Identification of Aedes aegypti Long Intergenic Non-coding RNAs and Their Association with Wolbachia and Dengue Virus Infection

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Abstract

Long intergenic non-coding RNAs (lincRNAs) are appearing as an important class of regulatory RNAs with a variety of biological functions. The aim of this study was to identify the lincRNA profile in the dengue vector Aedes aegypti and evaluate their potential role in host-pathogen interaction. The majority of previous RNA-Seq transcriptome studies in Ae. aegypti have focused on the expression pattern of annotated protein coding genes under different biological conditions. Here, we used 35 publically available RNA-Seq datasets with relatively high depth to screen the Ae. aegypti genome for lincRNA discovery. This led to the identification of 3,482 putative lincRNAs. These lincRNA genes displayed a slightly lower GC content and shorter transcript lengths compared to protein-encoding genes. Ae. aegypti lincRNAs also demonstrate low evolutionary sequence conservation even among closely related species such as Culex quinquefasciatus and Anopheles gambiae. We examined their expression in dengue virus serotype 2 (DENV-2) and Wolbachia-infected and non-infected adult mosquitoes and Aa20 cells. The results revealed that DENV-2 infection increased the abundance of a number of host lincRNAs, from which some suppress viral replication in mosquito cells. RNAi-mediated silencing of lincRNA_1317 led to enhancement in viral replication, which possibly indicates its potential involvement in the host anti-viral defense. A number of lincRNAs were also differentially expressed in Wolbachia-infected mosquitoes. The results will facilitate future studies to unravel the function of IncRNAs in insects and may prove to be beneficial in developing new ways to control vectors or inhibit replication of viruses in them.

Author Summary

Aedes aegypti is a major vector of several viruses such as dengue and Zika viruses. Understanding the intricate interaction of viruses with mosquito vectors and the factors involved in virus replication are essential for developing effective arbovirus control strategies. In
this study, we report a comprehensive list of long intergenic non-coding RNAs encoded by the genome of *Ae. aegypti* for the first time. In addition, we show that a number of these long non-coding RNAs are differentially expressed in mosquitoes infected with dengue virus, which could be involved in DENV-mosquito interaction. The outcomes provide a new avenue to explore mosquito biology and mosquito-virus interactions that may lead to the discovery of molecules that could be beneficial for vector manipulation.

**Introduction**

Dengue and Zika viruses are related mosquito-borne viruses that have a common primary vector, *Aedes aegypti* and infect millions of people worldwide [1,2]. Recent outbreaks of Dengue and Zika in South America pose a serious risk for other tropical regions in the world as *Ae. aegypti* is one of the most abundant mosquito species in these areas [2]. Although certain vaccines have been licensed in some countries, there are no efficient specific therapeutics available for either diseases, hence, the best protection against their global spreading is an efficient vector control program [3,4].

The genome sequence of *Ae. aegypti* is available, however, it has not been fully annotated. Only 2% of its large genome (1.376 Mb) has been annotated as protein coding genes and it reflects the presence of great proportions of non-coding transcripts as well as repetitive elements [5]. Transcriptomic changes, including those of non-coding transcripts, could provide a genome scale insight into host-pathogen interactions. Previous studies identified a series of small ncRNAs in *Ae. aegypti* and demonstrated their interaction with arboviruses [6–9], but our knowledge about their long ncRNAs is limited.

RNA transcripts longer than 200 nucleotides, which do not contain an open reading frame of longer than 100 amino acids, are simply defined as long ncRNA [10]. Generally, they are classified by their location relative to their neighboring protein-coding genes and include the long intergenic ncRNA (lincRNA), intronic lncRNA, antisense lncRNA and enhancer RNA [10]. Although a number of mammalian lncRNAs have been characterized and identified in the last few years, genome-wide identification of this class of ncRNAs has only recently become possible with the arrival of deep sequencing technologies. An expanding body of evidence reveals that lncRNAs, once described as dark matter, are involved in many biological processes such as genomic imprinting and cell differentiation [11]. They also play important roles in epigenetic and non-epigenetic based gene regulation [12]. Relatively, little is known about their involvement in activation and differentiation of immune cells, but new discoveries have revealed the involvement of lncRNA in defense systems [13]. Previous works have also outlined their quick responses to different stimuli and stress factors [14–17]. In addition, it has been shown that some lncRNAs enhance virus replication or decrease antiviral immunity [18].

Although in most host-virus interaction studies typically protein-coding genes have been the center of attention, there are few examples of virus and host lncRNA interactions in human and mouse models [18,19]. For instance, Hepatitis B virus (HBV) infection altered IncRNA profiles in patients, with about 4% of human IncRNAs showing more than 2-fold changes in HBV infected liver tissue [20]. Winterling et al. (2014) identified a virus inducible IncRNA, which is induced by vesicular stomatitis virus and several strains of Influenza A virus (IAV) [18].

The sequence and structure of lncRNAs are important in their function, in particular for their interaction with DNA, RNA, or proteins. In case of extensive base-pairing of IncRNA with target mRNA, translation can be stabilized, while partial base-pairing may accelerate
mRNA decay or inhibit translation of the target mRNA [21]. It has been shown that some lncRNAs interact with other small ncRNAs such as miRNAs. For example, in silkworm, 69 lncRNAs originating from 33 gene loci, may serve as miRNA precursors, and 104 lncRNAs may function as competing endogenous RNAs (ceRNAs) [22]. LncRNAs are also targeted by miRNAs similar to mRNAs and reduce their stability. They may also act as sponge or decoy of miRNAs, and compete with miRNAs for binding to mutual target mRNAs [21].

