the same expression patterns in libraries from the same tick life stage (NGP1 and NGP2; FD1 and FD2). Both transcripts were highly expressed in the

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#### (See figure on previous page.)

**Figure 2 Comparative analyses of the sialotranscriptome of** *Amblyomma triste* ticks. a) Venn diagram showing the percentage of common and unique reads in the salivary glands of *A. triste* during blood feeding in four parasitological conditions. NGP1: nymphs fed on naïve guinea pig; NGP2: nymphs fed on guinea pig undergoing a second tick infestation; FD1: females fed on naïve dogs and FD2: females fed on dogs undergoing a second tick infestation; GP1: nymphs fed on guinea pig undergoing a second tick infestation; FD1: females fed on naïve dogs and FD2: females fed on dogs undergoing a second tick infestation. b) Abundance of some transcripts coding for putative secreted proteins across the different conditions addressed herein. All of these protein families were differently expressed between libraries, NGP1 vs. NGP2 and FD1 vs. FD2, according to a chi-square test (*P* < 0.05) using the number of reads for each library in a given functionally annotated contig. \*Metalloproteases of the reprolysin family.

nymphal stage, which had 10-to-20-fold more reads than the female libraries (Figure 2b). The potential high expression of these two types of proteins on nymphs fed on guinea pig hosts, either once or twice infested, may explain at molecular level why specific tick life stages have different performances depending on the host species [46]. For instance, *A. triste* nymphs are more efficient parasites when they are fed on guinea pigs, whereas capybaras are best for *A. triste* female ticks.

Conversely, other transcripts were more abundant in the salivary glands of female ticks, such as those encoding metalloproteases, TIL domain-containing proteins and histamine-binding protein (Figure 2b). In I. scapularis, metalloproteases in saliva inhibit endothelial cell proliferation, angiogenesis and fibrinogenolysis [68,94], processes important for wound healing at the tick bite site. In addition, metalloproteases are involved in the metabolism of several peptides that regulate inflammation and immunity [69]. Thus, tick metalloproteases are essential for a complete blood feeding. Another secreted protein family important for the "tick hematophagy cocktail" contains the histamine-binding proteins, which were more abundant in female tick libraries. Because dogs are not the preferred hosts for adult ticks, the increased expression of these two proteins (TIL and HBP) in A. triste females might be a mechanism to overcome an impaired blood meal, in which females struggle to feed properly on the dog host. Lastly, the TIL domaincontaining proteins were also more abundant in female libraries. These cysteine-rich antimicrobial proteins inhibit serine proteases that are important for bacterial growth [73,95].

While we cannot confirm that the expression patterns are stage-specific or host-specific because expression data from nymphs fed on dogs and from females fed on guinea pigs is not available, the transcripts highlighted above were consistent in the pairs of libraries (NGP1 and NGP2; FD1 and FD2), suggesting that both the life stage and host may be driving gene expression in salivary glands.

## Amblyomma cajennense: the contribution of this new sialotranscriptome

The salivary glands of *A. cajennense* female ticks have been the subject of two other gene expression studies, conducted by Batista et al. ([58], here designated LIBEST\_019946) and Anatriello et al. [96], but the most recent study has not made the identified sequences publicly available in databases [96]. In addition to a new sialotranscriptome from *A. cajennense* female ticks fed on rabbits containing over 4,600 coding sequences, we briefly report a reanalysis of the sequences from Anatriello's [96] work (here designated LIBEST\_USP-RP dataset). We then compared these two datasets to reveal how our new sialotranscriptome enriched the pre-existing data.

The LIBEST\_USP-RP dataset originated from the salivary glands of *A. cajennense* female ticks fed on horses [76,96] and contains 1,147 ESTs. The dataset was reanalyzed with a previously described bioinformatic pipeline [36], and the annotated high-quality sequences clustered into 505 contigs with stringent similarity parameters (Additional file 4: AF4). Nearly 22% of the ESTs were annotated as putatively secreted and belonged to protein families consisting of proteases, protease inhibitors and cement-like proteins. However, the function of about half of these ESTs is unknown.

The new sialotranscriptome (Additional file 3: AF3, here designated the RNA-seq (454) dataset) contained many more sequences than the LIBEST\_019946 and LIBES-T\_USP-RP datasets. The two LIBEST datasets contain less than one megabase (MB) of content, whereas the RNA-seq (454) dataset produced over 69 MB, yielding approximately 180 times more sequences (Table 3). From the 4,604 CDSs annotated for RNA-seq (454), almost 30% (1,336) were shared with the other two LIBEST datasets (Figure 3a, see magenta, cyan and white overlaps). This ratio of shared sequences was calculated based on blastn searches using LIBEST\_019946 and LIBEST\_USP-RP as databases, and every match with an E-value less than  $10^{-10}$  was counted. In fact, the ratio of shared sequences may be lower because the counting was redundant, i.e., many CDSs from the new sialotranscriptome matched the same sequences from one or both LIBEST datasets. Because the CDSs were functionally annotated, we ascertained the main functional categories for each shared sequence. Whereas housekeeping proteins were most prevalent (67.9%) in the RNA-seq (454) and LIB-EST\_019946 overlap, secreted proteins were the majority of sequences (52.4%) shared by the RNA-seq (454) and LIBEST\_USPRP overlapping reads (Figure 3a, pie charts at the bottom). Thus, most transcripts shared by

