Towards a Semen Proteome of the Dengue Vector Mosquito: Protein Identification and Potential Functions

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Abstract

Background: No commercially licensed vaccine or treatment is available for dengue fever, a potentially lethal infection that impacts millions of lives annually. New tools that target mosquito control may reduce vector populations and break the cycle of dengue transmission. Male mosquito seminal fluid proteins (Sfps) are one such target since these proteins, in aggregate, modulate the reproduction and feeding patterns of the dengue vector, Aedes aegypti. As an initial step in identifying new targets for dengue vector control, we sought to identify the suite of proteins that comprise the Ae. aegypti ejaculate and determine which are transferred to females during mating.

Methodology and Principal Findings: Using a stable-isotope labeling method coupled with proteomics to distinguish male- and female-derived proteins, we identified Sfps and sperm proteins transferred from males to females. Sfps were distinguished from sperm proteins by comparing the transferred proteins to sperm-enriched samples derived from testes and seminal vesicles. We identified 93 male-derived Sfps and 52 predicted sperm proteins that are transferred to females during mating. The Sfp protein classes we detected suggest roles in protein activation/inactivation, sperm utilization, and ecdysteroidogenesis. We also discovered that several predicted membrane-bound and intracellular proteins are transferred to females in the seminal fluids, supporting the hypothesis that Ae. aegypti Sfps are released from the accessory gland cells through apocrine secretion, as occurs in mammals. Many of the Ae. aegypti predicted sperm proteins were homologous to Drosophila melanogaster sperm proteins, suggesting conservation of their sperm-related function across Diptera.

Conclusion and Significance: This is the first study to directly identify Sfps transferred from male Ae. aegypti to females. Our data lay the groundwork for future functional analyses to identify individual seminal proteins that may trigger female post-mating changes (e.g., in feeding patterns and egg production). Therefore, identification of these proteins may lead to new approaches for manipulating the reproductive output and vectorial capacity of Ae. aegypti.

Introduction

Male seminal fluid proteins (Sfps) influence female reproductive and feeding behaviors in a range of insects studied to date (reviewed in [1],[2]). Therefore, these proteins may provide targets or pathways that can be manipulated to reduce pathogen transmission by blood-feeding arthropods. The Aedes aegypti mosquito transmits several pathogens of concern to human health, including the viruses that cause dengue and dengue hemorrhagic fever (DHF) [3]. Dengue, the most important mosquito-borne virus impacting human health, is a re-emerging disease in the tropical regions of the world. There is currently no vaccine against, or cure for, dengue, although research in this area is ongoing ([4–6]). Therefore, prevention of dengue infection depends heavily on control of its mosquito vector.

Understanding mosquito reproductive biology is critical to developing effective vector control methods. Previous research on Ae. aegypti suggests that mating and, specifically, male-derived proteins may play an important role in modulating female reproduction and feeding behavior. Upon mating, female Ae. aegypti undergo a series of time-dependent behavioral and physiological changes. Relative to virgin females, mated females show physiological changes. Relative to virgin females, mated females
Author Summary

Dengue is a potentially lethal infection that impacts millions of humans annually. This disease is caused by flaviviruses transmitted by infected female *Aedes aegypti* mosquitoes during blood feeding. No commercial vaccine or treatment is available for dengue infection. One way to break the disease transmission cycle is to develop new tools to reduce dengue vector populations. Seminal fluid proteins (Sfps) produced in the reproductive glands of male mosquitoes and transferred to females in the ejaculate during mating could be the target of such a tool. In related insects, Sfps modulate female reproduction and feeding patterns. Here we report 145 proteins that are transferred to females in the *Ae. aegypti* ejaculate. The proteins, which include Sfps and sperm proteins, fall into biochemical classes that suggest important potential roles in mated females. Of particular interest are proteins that could play roles in fertility and hormonal activity (including pathways involved in egg development and utilization of the blood meal). Our results lay important groundwork for new control strategies by identifying candidate proteins that may alter the reproductive biology or blood-feeding patterns of female *Ae. aegypti* and ultimately reduce the global burden of dengue.

We adapted this method to blood-feeding mosquitoes and discovered a set of proteins transferred from male to female *Ae. aegypti*. Among the Sfps we identified are potential modulators of protein activation/inactivation, sperm utilization, and ecdysteroidogenesis. Few of the *Ae. aegypti* Sfps we detected are homologs of known or predicted Sfps in other insect species, although many are in protein classes that are conserved across seminal fluid of a wide range of taxa ([37],[39]). Furthermore, our finding of intracellular and membrane-bound proteins in the transferred Sfps supports the hypothesis that *Ae. aegypti* Sfps are secreted, at least in part, through apocrine processes (pinching off of the apical portion of the cells into vesicles containing Sfps; [39]) in the accessory glands.

In the process of identifying the Sfps, we also identified a subset of 52 putative *Ae. aegypti* sperm proteins. The *D. melanogaster* homologs of many of the predicted *Ae. aegypti* sperm proteins are also sperm proteins ([40]), suggesting conservation of sperm-related function across Diptera. Some of the proteins we have identified may be useful targets for control of *Ae. aegypti* and may be applicable to other mosquito vectors.

Methods

**15N-Labeling of female *Ae. aegypti* proteins**

Overview. In order to distinguish male- and female-derived proteins in the mated female reproductive tract, we adapted a stable-isotope labeling technique that had been developed for *D. melanogaster* ([36]). Stable-isotope labeling of proteins makes their peptides unidentifiable by standard mass spectrometry analysis. Therefore, we mated labeled females to unlabeled males in order to identify male-derived proteins in the female reproductive tract. The steps involved in this process (described in detail below) are: stable-isotope labeling of yeast as food for mosquito larvae, rearing of mosquitoes on labeled yeast, mating of labeled females to unlabeled males, extraction of proteins from mated female reproductive tracts, and mass spectrometric analysis of the extracted proteins.

**Yeast labeling.** *Saccharomyces cerevisiae* strain D273-10B was used for all experiments. Two batches of yeast were prepared: one labeled with 15N and one unlabeled. Isotopic labeling was performed following the methods of Findlay et al. [36] with adjustments for incorporating the label into mosquitoes. Briefly, yeast was grown to saturation (~16 h at 30°C) in 200 mL of minimal medium consisting of 20 g of glucose, 1.7 g of yeast nitrogen base without amino acids and either 5 g of 15N labeled ammonium sulfate (>99% 15N-enrichment; Cambridge Stable Isotopes, Andover MA, USA) or 5 g of unlabeled ammonium sulfate, in sterile water. The following day, 800 mL of additional medium was added and the yeast was grown for another 24 h. Yeast was harvested by centrifugation at 5,000 rpm at 4°C for 10 min, and the pellet was re-suspended in 30 mL of sterile water and centrifuged such that 15 mL of excess water was removed to yield a final volume of 15 mL of yeast slurry. The yeast slurry was stored at 4°C for less than two weeks before use.

**Mosquito rearing.** The Liverpool strain of *Aedes aegypti* L was used for all experiments. Initially, eggs were hatched under vacuum and 200 first instar larvae were transferred to shallow pans containing 1 L of sterile water. Approximately 1 mL of the yeast slurry was added to each larval rearing pan every 1–2 days until pupation occurred. Preliminary data showed that females from the 15N-labeled treatment required additional nutrition beyond the yeast slurry (as assessed by their inability to fly and, thus, mate). Therefore, we supplemented the rearing pans with 200 mL of inoculum of rearing-water from a previous cohort of larvac grown under the same treatment (i.e., unlabeled or 15N-
labeled yeast). The resulting larvae produced females that were able to fly and mated readily. Unlabeled males from the same mosquito strain were reared in pans of 200 larvae/1 L water and fed a 1:1 mixture of Brewer’s yeast and lactalbumin until pupation. All pupae were placed individually into vials and held for emergence. All adult mosquitoes were maintained on 20% sucrose solution on soaked cotton wicks. Reproductive tract samples were dissected from unlabeled and 15N-labeled virgin females to test the effectiveness of the labeling technique.

**Mating.** 15N-labeled virgin females (3–5 days after emergence) were individually introduced into a 5 L bucket container that contained 20–40 unlabeled males (4–6 days after emergence). If a successful mating event was observed, both the female and male were removed from the bucket at the termination of mating as the pair began to separate. After mating, each female was placed individually in a test-tube and stored on ice for dissection. Females that did not mate within approximately 5 min were removed from the male cage and discarded; a new female was introduced into the male cage for mating. Dissections of the 15N-labeled females mated to unlabeled males were conducted within 30 min of mating.

**Dissections.** Female lower reproductive tracts (i.e., spermathecae, bursa, common oviduct) were dissected in MOPS buffer (80 mM NaCl, 10 mM KCl, 1 mM CaCl2, 0.2 mM MgCl2, 10 mM MOPS) on ice and homogenized in 20 μl Dulbecco’s PBS (DPBS) with protease inhibitors (Roche Complete Protease Inhibitor Tablets, Indianapolis, IN). Reproductive tracts from 20 females were pooled for each biological replicate. Samples were centrifuged at 12,000 rpm for 30 min at 4°C. The supernatant was transferred to a separate tube, the pellet was resuspended in 20 μl Dulbecco’s PBS with protease inhibitors, and 20 μl of 2× SDS sample buffer (125 mM Tris–HCl pH 6.8, 20% glycerol, 4% SDS, 10% β-mercaptoethanol, 0.001% bromophenol blue) was added to each sample. The samples were boiled for 4 min and stored at −80°C.

**Identification of proteins in sperm derived from seminal vesicle and testes.** In order to distinguish Sfps from sperm proteins, we conducted proteomic analyses of sperm-enriched samples derived from the seminal vesicles or the testes. Seminal vesicles or testes from 40 males were dissected in DPBS with protease inhibitors, on ice. The tissues were then placed in a fresh tube containing the sperm was transferred to a microcentrifuge tube containing 500 μl DPBST (DPBS with 0.1% Tween-20). The tissues were then placed in a fresh tube containing the sperm was transferred to a microcentrifuge tube containing 500 μl DPBST (DPBS with 0.1% Tween-20). The samples were spun at 20,800 x g for 5 min at 4°C. The supernatant was discarded and the pellet was washed twice in DPBST. After the final wash, the pellet was resuspended in 2× SDS sample buffer. We considered the resulting samples as “sperm-enriched” as they contained not only sperm, but likely also some tissue, tissue secretions, and Sfps.