Library	Tick life stage	Host	No. of sequences	Average size of sequence <sup>a</sup>	Number of Contigs	Total yield in MB <sup>b</sup>
LIBEST_019946 <sup>c</sup>	Adult fed females	Rabbits	1,769	472	1,234	0.79
LIBEST_USP-RP <sup>d,e</sup>	Adult fed females	Horses	1,147	493	505	0.57
RNA-seq (454) <sup>f</sup>	Adult fed females	Rabbits	180,857	384	4,604	69.4

Table 3 General characteristics of cDNA libraries from salivary glands of *Amblyomma cajennense* produced in different conditions

<sup>a</sup>size in bases; <sup>b</sup>size in mega (10<sup>6</sup>) bases; <sup>c</sup>described by Batista *et al.*[58]; <sup>d</sup>produced by Anatriello *et al.* [96] and described herein; <sup>e</sup>for details see Additional file 4: AF4. <sup>f</sup>Shown values were calculated considering the two Roche 454 libraries described herein (for details regarding contigs see Additional file 3: AF3).

the datasets 454-RNA-seq and LIBEST\_USPRP potentially compose the saliva of *A. cajennense*.

In addition to its 70% novel sequences (Additional file 3: AF3), the new sialotranscriptome supplements the existing sequences for this tick species. For example, we performed protein sequence alignments using the translated nucleotide sequences from the three datasets and two important protein families of protease inhibitors. The CDS Ac-17015 was classified as a putative BPTI/Kunitz-type serine protease inhibitor, which contains five in tandem Kunitz (KU) domains within its 445 amino acids (aa). A five-KU domain protein, Penthalaris, is a tick anticoagulant found in the salivary glands of the Lyme disease vector, Ixodes scapularis [98]. Not only was Ac-17015 very similar to the aligned region of sequence ACGLSP02\_C10 (533 nt, 177 aa; 89.2% identity and 91.4% similarity) from the LIBEST\_USPRP dataset, but it also complemented the existing database record with the full-length sequence containing both the N- and C-termini of the putative protein (Figure 3B). Although only one KU domain was detected in the incomplete sequence of the serine protease inhibitor in the LIBEST\_USPRP dataset, this putative protein could be considered a Pentalaris-type protein because it presented five KU domains.

Another secreted protease inhibitor addressed by sequence alignment was a member of the Cystatins, a single domain protein family of cysteine protease inhibitors in vertebrates, invertebrates and plants that affect a broad range of biological process [99]. Tick cystatins have been observed in the midgut and/or salivary glands and are most likely important during feeding and blood digestion [100]. The CDS Ac-103849 was classified as a secreted cystatin (136 aa) and was very similar to the aligned region of the ACAH15A10 sequence (505 nt, 167 aa; 59.5% identity and 73.5% similarity) from the LIBEST\_019946 dataset (Figure 3c). In addition to the putative 5' untranslated region, the translated nucleotide sequence of ACAH15A10 had a stop codon (not shown in the alignment) at 10 residues before the initial methionine, most likely due to sequencing error. This is an example of how the new A. cajennense sialotranscriptome will complete the existing expression data from salivary glands of this tick. However, we must mention that the two LIBEST datasets also contain important unique sequences produced by biological variations at the tick-host interface, such as tick population, host species and time of blood feeding.

## Phylogenetic analyses of salivary coding sequences with potential immunomodulatory functions

The three Amblyomma sialotranscriptomes contained the cysteine protease inhibitors called thyropins, which contain thyroglobulin type-1 (Thyr-1) domains and inhibit either cysteine or cation-dependent proteinases [101]. During tick feeding, antigens are exposed to hosts and enter the endocytic pathway of antigen presenting cells (APC), where they are processed and loaded onto the major histocompatibility complex (MHC) class II. Proper folding and trafficking of MHCII is associated with the invariant chain (Ii) molecule, whose Thyr-1 domain-containing isoforms regulate endosomal cathepsins, cysteine proteases implicated in antigen processing [102]. The significant homology between Ii Thyr-1 domains and thyropins suggests that this protein family controls host immune factors. The A. triste, A. parvum and A. cajennense sialotranscriptomes contained 11 fulllength thyropin sequences that were compared with other 15 thyropin sequences from A. maculatum, A. variegatum, Ixodes ricinus, Rhipicephalus pulchellus, R. sanguineus and Ornithodoros moubata ticks (Figure 4). The phylogram showed the divergence between hard and soft ticks; the O. moubata thyropin sequence (Om-2539) presented low similarity with proteins from hard ticks and clustered as an external group. Two robust super clades (defined by related proteins belonging to clades with > 86% bootstrap support) were composed of three polyspecific clades (with > 72% bootstrap support). Most Amblyomma thyropins are more closely related to each other, and some species-specific thyropin sequences were very similar (e.g., Ap-10274, Ap-10275 and Ap-10276 on clade IV), which could represent alleles from the same gene or products of gene duplication. Thyropin sequences found in A. triste and A. cajennense clustered in clade I and presented higher similarities with Rhipicephalus sequences than with proteins in clade IV, which is composed of sequences from all three Amblyomma sialotranscriptomes. We observed two consistent subclusters in clade I and IV, formed by A. triste and A. maculatum sequences, which corroborated the close relationship between these tick species previously described by Estrada-