**Protein separation and identification.** Both the supernatant and the pellet samples were analyzed using two independent biological replicates of each sample type, except for the virgin females for which only one sample each was used for verification of our techniques. Protein separation and identification was conducted as previously described [35]. Briefly, proteins from each sample were separated by electrophoresis on a one-dimensional 5–15% gradient SDS polyacrylamide gel and visualized using Simply-Blue SafeStain (Invitrogen, Carlsbad, CA), The Cornell University Life Sciences Core Proteomics and Mass Spectrometry facility conducted in-gel digestion, tryptic peptide extractions, and Nano-LC-MS/MS. The nanoLC was carried out using an LC Packings Ultimate integrated capillary HPLC system equipped with a Switchos valve switching unit (Dionex, Sunnyvale, CA), which was connected in-line to a hybrid triple quadrupole linear ion trap mass spectrometer, 4000 Q Trap (ABI/MD/SCIEX, Framingham, MA) equipped with Micro Ion Spray Head ion source. The resulting MS and MS/MS data were submitted for database searching using the MASCOT search engine version 2.3 (Matrix Science, Inc., Boston, MA) or ProteinPilot software 1.0 (Applied Biosystems, Foster City, CA) against three databases (see “Databases” section below): the Ae. aegypti predicted peptide (AaegL1.2 Gene Build; hereafter “Vectorbase”) database (including supplemental peptides from AaegL1.1 Gene Build, http://aegypti.vectorbase.org/index.php; MASCOT program, Matrix Science, Boston, MA), a 6-frame translation of the Aedes genome (version AaegL1.2; MASCOT program; hereafter “6-frame translation”), and a database of small (<150 amino acid) predicted peptides (ProteinPilot; hereafter “small peptides”). The default search settings used for protein identification were: one mis-cleavage for full tryptic with variable carbamidomethyl modification of cysteine, and a methionine oxidation. For the searches using MASCOT, the false discovery rate (FDR) was estimated for a measure of random identification from the same database. To estimate the FDR, an automatic decoy database search was performed in which a database of random sequences was generated and tested for raw spectra along with the real database.

Protein identifications were based on a significance threshold of <0.05. Additionally, we only considered proteins to be high confidence hits if either two different peptides from the same sample exceeded the significance threshold or if one peptide hit exceeded the significance threshold in two independent biological replicates (single or multiple peptide hits are reported in Tables S1 and S2). Hits from the 6-frame translation and the small peptides database were compared to the Vectorbase database using BLASTP, and queries with a significant match were removed. Any remaining hits from the 6-frame translation were searched for a predicted gene using the GeneID 1.2 prediction program (http://genome.crg.es/geneid.html) and only these hits were used in further analyses. We tested for predicted secretion signal sequences using SignalP 3.0 (www.cbs.dtu.dk/services/SignalP/) and for predicted protein domains using SMART (http://smart.embl-heidelberg.de/), and Pfam (http://pfam.sanger.ac.uk/). The relative quantitation of identified proteins in each biological sample was estimated using the exponentially modified protein abundance index (emPAI, [44]) and are reported in Tables S1 and S2.

**Databases.** All of the databases are derived from sequencing of the Liverpool strain. The Vectorbase database is based on 8× coverage and includes 4,758 supercontigs, 1.3 gigabases, 15,988 protein-encoding genes. Any remaining hits from the 6-frame translation were searched for a predicted gene using the GeneID 1.2 prediction program (http://genome.crg.es/geneid.html) and only these hits were used in further analyses. We tested for predicted secretion signal sequences using SignalP 3.0 (www.cbs.dtu.dk/services/SignalP/) and for predicted protein domains using SMART (http://smart.embl-heidelberg.de/), and Pfam (http://pfam.sanger.ac.uk/). The relative quantitation of identified proteins in each biological sample was estimated using the exponentially modified protein abundance index (emPAI, [44]) and are reported in Tables S1 and S2.

**Identification of homologs to Ae. aegypti Sfp and sperm protein-encoding genes.** By using BLASTP, we identified homologs of the Ae. aegypti proteins in the genomes of three other Diptera (Culex quinquefasciatus, Anopheles gambiae, and D. melanogaster)
based on amino acid similarity. The divergence time between *Aedes* and *Drosophila* is predicted to be 250 million years ago ([46]). The *Anopheles* and *Aedes* genera are predicted to have diverged approximately 226 million years ago ([47]). Divergence time for *Aedes* and *Caicus* is more recent than the divergence between *Aedes* and *Anopheles* ([47]). For the proteins identified from the *Ae. aegypti* Gene Build database, we defined homologs as reciprocal best BLASTP hits with ≥30% identity and E values ≤10^-3. For proteins identified from the Supplemental predicted peptides, the 6-frame translation and the small peptides databases, we used unidirectional BLASTP searches of the *Ae. aegypti* hits against the databases from the other three species.

Results and Discussion

Isotope labeling technique

In order to identify male-derived proteins that are transferred to females during mating, we used a whole-organism isotope labeling method. The principle of this method is to mate males to females whose proteins are labeled with the stable isotopes so as to exclude the female proteins from proteomic identification. Specifically, female proteins are labeled with 15N, which shifts their masses upward such that the masses of female-derived peptides do not match those expected in a standard search (uncorrected for 15N) of a predicted protein database. The method was developed by Krijgsveld et al. [48] and first used to identify SPFs by Findlay et al. [36] in *D. melanogaster*. We adapted this method to label female-derived proteins in *Ae. aegypti*. As with *D. melanogaster*, we reared larvae on yeast whose only nitrogen source was 15N. However, in order to generate females that could fly and mate, we had to supplement the larvae with an inoculum of rearing-water from larvae previously reared on 15N-labeled yeast (see Methods for more detail). To verify that the 15N-labeling technique sufficiently labeled female-derived proteins, we conducted nanoLC-MS/MS on protein samples from the reproductive tracts of labeled and unlabeled virgin females. We initially analyzed protein samples from two arbitrarily-chosen molecular weight ranges (~30 kD to 50 kD and ~98 kD to 120 kD) of both types of females. Using the Vectorbase database, we identified 115 proteins from the unlabeled female samples (Table S3) and no proteins from the labeled female samples. To further verify the labeling technique, we conducted nanoLC-MS/MS on the remaining gel sections from the labeled virgin female sample. We did not identify proteins from any of these samples. These results demonstrate that any proteins we identify in the reproductive tracts of labeled females mated to unlabeled males are highly likely to be male-derived. Furthermore, the technique we developed for *in vivo* stable-isotope labeling of *Ae. aegypti* proteins could be applied to other studies (e.g., to quantify proteomic changes in females in response to mating and/or blood-feeding or to distinguish mosquito-derived proteins from those of their pathogens, parasites, and/or endosymbionts; [49]).

Proteins transferred to females during mating

In our search against the Vectorbase database, we identified 128 proteins in the reproductive tracts of labeled females after mating with unlabeled males (Tables 1 and 2). Since the *Ae. aegypti* genome sequence is relatively new and thus the current annotation might still be missing actual genes, we also searched our mass spectrometry results against a 6-frame translation database of the *Ae. aegypti* contigs (Version AaegL1.2) and against a database of predicted small peptides (<150 amino acids). In our search against the 6-frame translation, we identified 12 novel predicted semen proteins (Tables 1 and 2). We identified 5 novel predicted small peptides from the small peptides database (Tables 1 and 2). The sequences of the unannotated predicted proteins and peptides are provided in Table S4. Thus, in total, we identified 145 male-derived predicted proteins that are transferred during mating to females. These proteins include SPFs and sperm proteins. For all searches, the FDR was ≤1%.

Nine of the identified proteins share identical amino acid sequence with other *Ae. aegypti* predicted proteins in the regions that were identified by our mass spectrometry analysis. As a result, we cannot distinguish amongst these proteins in our samples. For simplicity, we have listed just one protein from each of these pairs or groups of indistinguishable proteins in Tables 1 and 2. However, we list the identities of all proteins in these pairs or groups in Table S3.

Comparison to previously identified *Ae. aegypti* reproductive gland proteins

Of the 145 transferred proteins we identified using the whole-organism isotope labeling method, 123 are newly-recognized components of *Ae. aegypti* semen. The remaining 22 were previously identified as putative SPFs ([35]), and we demonstrate here that they are transferred to females during mating. Thirty-one additional proteins were identified as putative SPFs in our previous study ([35]), but we did not detect them as transferred in our current study. Those proteins may not be transferred to females, or may be transferred at quantities below our detection threshold or with post-translational modifications that render them unidentifiable by standard mass spectrometry. Of the 123 newly-recognized seminal proteins, 84 were previously identified from the reproductive glands of *Ae. aegypti*, however they were not designated as putative SPFs because they lacked predicted secretion signal sequences ([35]; L. Sirot, M. Wolfiner, L. Harrington, unpubl. data).

Distinguishing seminal fluid proteins from sperm proteins

In order to distinguish SPFs from sperm proteins among those transferred to females, we conducted a proteomics analysis of sperm-enriched samples from the seminal vesicles (SVs) and testes of virgin males. Sperm-enriched samples were obtained by releasing sperm from these organs, pelleting the sperm by centrifugation, and washing them repeatedly, as in Dorus et al. [40] (see Methods for details). We found 101 proteins that overlapped between our sperm-enriched samples from seminal vesicles and our sperm-enriched samples from testes. Of these 101 putative sperm proteins, 52 were detected as transferred to females during mating, providing a high-confidence subset of putative *Ae. aegypti* sperm proteins (Table 2).

Of the 145 total transferred proteins (see section “Proteins transferred to females during mating”), 16 were isolated from only one of the sperm-enriched tissues (SV; 5; testes; 11) and therefore were considered SV- or testes-derived SPFs, respectively, although we recognize that these could be sperm proteins. Additionally, 77 of the transferred proteins did not overlap with either sperm-enriched sample. Together, the 3 SV-derived SPFs, 11 testes-derived SPFs, and 77 of the 145 total transferred proteins that did not overlap with the sperm-enriched samples comprise a total of 93 proteins assigned with high-confidence as *Ae. aegypti* SPFs (Table 1).

Seminal fluid proteins

The *Ae. aegypti* SPFs identified represent a wide-range of predicted protein classes including proteolytic regulators, lectins, lipases, oxidoreductases, a cysteine-rich secretory protein (CRISP)
Table 1. Predicted seminal fluid proteins transferred in *Aedes aegypti* ejaculate.