**Figure 3 Comparison between distinct sialotranscriptome of Amblyomma cajennese tick. a)** Venn diagram showing the proportion of CDSs (4,604 annotated in Additional file 3: AF3) identified in the RNA-seq (454) dataset and shared with the LIBEST\_019946 (Batista et al., 2008) and LIBEST\_USP-RP datasets. The values were based on the number of CDSs with significant matches to the other two datasets using the blastn algorithm. The magenta, cyan and white overlaps represent the sequences shared between the datasets. Approximately 30% of the CDSs (1,336) were contained sequences shared between two or more libraries. The proportion of shared proteins putatively secreted in the saliva, with housekeeping or unknown functions, are displayed in the small gray pie charts at the bottom. b) Sequence alignment of CDS Ac-1705 and ACGLSP02\_C10 of the LIBEST\_USP-RP dataset. This CDS was annotated as a putative Pentalaris protein, a serine protease inhibitor belonging to Kunitz domain superfamily. Horizontal brackets indicate the Kunitz domains (KU, 53 amino acids) along the sequences (I to V), as identified by the SMART protein database (smart00131). Dashed brackets indicate SMART hits with low scores, whereas solid brackets indicate SMART hits with high similarity scores. c) Sequence alignment of CDS Ac-103849 and ACAH15A10 of the LIBEST\_019946 dataset. This CDS was annotated as a secreted Cystatin, a cysteine protease inhibitors. Horizontal brackets indicate the Cystatin-like domain (CY, 107 amino acids) identified by the SMART protein database (smart00043). In *b* and *c*, upper arrows indicate the peptide cleavage sites predicted by SignalP Server [97]. Side numbers on block alignments indicate the amino acid position in each sequence. The alignments were produced with Clustal-W using a PAM 250 matrix for amino acid coloring in BioEdit software.

Peña et al. [103] through 16S rDNA analysis. The divergence between Amblyomma thyropins in clades I and IV is reflected in features such as potential glycosylation sites. A detailed analysis of A. cajennense thyropin sequences (performed by the NetOGlyc 4.0 Server) demonstrated that CDS Ac-36285 has one and Ac-103065 has no potential glycosylation sites, while Ac-50867 has 32 potential glycosylation sites (Additional file 3: AF3, column AD). These differences in glycosylation might impact protein functions. Notably, A. maculatum, A. variegatum and R. *microplus* thyropins possess potential galactosylation sites, suggesting Thyr-1 and mucin domains [43].

Immune modulation is an important requirement for tick feeding because contact with hosts leads to the recognition of tick antigens by immune cells and subsequent activation of both cellular and humoral responses. Suppression of the immune system of hosts facilitates feeding and creates a convenient microenvironment for enhanced pathogen transmission [104]. Ticks develop characteristics to overcome barriers imposed by distinct hosts during their life cycle, and divergent DAP-36 immunosuppressant proteins found in heteroxenous ticks have a general role in immune modulation [105]. This family of proteins has been identified in tick sialotranscriptomes [39,43,105] and as immunosuppressants in Dermacentor andersoni salivary glands [106]. Fifteen full-length DAP-36 sequences were employed in a phylogenetic analysis with sequences from other hard ticks





downloaded from NCBI protein database, including A. maculatum, A. variegatum, Ixodes ricinus and Dermacentor andersoni (Figure 5). Although the overall analysis showed divergence within the hard ticks, conserved homologues between A. triste and A. cajennense were shown by the super clade (clade I and II). Notably, clade VIII indicated an earlier deviation of I. ricinus DAP-36 proteins from other members of the Ixodidae family. As with thyropins, ticks may have developed DAP-36 immunosuppressant proteins through gene duplication, and members of the same clade might have similar functions. Interestingly, CDS Ap-15676 showed little similarity to other Amblyomma sequences and was clustered as an external group, suggesting that DAP-36 proteins are divergent within the Amblyomma genus. Based on sequences that are >20% divergent at the amino acid level, three putative genes in A. triste and A. parvum and two

## Conclusions

We generated the first detailed transcriptional profile of salivary glands from two important hard ticks from South America, *A. parvum* and *A. triste*, which are reported reservoirs and potential vectors of pathogenic agents such as *Rickettsia*. Additionally, we obtained a larger and more accurate sialotranscriptome of the vector of Brazilian spotted fever, the *A. cajennense* tick. This work extensively supplements the current biological knowledge of the *Amblyomma* genus through the tens of thousands of sequences deposited and freely available for further characterization. Transcripts encoding the protein families important for tick-host-pathogen interactions were common among the libraries and shared



similarities with other heteroxenous hard ticks. Thus, these components are good targets for controlling tick infestations and, consequently, decreasing the incidence of tick-borne diseases.

## **Additional files**

Additional file 1: AF1. Annotated sialotranscriptome of the *Amblyomma triste tick* (xlsx), available at http://exon.niaid.nih.gov/ transcriptome/sm/AF1-Atris-web.xlsx

Additional file 2: AF2. Annotated sialotranscriptome of the *Amblyomma parvum tick* (xlsx), available at http://exon.niaid.nih.gov/ transcriptome/sm/AF2-Aparv-web.xlsx

Additional file 3: AF3. Annotated sialotranscriptome of the *Amblyomma cajennense* ticks, available at http://exon.niaid.nih.gov/ transcriptome/sm/AF3-Acaj-web.xlsx

Additional file 4: AF4. Functional classification of the LIBEST\_USP-RP dataset.

Additional file 5: AF5. Accession numbers of the public sequences from other ticks used in phylogenetic analyses.

### Abbreviations

aa: Amino acids; APC: Antigen presenting cells; cDNA: Complementary deoxyribonucleic acid; CDS(s): Coding sequence(s); ESTs: Expressed sequence tags; GRP: Glycine rich proteins; H: Housekeeping; HBP: Histamine binding protein; HCL: Hierarchical clustering; KU: Kunitz domains; LP: *Leishmania*-related proteins; MHC: Major histocompatibility complex; NGP1: *Amblyomma triste* nymphs fed on naïve guinea pigs; NGP2: *Amblyomma triste* nymphs fed on guinea pigs; NGP2: *Amblyomma triste* nymphs fed on dogs; FD1: *Amblyomma triste* females fed on acive dogs; FD2: *Amblyomma triste* females fed on dogs undergoing a second infestation; FD: Female ticks fed on rabbits; NSG: Next generation sequencing; nt: Nucleotide; PBS: Phosphate buffered saline; SG(s): Salivary gland(s); TE: Transposable elements; Thyr-1: Thyroglobulin type-1; TIL: Trypsin inhibitor-like domain; U: Unknown; US: Unknown secreted; V: Viral; S: Secreted.

### Competing interests

The authors declare that they have no competing interests.

### Authors' contributions

IKFMS, CM, MPJZ, SRM and BRF conceived and designed the experiments; MMM, HNSM and EA performed the tick infestations. GRG, SRM, EA, MMM and HNSM performed the experiments; GRG, LGG, JMCR, EA and SRM analyzed the data. GRG, LGG, IKFMS and SRM wrote the paper. All authors read and approved the final manuscript.

### Authors' information

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#### References

- Ribeiro JM: Blood-feeding arthropods: live syringes or invertebrate pharmacologists? Infect Agents Dis 1995, 4:143–152.
- Fontaine A, Diouf I, Bakkali N, Missé D, Pagès F, Fusai T, Rogier C, Almeras L: Implication of haematophagous arthropod salivary proteins in hostvector interactions. *Parasit Vectors* 2011, 4:187.
- Stibrániová I, Lahová M, Bartíková P: Immunomodulators in tick saliva and their benefits. Acta Virol 2013, 57:200–216.
- Nava S, Guglielmone AA, Mangold AJ: An overview of systematics and evolution of ticks. Front Biosci (Landmark Ed) 2009, 14:2857–2877.
- Martins TF, Onofrio VC, Barros-Battesti DM, Labruna MB: Nymphs of the genus Amblyomma (Acari: Ixodidae) of Brazil: descriptions, redescriptions, and identification key. *Ticks Tick-Borne Dis* 2010, 1:75–99.
- Szabó MPJ, Olegário MMM, Santos ALQ: Tick fauna from two locations in the Brazilian savannah. Exp Appl Acarol 2007, 43:73–84.
- Veronez VA, Freitas BZ, Olegário MMM, Carvalho WM, Pascoli GVT, Thorga K, Garcia MV, Szabó MPJ: Ticks (Acari: Ixodidae) within various phytophysiognomies of a Cerrado reserve in Uberlândia, Minas Gerais, Brazil. Exp Appl Acarol 2010, 50:169–179.
- Lopes CM, Leite RC, Labruna MB, de Oliveira PR, Borges LM, Rodrigues ZB, de Carvalho HA, de Freitas CM, Vieira Júnior CR: Host specificity of Amblyomma cajennense (Fabricius, 1787) (Acari: Ixodidae) with comments on the drop-off rhythm. *Mem Inst Oswaldo Cruz* 1998, 93:347–351.
- Rojas R, Marini MA, Zanatta Coutinho AT: Wild birds as hosts of Amblyomma cajennense (Fabricius, 1787) (Acari: Ixodidae). Mem Inst Oswaldo Cruz 1999, 94:315–322.
- Van der Heijden KM, Szabó MPJ, Egami MI, Pereira MC, Matushima ER: Histopathology of tick-bite lesions in naturally infested capybaras (Hydrochoerus hydrochaeris) in Brazil. Exp Appl Acarol 2005, 37:245–255.
- Szabó MPJ, Pinter A, Labruna MB: Ecology, biology and distribution of spotted-fever tick vectors in Brazil. Front Cell Infect Microbiol 2013, 3:27.
- Labruna MB: Ecology of rickettsia in South America. Ann N Y Acad Sci 2009, 1166:156–166.
- Beati L, Nava S, Burkman EJ, Barros-Battesti DM, Labruna MB, Guglielmone AA, Cáceres AG, Guzmán-Cornejo CM, León R, Durden LA, Faccini JLH: Amblyomma cajennense (Fabricius, 1787) (Acari: Ixodidae), the Cayenne tick: phylogeography and evidence for allopatric speciation. *BMC Evol Biol* 2013, 13:267.
- Nava S, Beati L, Labruna MB, Cáceres AG, Mangold AJ, Guglielmone AA: Reassessment of the taxonomic status of Amblyomma cajennense () with the description of three new species, Amblyomma tonelliae n. sp., Amblyomma interandinum n. sp. and Amblyomma patinoi n. sp., and reinstatement of Amblyomma mixtum, and Amblyomma sculptum (Ixodida: Ixodidae). *Ticks Tick-Borne Dis* 2014, 5:252–276.
- Pacheco RC, Moraes-Filho J, Nava S, Brandão PE, Richtzenhain LJ, Labruna MB: Detection of a novel spotted fever group rickettsia in Amblyomma parvum ticks (Acari: Ixodidae) from Argentina. *Exp Appl Acarol* 2007, 43:63–71.
- Olegario MM, Gerardi M, Tsuruta SA, Szabo MP: Life cycle of the tick Amblyomma parvum Aragao, 1908 (Acari: Ixodidae) and suitability of domestic hosts under laboratory conditions. *Vet Parasitol* 2011, 179:203–208.
- Tomassone L, Nuñez P, Gürtler RE, Ceballos LA, Orozco MM, Kitron UD, Farber M: Molecular detection of Ehrlichia chaffeensis in Amblyomma parvum ticks, Argentina. *Emerg Infect Dis* 2008, 14:1953–1955.