| Molecular function | Predicted protein class | Aa$^a$ | Molecular function (cont.) | Predicted protein class | Aa $^a$ |
|--------------------|-------------------------|-------|---------------------------|-------------------------|--------|
| Binding            | Annexin                 | 11302 | Proteolysis/ Catalysis (cont.) | Protease               | 01588  |
|                    | Calcyphosine            | 08489 |                          |                         |        |
|                    | Dipeptidase             | 08893 |                          |                         |        |
|                    | Fibrinogen              | 01713 |                          |                         |        |
|                    | Kakapo                  | 02829 |                          |                         |        |
|                    | Lectin                  | 04679 |                          |                         |        |
|                    | Mitochondrial brown fat uncoupling protein | 07046 |                          |                         |        |
|                    | Moesin                  | 07915 |                          |                         |        |
|                    | Mucin                   | 00718 |                          |                         |        |
|                    | Odorant-binding         | AaegSfp1 |                          | Protease inhibitor | 02715  |
|                    | Phosphatidyethanol-binding protein | 11263 |                          |                         |        |
|                    | Tubulin β-chain         | 02848 |                          | Thioesterase            | 00479  |
|                    | None                    | 00479 |                          | Transferase             | 03746  |
|                    |                          | 08274 |                          |None                      | 02793  |
|                    |                          | 09201 |                          |                          | 13559  |
| Oxidoreductase     | Catalase                | 13407 |                          |                          | 17451  |
|                    | Decarboxylase           | 05790 |                          |                          | 17460  |
|                    | Dehydrogenase           | 04338 | Structural               | Actin                   | 01928  |
|                    |                          | 05308 |                          |                          | 05964  |
|                    |                          | 06928 |                          | Vitellogenin             | 05815  |
|                    |                          | 10464 |                          | None                     | 03348  |
|                    |                          | 12014 | Transport                | Cytochrome c oxidase subunit | 00929  |
|                    | NADH-ubiquinone oxidoreductase subunit | 05946 |                          |                          | 13751  |
|                    | Peroxidase              | 04112 |                          | Glutamate receptor       | 09813  |
| Proteolysis/ Catalysis | Aminopeptidase       | 02399 | Mitochondrial glutamate carrier | 11276  |
|                    |                          | 02978 | Sodium/calcium exchanger |                          | 12480  |
|                    |                          | 07201 | Other                    | Niemann-Pick Type C-2   | 09760  |
|                    | Asparaginase            | 02796 | Rab GDP-dissociation inhibitor |                        | 12904  |
|                    | ATP synthase subunit    | 06516 | Venom allergen            |                          | 09239  |
|                    |                          | 07777 | None                     |                          | 04944  |
|                    |                          | 11025 |                          |                          | 05219  |
|                    |                          | 12035 |                          |                          | 10824  |
|                    |                          | 12819 |                          |                          |        |
|                    |                          | 14053 |                          |                          |        |
|                    |                          | 14580 |                          |                          |        |
|                    | Glutathione transferase | 11741 |                          |                          |        |
Table 1. Cont.

| Molecular function | Predicted protein class | Aa* | Molecular function (cont.) | Predicted protein class | Aa |
|--------------------|-------------------------|-----|---------------------------|-------------------------|----|
| Hydrolyase         | 03666                   |     |                           |                         |    |
| Kinase             | 06485                   |     |                           |                         |    |
| Lipase             | 12359                   |     |                           |                         |    |
| Mannosidase        | 12731                   |     |                           |                         |    |
|                    | 07063                   |     |                           |                         |    |
|                    | 05763                   |     |                           |                         |    |

*5-digit numbers are the Vectorbase database identification numbers without the proceeding “AAEL”. Numbers with “Supp” prefix refer to proteins from the Supplementary predicted peptide database from AeagL1.1Gene Build. Numbers with the prefix “AaegSfp” refer to proteins from either the 6-frame translation or the small peptides databases. The amino acid sequences for all of the “Supp” and “AaegSfp” predicted proteins are given in Table S4.

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and a venom allergen, and fall into a variety of Gene Ontology predicted molecular function classes (Table 1; Fig 1A). Unlike Sfps in Drosophila ([31],[36],[50]) and An. gambiae ([33],[51]), in which some groups of Sfps tend to be spatially clustered, we found little evidence for spatial clustering of the 93 Sfp genes in the Ae. aegypti genome, with one exception. Four Sfp genes (AAEL006403; AAEL006414; AAEL006421; AAEL006429) clustered within a 23 kbp region on supercontig 1.204. One gene in this region (AAEL006430) encodes a protein that was not detected in the present study and may either not be transferred or may be transferred at a level that was undetectable by our methods. The proteins encoded by all five of the genes in this region have predicted trypsin domains and their shared amino acid sequence identities range from 36% to 64%.

As might be expected, the genes encoding the Sfps identified in this study tend to have highly male-biased expression when gene expression of whole males is compared to gene expression of whole females ([52]). Half (26 of 51) of the genes for which microarray data are available have significantly higher expression (at P<0.001) in males than in non-blood-fed females (O. Marinotti, pers. comm.), as compared to a genome average of 16% (Chi-square test; X² = 47.7; P<0.001; [52]). By comparison, 63% of the D. melanogaster Sfps identified by [36] have significantly higher expression (at P≤0.01) in males than in females as compared to a genome average of 12% (X² = 545.9; P<0.001; [53]). Surprisingly, transcript levels of two Ae. aegypti Sfp-encoding genes (encoding a predicted kinase, AAEL012359, and a predicted zinc metalloprotease, AAEL012217) are significantly higher in non-blood-fed females than in males.

Sequences comparisons to other Diptera. Table S1 shows the extent to which homologs of the 93 putative Sfps can be detected in three other Dipteran genomes (Cx. quinquens, An. gambiae, and D. melanogaster). In comparison to known or predicted Sfp- or male AG-encoding genes from An. gambiae or D. melanogaster, Ae. aegypti Sfp genes generally had little sequence similarity. None of the An. gambiae homologs to Ae. aegypti Sfps is among the previously-identified An. gambiae AG genes ([33],[51]) or mating plug protein-encoding genes ([33]). Only two D. melanogaster (CG31704, and CG5162) homologs to Ae. aegypti Sfps (AaegSfp3 and AAEL005815, respectively; range of percent identity: 41–59%) are among the known D. melanogaster Sfps or AG genes (Table S1; [31],[31],[36],[54]). Additionally, three Ae. aegypti Sfps (AAEL005815; AAEL004112; AAEL000718) share sequence similarity (range: 30–84% identity) with four of the 13 AG genes identified from the Mediterranean fruit fly, Ceratitis capitata (clones 11c, 18a, 30c, 33a; [55]).