- Pacheco RC, Echaide IE, Alves RN, Beletti ME, Nava S, Labruna MB: Coxiella burnetii in ticks, Argentina. *Emerg Infect Dis* 2013, 19:344–346.
- Nava S, Szabó MPJ, Mangold AJ, Guglielmone AA: Distribution, hosts, 165 rDNA sequences and phylogenetic position of the Neotropical tick Amblyomma parvum (Acari: Ixodidae). Ann Trop Med Parasitol 2008, 102:409–425.
- 20. Nava S, Mangold AJ, Guglielmone AA: The natural hosts for larvae and nymphs of Amblyomma neumanni and Amblyomma parvum (Acari: Ixodidae). *Exp Appl Acarol* 2006, **40**:123–131.
- Szabó MPJ, Labruna MB, Pereira MC, Duarte JMB: Ticks (Acari: Ixodidae) on wild marsh-deer (Blastocerus dichotomus) from Southeast Brazil: infestations before and after habitat loss. J Med Entomol 2003, 40:268–274.
- Szabó MPJ, Castro MB, Ramos HGC, Garcia MV, Castagnolli KC, Pinter A, Veronez VA, Magalhães GM, Duarte JMB, Labruna MB: Species diversity and seasonality of free-living ticks (Acari: Ixodidae) in the natural habitat of wild Marsh deer (Blastocerus dichotomus) in Southeastern Brazil. Vet Parasitol 2007, 143:147–154.
- Venzal JM, Estrada-Peña A, Castro O, de Souza CG, Félix ML, Nava S, Guglielmone AA: Amblyomma triste Koch, 1844 (Acari: Ixodidae): hosts and seasonality of the vector of Rickettsia parkeri in Uruguay. Vet Parasitol 2008, 155:104–109.
- Nava S, Mangold AJ, Mastropaolo M, Venzal JM, Fracassi N, Guglielmone AA: Seasonal dynamics and hosts of Amblyomma triste (Acari: Ixodidae) in Argentina. Vet Parasitol 2011, 181:301–308.
- Venzal JM, Portillo A, Estrada-Peña A, Castro O, Cabrera PA, Oteo JA: Rickettsia parkeri in Amblyomma triste from Uruguay. *Emerg Infect Dis* 2004, 10:1493–1495.
- 26. Pacheco RC, Venzal JM, Richtzenhain LJ, Labruna MB: Rickettsia parkeri in Uruguay. *Emerg Infect Dis* 2006, **12**:1804–1805.
- Silveira I, Pacheco RC, Szabó MPJ, Ramos HGC, Labruna MB: Rickettsia parkeri in Brazil. Emerg Infect Dis 2007, 13:1111–1113.
- Nava S, Elshenawy Y, Eremeeva ME, Sumner JW, Mastropaolo M, Paddock CD: Rickettsia parkeri in Argentina. Emerg Infect Dis 2008, 14:1894–1897.
- Cicuttin G, Nava S: Molecular identification of Rickettsia parkeri infecting Amblyomma triste ticks in an area of Argentina where cases of rickettsiosis were diagnosed. *Mem Inst Oswaldo Cruz* 2013, 108:123–125.
- 30. Francischetti IMB: Platelet aggregation inhibitors from hematophagous animals. *Toxicon* 2010, 56:1130–1144.
- Chmelar J, Calvo E, Pedra JHF, Francischetti IMB, Kotsyfakis M: Tick salivary secretion as a source of antihemostatics. J Proteomics 2012, 75:3842–3854.
- Bowman AS, Sauer JR: Tick salivary glands: function, physiology and future. Parasitology 2004, 129(Suppl):S67–S81.
- Francischetti IMB, Sa-Nunes A, Mans BJ, Santos IM, Ribeiro JMC: The role of saliva in tick feeding. Front Biosci (Landmark Ed) 2009, 14:2051–2088.
- Nuttall PA, Trimnell AR, Kazimirova M, Labuda M: Exposed and concealed antigens as vaccine targets for controlling ticks and tick-borne diseases. *Parasite Immunol* 2006, 28:155–163.
- Ribeiro JMC, Labruna MB, Mans BJ, Maruyama SR, Francischetti IMB, Barizon GC, de Miranda Santos IKF: The sialotranscriptome of Antricola delacruzi female ticks is compatible with non-hematophagous behavior and an alternative source of food. *Insect Biochem Mol Biol* 2012, 42:332–342.
- Ribeiro JM, Alarcon-Chaidez F, Francischetti IM, Mans BJ, Mather TN, Valenzuela JG, Wikel SK: An annotated catalog of salivary gland transcripts from Ixodes scapularis ticks. *Insect Biochem Mol Biol* 2006, 36:111–129.
- Chmelar J, Anderson JM, Mu J, Jochim RC, Valenzuela JG, Kopecký J: Insight into the sialome of the castor bean tick, Ixodes ricinus. *BMC Genomics* 2008, 9:233.
- Francischetti IMB, Anderson JM, Manoukis N, Pham VM, Ribeiro JMC: An insight into the sialotranscriptome and proteome of the coarse bontlegged tick, Hyalomma marginatum rufipes. J Proteomics 2011, 74:2892–2908.
- Anatriello E, Ribeiro JM, de Miranda-Santos IK, Brandao LG, Anderson JM, Valenzuela JG, Maruyama SR, Silva JS, Ferreira BR: An insight into the sialotranscriptome of the brown dog tick, Rhipicephalus sanguineus. BMC Genomics 2010, 11:450.
- Alarcon-Chaidez FJ, Sun J, Wikel SK: Transcriptome analysis of the salivary glands of Dermacentor andersoni Stiles (Acari: Ixodidae). Insect Biochem Mol Biol 2007, 37:48–71.

- Ribeiro JM, Anderson JM, Manoukis NC, Meng Z, Francischetti IM: A further insight into the sialome of the tropical bont tick, Amblyomma variegatum. BMC Genomics 2011, 12:136.
- Metzker ML: Sequencing technologies the next generation. Nat Rev Genet 2010, 11:31–46.
- 43. Karim S, Singh P, Ribeiro JM: A deep insight into the sialotranscriptome of the gulf coast tick, amblyomma maculatum. *PLoS One* 2011, 6:e28525.
- Schwarz A, von Reumont BM, Erhart J, Chagas AC, Ribeiro JMC, Kotsyfakis M: De novo lxodes ricinus salivary gland transcriptome analysis using two next-generation sequencing methodologies. *FASEB J* 2013, 27:4745–4756.
- 45. Iwanaga S, Isawa H, Yuda M: Horizontal gene transfer of a vertebrate vasodilatory hormone into ticks. *Nat Commun* 2014, **5:**3373.
- Labruna MB, Fugisaki EYM, Pinter A, Duarte JMB, Szabó MJP: Life cycle and host specificity of Amblyomma triste (Acari: Ixodidae) under laboratory conditions. *Exp Appl Acarol* 2003, 30:305–316.
- Nunes PH, Bechara GH, Camargo-Mathias MI: Morphological changes in the salivary glands of Amblyomma cajennense females (Acari: Ixodidae) in different feeding stages on rabbits at first infestation. *Exp Appl Acarol* 2008, 45:199–209.
- Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ: Basic local alignment search tool. J Mol Biol 1990, 215:403–410.
- Ashburner M, Ball CA, Blake JA, Botstein D, Butler H, Cherry JM, Davis AP, Dolinski K, Dwight SS, Eppig JT, Harris MA, Hill DP, Issel-Tarver L, Kasarskis A, Lewis S, Matese JC, Richardson JE, Ringwald M, Rubin GM, Sherlock G: Gene ontology: tool for the unification of biology, The Gene Ontology Consortium. Nat Genet 2000, 25:25–29.
- Bairoch A, Boeckmann B: The SWISS-PROT protein sequence data bank, recent developments. Nucleic Acids Res 1993, 21:3093–3096.
- Marchler-Bauer A, Panchenko AR, Shoemaker BA, Thiessen PA, Geer LY, Bryant SH: CDD: a database of conserved domain alignments with links to domain three-dimensional structure. *Nucleic Acids Res* 2002, 30:281–283.
- Schultz J, Copley RR, Doerks T, Ponting CP, Bork P: SMART: a web-based tool for the study of genetically mobile domains. *Nucleic Acids Res* 2000, 28:231–234.
- Tatusov RL, Fedorova ND, Jackson JD, Jacobs AR, Kiryutin B, Koonin EV, Krylov DM, Mazumder R, Mekhedov SL, Nikolskaya AN, Rao BS, Smirnov S, Sverdlov AV, Vasudevan S, Wolf YI, Yin JJ, Natale DA: The COG database: an updated version includes eukaryotes. *BMC Bioinformatics* 2003, 4:41.
- Bateman A, Birney E, Durbin R, Eddy SR, Howe KL, Sonnhammer EL: The Pfam protein families database. *Nucleic Acids Res* 2000, 28:263–266.
- Morgan M, Anders S, Lawrence M, Aboyoun P, Pagès H, Gentleman R: ShortRead: a bioconductor package for input, quality assessment and exploration of high-throughput sequence data. *Bioinformatics* 2009, 25:2607–2608.
- Eisen MB, Spellman PT, Brown PO, Botstein D: Cluster analysis and display of genome-wide expression patterns. *Proc Natl Acad Sci U S A* 1998, 95:14863–14868.
- Saeed AI, Sharov V, White J, Li J, Liang W, Bhagabati N, Braisted J, Klapa M, Currier T, Thiagarajan M, Sturn A, Snuffin M, Rezantsev A, Popov D, Ryltsov A, Kostukovich E, Borisovsky I, Liu Z, Vinsavich A, Trush V, Quackenbush J: TM4: a free, open-source system for microarray data management and analysis. *Biotechniques* 2003, 34:374–378.
- Batista IF, Chudzinski-Tavassi AM, Faria F, Simons SM, Barros-Batestti DM, Labruna MB, Leao LI, Ho PL, Junqueira-de-Azevedo IL: Expressed sequence tags (ESTs) from the salivary glands of the tick Amblyomma cajennense (Acari: Ixodidae). *Toxicon* 2008, 51:823–834.
- Boguski MS, Lowe TM, Tolstoshev CM: dbEST-database for "expressed sequence tags". Nat Genet 1993, 4:332–333.
- 60. dbEST: Expressed Sequence Tags Database. [http://www.ncbi.nlm.nih.gov/ dbEST/]
- 61. ORF Finder: A web Based Interactive Tool for Predicting and Analysing ORFs from Nucleotide Sequences. [http://www.ncbi.nlm.nih.gov/gorf/gorf.html]
- Thompson JD, Higgins DG, Gibson TJ: CLUSTAL W: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice. Nucleic Acids Res 1994, 22:4673–4680.
- 63. BioEdit: Biological Sequence Alignment Editor for Windows 95/98/NT/2K/XP/7. [http://www.mbio.ncsu.edu/BioEdit/bioedit.html]
- 64. Tamura K, Peterson D, Peterson N, Stecher G, Nei M, Kumar S: MEGA5: molecular evolutionary genetics analysis using maximum likelihood,

evolutionary distance, and maximum parsimony methods. *Mol Biol Evol* 2011, **28**:2731–2739.