Below, we discuss (i) the unannotated, newly-identified predicted proteins from the 6-frame translation and small peptides database, (ii) new insights into the mode of Ae. aegypti Sfp secretion from Vectorbase identified proteins, and (iii) the potential biological functions of a subset of Sfps in modulating reproductive and physiological processes within the mated female. In our previous report on Ae. aegypti Sfps ([35]), we discussed the potential roles of these proteins in a number of processes including protein folding, antimicrobial activity, and sperm utilization by females. Although the proteins we have identified in the current report include many in the same classes and predicted functions as our previous report (and include 22 of the same proteins), we have specifically chosen to highlight proteins that suggest functions in the mated female that were not discussed in our previous report ([35]). These protein classes are not necessarily the most highly represented of the predicted Sfps.

Unannotated seminal fluid proteins. Based on comparisons to the 6-frame translation and small peptides databases, we discovered 14 previously unannotated predicted Sfps (Table 1). Of the 9 predicted proteins from the 6-frame translation, only one (AaegSfp2) had a significant hit to the SMART or PFAM database. This protein includes a predicted secretion signal sequence and a Kazal-type serine protease inhibitor domain. Interestingly, another predicted secreted protein of the 9 hits (AaegSfp8) shared a high degree of sequence similarity (70% identity; E value = 2e-74) with one of the Vectorbase predicted proteins, AAEL010824. AAEL010824 and AaegSfp8 are located on the same contig within 34 Kb of each other. Two of the other hits from the 6-frame database also had predicted secretion signal sequences. Of the 5 hits to the small peptides database, we found one predicted secreted Kazal-type serine protease inhibitor (AaegSfp3), one predicted secreted odorant-binding protein (AaegSfp1), and three hits with no predicted secretion signal sequence and no predicted protein domains.

Mode of secretion. Sixty-two of the Sfps that we identified are predicted intracellular or membrane-bound proteins (e.g., ATPases, dipeptidyl peptidase, gamma glutamyl transpeptidase, glutathione S-transferase, angiotensin converting enzyme). Predicted intracellular proteins also have been reported in the seminal fluid of other organisms including bed bugs ([56]), honey bees ([57]), and humans ([58],[59]), and in the AGs and seminal fluids of D. melanogaster ([60]; G. Findlay & W. Swanson, pers. comm.). In bed bugs, Sfps of predicted intracellular origin have been considered potential cell contaminants ([56]); whereas, in honeybees, it has been suggested that these proteins may be
secreted through non-standard secretion routes ([57]). In *Ae. aegypti*, our finding of intracellular and membrane-bound Sfps is consistent with the hypothesized modes of secretion in the AGs of this species: based on cytological studies using light and electron microscopy, cells in the anterior portion of the glands are thought to secrete proteins by pinching off apically ("apocrine secretion"; [39],[61]), whereas cells in the posterior portion of the glands are thought to secrete proteins through granules and/or via rupture of the cell membrane ("holocrine secretion"; [61],[62]; but see [39]). Inside the female reproductive tract, the ejaculate contains "vacuoles" that grow over time after mating and then disappear by 24 h post-mating ([62]). Four of the Sfps we identified are subunits of the membrane-bound vacuolar-type proton ATPase and thus may be part of these vacuoles. Future studies could investigate whether the male-derived vacuolar ATPase is functional in the ejaculate vacuole membrane in the female reproductive tract and therefore may play a role in regulating the release of the contents of the vacuoles.

Importantly, our results emphasize that studies of Sfps in other species should not exclude proteins whose sequence (alone) suggest that they are intracellular and membrane-bound proteins. For species in which intracellular and/or membrane-bound proteins are found in the seminal fluid (e.g., bed bugs and honey bees), further research should be conducted on the mode of secretion of the male reproductive gland cells. Furthermore, secretion by *Ae. aegypti* accessory glands could serve as a model for secretion in systems such as the male reproductive glands of mammals, since mammalian prostasomes, exosomes, and epididymosomes contain many of the same classes as found among *Ae. aegypti* Sfps ([63]).

**Proteolysis.** Across a wide range of organisms, seminal fluid is rich in proteolysis regulators (e.g., [37],[38],[64–67]). *Ae. aegypti* is no exception. Thirteen of the 93 Sfps we identified in *Aedes aegypti* Table 2. Predicted sperm proteins transferred in *Aedes aegypti* ejaculate.

| Molecular function | Predicted protein class | Aa* | Molecular function (cont.) | Predicted protein class | Aa |
|--------------------|-------------------------|-----|---------------------------|-------------------------|----|
| **Binding**        | Aminopeptidase          | 06975 | Proteolysis/ Catalysis (cont.) | Protease inhibitor | AaegSp1 |
| Heat shock protein 70 | Supp4130 | None | | | 06509 |
| Histone            | 00490                   | | | | 10754 |
| Reticulocalbin     | 14589                   | Structural | Actin | | 01673 |
| Tubulin α-chain    | 06642                   | | | | 11197 |
| Myosin             | 13229                   | | | | 12543 |
| None               | 00637                   | Tubulin β-chain | 02851 | | 05052 |
| | 08779                   | | | | 04855 |
| | 10149                   | Transport | ADP, ATP carrier | | 04457 |
| | 10882                   | Cytochrome c | | | 05170 |
| | 14231                   | Cytochrome c oxidase subunit | | | 03675 |
| Oxido-reductase    | Dehydrogenase           | 00454 | Ubiquinol-cytochrome c reductase unit | | 03675 |
| | | 02881                   | | | 05269 |
| | | 03757                   | Voltage-dependent anion-selective channel | | 01872 |
| | | 08166                   | None | | 07195 |
| Proteolysis/Catalysis | Aconitase             | 03734 | None | | 09707 |
| | Aminopeptidase         | 00108                   | | | 12282 |
| | ATP synthase subunit   | 02827                   | | | 17096 |
| | | 05173                   | Supp4104 | | |
| | | 05610                   | Supp7141 | | |
| | | 05798                   | AaegSp2 | | |
| | | 08787                   | AaegSp3 | | |
| | | 08848                   | | | |
| | | 12175                   | | | |
| Kinase             | 06042                   | | | |
| Protease           | 03308                   | | | |