- Jones DT, Taylor WR, Thornton JM: The rapid generation of mutation data matrices from protein sequences. *Comput Appl Biosci* 1992, 8:275–282.
- Dantas-Torres F: Ticks as vectors of Leishmania parasites. Trends Parasitol 2011, 27:155–159.
- Parola P, Paddock CD, Socolovschi C, Labruna MB, Mediannikov O, Kernif T, Abdad MY, Stenos J, Bitam I, Fournier P-E, Raoult D: Update on tick-borne rickettsioses around the world: a geographic approach. *Clin Microbiol Rev* 2013, 26:657–702.
- Francischetti IM, Mather TN, Ribeiro JM: Cloning of a salivary gland metalloprotease and characterization of gelatinase and fibrin (ogen) lytic activities in the saliva of the Lyme disease tick vector lxodes scapularis. *Biochem Biophys Res Commun* 2003, 305:869–875.
- Turner AJ, Isaac RE, Coates D: The neprilysin (NEP) family of zinc metalloendopeptidases: genomics and function. *Bioessays* 2001, 23:261–269.
- Anguita J, Ramamoorthi N, Hovius JWR, Das S, Thomas V, Persinski R, Conze D, Askenase PW, Rincón M, Kantor FS, Fikrig E: Salp15, an ixodes scapularis salivary protein, inhibits CD4(+) T cell activation. *Immunity* 2002, 16:849–859.
- Hojgaard A, Biketov SF, Shtannikov AV, Zeidner NS, Piesman J: Molecular identification of Salp15, a key salivary gland protein in the transmission of lyme disease spirochetes, from Ixodes persulcatus and Ixodes pacificus (Acari: Ixodidae). J Med Entomol 2009, 46:1458–1463.
- Lai R, Takeuchi H, Lomas LO, Jonczy J, Rigden DJ, Rees HH, Turner PC: A new type of antimicrobial protein with multiple histidines from the hard tick, Amblyomma hebraeum. *FASEB J* 2004, 18:1447–1449.
- Fogaça AC, Almeida IC, Eberlin MN, Tanaka AS, Bulet P, Daffre S: Ixodidin, a novel antimicrobial peptide from the hemocytes of the cattle tick Boophilus microplus with inhibitory activity against serine proteinases. *Peptides* 2006, 27:667–674.
- Chrudimská T, Slaninová J, Rudenko N, Růžek D, Grubhoffer L: Functional characterization of two defensin isoforms of the hard tick Ixodes ricinus. *Parasit Vectors* 2011, 4:63.
- Frauenschuh A, Power CA, Déruaz M, Ferreira BR, Silva JS, Teixeira MM, Dias JM, Martin T, Wells TNC, Proudfoot AEI: Molecular cloning and characterization of a highly selective chemokine-binding protein from the tick Rhipicephalus sanguineus. J Biol Chem 2007, 282:27250–27258.
- 76. Maruyama SR, Anatriello E, Anderson JM, Ribeiro JM, Brandao LG, Valenzuela JG, Ferreira BR, Garcia GR, Szabo MP, Patel S, Bishop R, de Miranda-Santos IK: The expression of genes coding for distinct types of glycine-rich proteins varies according to the biology of three metastriate ticks, Rhipicephalus (Boophilus) microplus, Rhipicephalus sanguineus and Amblyomma cajennense. *BMC Genomics* 2010, 11:363.
- Andersen JF, Pham VM, Meng Z, Champagne DE, Ribeiro JMC: Insight into the sialome of the Black Fly, Simulium vittatum. J Proteome Res 2009, 8:1474–1488.
- Campbell CL, Vandyke KA, Letchworth GJ, Drolet BS, Hanekamp T, Wilson WC: Midgut and salivary gland transcriptomes of the arbovirus vector Culicoides sonorensis (Diptera: Ceratopogonidae). Insect Mol Biol 2005, 14:121–136.
- Paesen GC, Siebold C, Dallas ML, Peers C, Harlos K, Nuttall PA, Nunn MA, Stuart DI, Esnouf RM: An ion-channel modulator from the saliva of the brown ear tick has a highly modified Kunitz/BPTI structure. J Mol Biol 2009, 389:734–747.
- Mason RG, Read MS: Some species differences in fibrinolysis and blood coagulation. J Biomed Mater Res 1971, 5:121–128.
- Lehane MJ: Peritrophic matrix structure and function. Annu Rev Entomol 1997, 42:525–550.
- Sangamnatdej S, Paesen GC, Slovak M, Nuttall PA: A high affinity serotonin- and histamine-binding lipocalin from tick saliva. Insect Mol Biol 2002, 11:79–86.
- Paesen GC, Adams PL, Harlos K, Nuttall PA, Stuart DI: Tick histaminebinding proteins: isolation, cloning, and three-dimensional structure. *Mol Cell* 1999, 3:661–671.
- 84. Lai R, Takeuchi H, Jonczy J, Rees HH, Turner PC: A thrombin inhibitor from the ixodid tick, Amblyomma hebraeum. *Gene* 2004, **342**:243–249.
- Koh CY, Kazimirova M, Trimnell A, Takac P, Labuda M, Nuttall PA, Kini RM: Variegin, a novel fast and tight binding thrombin inhibitor from the tropical bont tick. J Biol Chem 2007, 282:29101–29113.