*5-digit numbers are the Vectorbase database identification numbers without the proceeding “AAEL0”. Numbers with “Supp” prefix refer to proteins from the Supplementary predicted peptide database from AaegL1.1 Gene Build. Numbers with the prefix “AaegSp” refer to proteins from either the 6-frame translation or the small peptides databases. The amino acid sequences for all of the “Supp” and “AaegSp” predicted proteins are given in Table S4.*

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ecdysteroids (ECDs) (e.g., [74]). ECDs are hormones that influence

sterol carriers are essential for the production of

Ae. aegypti putative male reproductive tracts and/or in mated females.

Sperm and sperm-associated proteins

From Ae. aegypti sperm-enriched tissue samples, we identified 101 sperm or sperm-associated proteins. Fifty-two were found among proteins transferred to females during mating (Table 2); the remaining 49 proteins were not detected as transferred (Table S6). These latter 49 proteins included 7 that have homologs found in the D. melanogaster sperm proteome ([40]). It is possible that some of these 49 Ae. aegypti proteins are components of somatic cells of the testis and/or SV tissues and play a role in spermatogenesis or sperm maintenance, whereas others could be sperm proteins whose abundance was too low for us to detect in the transferred samples or whose post-translational modifications rendered them unidentifiable by standard mass spectrometry. In the remainder of this section, we will only discuss the 52 proteins detected as transferred (hereafter referred to as “putative sperm proteins”).

Sequence comparisons to other Diptera. Table S2 shows the extent to which homologs of the 52 putative sperm proteins can be detected in three other Dipteran genomes (Cx. quinquefasciatus, An. gambiae, and D. melanogaster). Of the 30 proteins with D. melanogaster homologs, 17 (57%) of the homologs were found among the identified sperm proteins of D. melanogaster ([40]). This level of homology between Ae. aegypti putative sperm proteins and D. melanogaster sperm proteins suggests that the sperm-related functions of these proteins are conserved between the two species. Non-sperm D. melanogaster homologs of Ae. aegypti putative sperm proteins may also serve reproductive functions, as suggested by gene expression patterns. Of the 13 D. melanogaster homologs not found in the D. melanogaster sperm proteome (Table S2), transcripts of 5 are enriched in male and female reproductive tissues, transcripts of 1 are enriched only in male reproductive tissues, and transcripts of 6 are enriched only in female reproductive tissues ([73]).

Unannotated sperm proteins. We discovered three previously unannotated predicted sperm proteins from the 6-frame translation database (Table 2). One predicted protein from the 6-frame translation, AaegSp1, contained a predicted secretion signal sequence and a Kazal serine protease inhibitor domain. The other two hits contained no predicted secretion signal sequence and no conserved protein domains.

Potential functions. The likely biological functions of putative Ae. aegypti sperm proteins include spermatogenesis, serving as structural components of mature sperm, and sperm locomotion (Fig. 1B). Spermatogenesis-related proteins that we found among Ae. aegypti putative sperm proteins include the heat shock protein 70, actin, and tubulin (α- and β-chains). Ae. aegypti putative sperm proteins that might contribute to sperm structure or motility include actin, tubulins, dyneins, ATP synthases, protein kinases and kinesin motor proteins ([82–86]). Ae. aegypti putative sperm proteins that are predicted mitochondrial enzymes include malate dehydrogenase, aconitase, cytochrome c oxidase, and ubiquinol-cytochrome c reductase. In other animals, these proteins generate the energy necessary for sperm locomotion via oxidative phosphorylation and the citric acid cycle ([87–92]).

Steroidogenesis. One of the Sfps we identified (AAEL009760) is a predicted sterol carrier in the Niemann-Pick type C-2 (NPC2) family. In insects, sterol carriers are essential for the production of ecdysteroids (ECDs) (e.g., [74]). ECDs are hormones that influence molting, gametogenesis, vitellogenesis, and other reproductive processes (reviewed in [75],[76]). In Ae. aegypti, ECDs are essential components of a signaling cascade linking blood meal intake with vitellogenesis (e.g., [77]; reviewed in [78], [79]). Although a role for the male-derived NCP2-like protein has not yet been determined in Ae. aegypti, findings in other Dipteran species suggest that male AGs can not only synthesize ECD (An. gambiae, [80]) but also stimulate ECD production in females (D. melanogaster, [81]). We suggest that AAEL009760 could potentially contribute to the regulation of ECD biosynthesis in Ae. aegypti male reproductive tracts and/or in mated females.