- Mulenga A, Khumthong R, Blandon MA: Molecular and expression analysis of a family of the Amblyomma americanum tick Lospins. *J Exp Biol* 2007, 210(Pt 18):3188–3198.
- 87. Mulenga A, Kim T, Ibelli AMG: Amblyomma americanum tick saliva serine protease inhibitor 6 is a cross-class inhibitor of serine proteases and papain-like cysteine proteases that delays plasma clotting and inhibits platelet aggregation. *Insect Mol Biol* 2013, **22**:306–319.
- Xu G, Fang QQ, Keirans JE, Durden LA: Ferritin gene coding sequences Are conserved among eight hard tick species (ixodida: ixodidae). Ann Entomol Soc Am 2004, 97:567–573.
- Prevot P-P, Adam B, Boudjeltia KZ, Brossard M, Lins L, Cauchie P, Brasseur R, Vanhaeverbeek M, Vanhamme L, Godfroid E: Anti-hemostatic effects of a serpin from the saliva of the tick lxodes ricinus. J Biol Chem 2006, 281:26361–26369.
- Chmelar J, Oliveira CJ, Rezacova P, Francischetti IMB, Kovarova Z, Pejler G, Kopacek P, Ribeiro JMC, Mares M, Kopecky J, Kotsyfakis M: A tick salivary protein targets cathepsin G and chymase and inhibits host inflammation and platelet aggregation. *Blood* 2011, 117:736–744.
- Graça-Souza AV, Maya-Monteiro C, Paiva-Silva GO, Braz GRC, Paes MC, Sorgine MHF, Oliveira MF, Oliveira PL: Adaptations against heme toxicity in blood-feeding arthropods. Insect Biochem Mol Biol 2006, 36:322–335.
- Galay RL, Aung KM, Umemiya-Shirafuji R, Maeda H, Matsuo T, Kawaguchi H, Miyoshi N, Suzuki H, Xuan X, Mochizuki M, Fujisaki K, Tanaka T: Multiple ferritins are vital to successful blood feeding and reproduction of the hard tick Haemaphysalis longicornis. J Exp Biol 2013, 216(Pt 10):1905–1915.
- Hackman RH: Structure and function in tick cuticle. Annu Rev Entomol 1982, 27:75–95.
- Francischetti IMB, Mather TN, Ribeiro JMC: Tick saliva is a potent inhibitor of endothelial cell proliferation and angiogenesis. *Thromb Haemost* 2005, 94:167–174.
- Zhang H, Zhang W, Wang X, Zhou Y, Wang N, Zhou J: Identification of a cysteine-rich antimicrobial peptide from salivary glands of the tick Rhipicephalus haemaphysaloides. *Peptides* 2011, 32:441–446.
- Anatriello E, Ferreira BR, Brandão LG, Valenzuela JG, Ribeiro JM, Silva JS, de Miranda Santos IKF: Comparative analysis of transcriptomes of salivary glands from ticks, Amblyomma cajennese and Rhipicephalus sanguineus [abstract]. Vet Immunol Immunopathol 2009, 128:223.
- Petersen TN, Brunak S, von Heijne G, Nielsen H: SignalP 4.0: discriminating signal peptides from transmembrane regions. Nat Methods 2011, 8:785–786.
- Francischetti IMB, Mather TN, Ribeiro JMC: Penthalaris, a novel recombinant five-Kunitz tissue factor pathway inhibitor (TFPI) from the salivary gland of the tick vector of Lyme disease, Ixodes scapularis. *Thromb Haemost* 2004, 91:886–898.
- Abrahamson M, Alvarez-Fernandez M, Nathanson C-M: Cystatins. Biochem Soc Symp 2003, 70:179–199.
- 100. Schwarz A, Valdés JJ, Kotsyfakis M: The role of cystatins in tick physiology and blood feeding. *Ticks Tick-Borne Dis* 2012, **3:**117–127.
- Lenarcic B, Bevec T: Thyropins-new structurally related proteinase inhibitors. Biol Chem 1998, 379:105–111.
- 102. Zavasnik-Bergant V, Schweiger A, Bevec T, Golouh R, Turk V, Kos J: Inhibitory p41 isoform of invariant chain and its potential target enzymes cathepsins L and H in distinct populations of macrophages in human lymph nodes. *Immunology* 2004, 112:378–385.
- 103. Estrada-Peña A, Venzal JM, Mangold AJ, Cafrune MM, Guglielmone AA: The Amblyomma maculatum Koch, 1844 (Acari: Ixodidae: Amblyomminae) tick group: diagnostic characters, description of the larva of A. parvitarsum Neumann, 16S rDNA sequences, distribution and hosts. *Syst Parasitol* 1901, 2005(60):99–112.
- 104. Brossard M, Wikel SK: Tick immunobiology. Parasitology 2004, 129(Suppl): S161–S176.
- 105. Aljamali MN, Hern L, Kupfer D, Downard S, So S, Roe BA, Sauer JR, Essenberg RC: Transcriptome analysis of the salivary glands of the female tick Amblyomma americanum (Acari: Ixodidae). Insect Mol Biol 2009, 18:129–154.
- Bergman DK, Ramachandra RN, Wikel SK: Characterization of an immunosuppressant protein from Dermacentor andersoni (Acari: Ixodidae) salivary glands. J Med Entomol 1998, 35:505–509.

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