Figure 1. Gene Ontology molecular function categories of Aedes aegypti seminal fluid and putative sperm proteins. A. Seminal fluid proteins; B. Putative sperm proteins. doi:10.1371/journal.pntd.0000989.g001

are predicted regulators of proteolysis. These include predicted trypsins, a zinc carboxypeptidase, a metalloprotease, a serine protease inhibitor (serpin), an angiotensin converting enzyme, and a proprotein convertase in the subtilisin/kexin type 4 (PCSK4) family (Tables 1 and S1). The predicted or known functions of seminal fluid proteolysis regulators include regulating the liquefaction of semen and/or mucus in the female reproductive tract ([68],[69]); protection of sperm from premature acrosome reactions ([67]); and activation and/or degradation of other reproductive proteins ([70]). We have previously discussed the potential role of predicted proteolysis regulators in Ae. aegypti seminal fluid ([35]). Here, we highlight the novel finding of a predicted proprotein convertase subtilisin/kexin type 4 (PCSK4) protein in insect seminal fluid.

PCSK4s can activate precursors of membrane receptors, peptide hormones, antibacterial peptides and neuropeptides through proteolytic processing ([71],[72]). Interestingly, the predicted PCSK4 (AAEL010725) found in this study shares close sequence similarity (56% identity; E-value: 2e-158) with the Ae. aegypti protein that processes vitellogenin (AAEL003652; [72]). To our knowledge, proprotein convertases have not been reported through proteolytic processing ([71],[72]). Interestingly, the predicted PCSK4 (AAEL010725) found in this study shares close sequence similarity (56% identity; E-value: 2e-158) with the Ae. aegypti protein that processes vitellogenin (AAEL003652; [72]). To our knowledge, proprotein convertases have not been reported previously in insect seminal fluid. Transcripts of the D. melanogaster gene encoding one protein in this class (CG10702) are enriched (relative to whole body) in the AGs ([73]). The protein they encode contains a predicted secretion signal sequence (SignalP; [41]), but it has not yet been detected in the seminal fluid in this species ([36]).
Summary and conclusion
Secretions of the reproductive glands of male *Ae. aegypti* have previously been shown to induce post-mating changes in female reproductive and feeding behavior ([22],[23],[26]). In order to lay the groundwork for identifying specific proteins causing these effects, we report here 145 male-derived proteins that are transferred to females during mating in *Ae. aegypti*. We distinguished 93 seminal fluid proteins from 52 predicted sperm proteins, thus contributing to the growing understanding of insect ejaculate proteomes ([33],[35],[36],[40],[57],[88],[93–95]). Twenty-two of these proteins were previously identified as male reproductive gland proteins ([35]), and we demonstrate here that they are transferred to the female.

The Sfps identified in this study suggest roles in protein activation/inactivation, ecdysteroidogenesis, and sperm utilization. Furthermore, our discovery that many predicted intracellular and membrane-bound proteins are transferred to females in the seminal fluid indicates that findings of such proteins in the seminal fluid of other species (e.g., [56],[57]) may also result from apocrine and/or holocrine secretion from the male reproductive glands ([39],[61]). The putative sperm proteins of *Ae. aegypti* show sequence conservation within Diptera and 17 of their *D. melanogaster* homologs are sperm proteins in that species ([40]) indicating potential conservation of sperm-related functions.

Genes encoding Sfps showed higher male-biased expression than the genome average. On the one hand, this is not unexpected because Sfps are made in the male reproductive tract and are then transferred to females. On the other hand, it is not necessarily predicted a priori that Sfp-encoding genes will be male-biased in their expression, and the way we identified the proteins was without bias regarding their genes’ expression. That 49% of the *Ae. aegypti* Sfp-encoding genes for which there are microarray data are not male-biased in expression will be important to bear in mind in designing future screens for Sfps.

Together, our results provide a foundation for functional analyses to associate individual Sfps with their function in the mated female. Once functions are identified for individual proteins, investigations of the pathways by which they induce effects on male and female reproductive biology could identify novel targets for control of *Ae. aegypti* and dengue transmission. Of particular interest is to determine how specific Sfps modulate female behavior and physiology (e.g., egg production and blood feeding) and to investigate candidate genes which increase the reproductive success of male *Ae. aegypti* that are to be used in genetic control strategies.

Supporting Information

**Table S1** Predicted seminal fluid proteins transferred in *Aedes aegypti* ejaculate

Found at: doi:10.1371/journal.pntd.0000989.s001 (0.18 MB DOC)

**Table S2** Predicted sperm proteins transferred in *Aedes aegypti* ejaculate

Found at: doi:10.1371/journal.pntd.0000989.s002 (0.11 MB DOC)

**Table S3** Proteins identified from unlabeled unmated *Aedes aegypti* female sample

Found at: doi:10.1371/journal.pntd.0000989.s003 (0.07 MB DOC)

**Table S4** Amino acid sequences of unannotated predicted sperm and seminal fluid proteins from *Aedes aegypti*

Found at: doi:10.1371/journal.pntd.0000989.s004 (0.07 MB DOC)

**Table S5** *Aedes aegypti* seminal fluid proteins and sperm proteins indistinguishable by peptides identified through mass spectrometry

Found at: doi:10.1371/journal.pntd.0000989.s005 (0.05 MB DOC)

**Table S6** Putative *Aedes aegypti* sperm proteins that were not detected as transferred to females during mating

Found at: doi:10.1371/journal.pntd.0000989.s006 (0.08 MB DOC)

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**Author Contributions**
Conceived and designed the experiments: LKS MEHH MFW LCH. Performed the experiments: LKS MEHH MK. Analyzed the data: LKS MEHH MFW LCH. Contributed reagents/materials/analysis tools: MFW LCH. Wrote the paper: LKS MCH MEHH JMCR PD. Contributed data/materials/analysis tools: PD. Generated databases of predicted peptides for protein discovery, and generated output for each of these proteins on predicted protein domains, homologs, and other features: JMCR.

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