In insects, only a few genes have been experimentally annotated as lncRNA. It has been estimated that more than 5000 loci potentially encode non-coding transcripts in *Drosophila melanogaster*, however, just seven loci (bxd, Hsro, pgc, roX1, rox2, sphinx and yar) have been annotated as functional regulatory lncRNAs by experimentally derived data [23,24]. We recently found that a number of lncRNAs in *Plutella xylostella*, a pest of cruciferous plants, were linked to the insect’s resistance to insecticides and might be involved in detoxification processes [14]. Jenkins et al (2015) identified 2,949 lncRNAs in the malaria mosquito vector, *Anopheles gambiae*, using RNA-Seq data [25]. They showed that in various *Anopheles* species, lncRNAs have considerably lower sequence conservation as compared with protein-coding genes. In another study, it has been shown that 43% of total midgut transcripts of *An. gambiae* are lncRNAs and 32% of them showed some level of homology to other species [26].

The current study generated a comprehensive list of *Ae. aegypti* lincRNAs, which will be a complement to the other ncRNAs (microRNAs and piRNAs) that have already been discovered in this medically important species. This work also helps to improve the present annotation of the genome of *Ae. aegypti*. We also examined the expression pattern of some selected lincRNAs in response to microbial challenge namely dengue virus serotype 2 (DENV-2) and *Wolbachia* infections to identify potential immune-related lincRNAs in *Ae. aegypti* [27,28]. The results help better understanding of mosquito-pathogen interactions providing new insights on the potential role of lncRNAs as candidates for exploitation to inhibit replication of mosquito-borne viruses.

**Methods**

**RNA-Seq Data preparation**

Previously sequenced RNA-Seq raw data of *Ae. aegypti* were downloaded from NCBI Sequences Read Archive and ArrayExpress Archive with accession numbers SRA048559, SRA058076, SRA244067 and ERP002530 [29–32]. Raw data were stripped of adapters using CLC Genomic Workbench version 7.5.1 and reads with quality score of above 0.05 and maximum 2 ambiguous sequences were retained for further analysis.

**Large gap mapping and transcript discovery**

The CLC Genomic workbench’s Transcript Discovery plugin was used for lincRNA discovery in the *Ae. aegypti* genome. New transcripts were identified by large gap mapping of 1,148,814,115 reads of 35 RNA-Seq libraries to the genomic reference (*AaegL3.3*). We implemented strict mapping criteria (mismatch, insertion and deletion costs: 2: 3: 3 respectively). The minimum similarity and length fraction of 0.9 between a mapped segment and the reference were allowed as part of the mapping criteria. The large gap mapper algorithm also requires each mapped segment to include at least 10% of the read with minimum length of 17 bases. We considered a gap with maximum of 50 Kbp distance between mapped read segments to span the introns from RNA-Seq data. The annotations were generated by inspecting mapping of reads and identifying likely regions corresponding to genes, including their exons and splice sites. The algorithm scans each gap in the read mapping to explore whether the gap is assigned to a valid splice site or can be relocated to a valid splice site without cost.
**lincRNA identification pipeline**

A rigorous filtering pipeline was developed to remove transcripts that may potentially encode proteins. The pipeline for *Ae. aegypti* lincRNA discovery is summarized in Fig 1. We identified 75,069 potential genes using the CLC Genomic Workbench transcript discovery algorithm. The genes that were annotated as known *Ae. aegypti* protein-coding genes were discarded and 30,865 potential genes were also checked for any exon or intron overlap with other known *Ae. aegypti* protein-coding genes. We selected 22,079 sequences, which were located more than 1kb away from any other known transcripts, for finding putative open reading frames (ORF). All possible six frames were produced for all selected sequences and then the translated sequences were subjected to a domain search to identify any putative conserved protein domains through Pfam v27.0 database [33]. We discarded 8,795 sequences with potential ORF above 100 aa or conserved protein domains. The remaining sequences were submitted to a coding potential assessment tool (CPAT), which utilizes a logistic regression model made with four sequence features: ORF size, ORF coverage, Fickett TESTCODE statistics and hexamer usage bias [34]. We applied the coding probability threshold of 0.3, which led to discarding 376 sequences as putative coding RNAs. We also implemented an expression threshold on our data to strengthen the identification pipeline. Sequences with more than 10 mappable reads in at least 17 out of 35 RNA-Seq libraries were considered as valid sequences and were kept for the next step. Any possible similarity with other known proteins was found by using BLASTx algorithm against nr and Swiss port database (E-value cut off 10\(^{-5}\)). Finally, 3,842 potential lincRNAs were identified and used for further analysis.

To identify *Ae. aegypti* putative lincRNAs that are regarded as small RNA associated lincRNAs, we used the Blast algorithm to search for *Ae. aegypti* precursor miRNA sequences in the predicted *Ae. aegypti* lincRNA dataset. We also used publicly available small RNA libraries from DENV-infected and non-infected samples (SRP026241) in this analysis for further characterization of lincRNA_1317. All known *Ae. aegypti* miRNA sequences were mapped to lincRNA_1317 for possible best fitting using RNAhybrid, which is a tool for finding the normalized minimum free energy (mfe) of RNA. We did not allow G:U pairing in the seed region (nucleotide 2–8) and required miRNA-lincRNA duplexes to have a helix in this region. Maximum 5nt were allowed as unpaired nucleotides in either side of an internal loop. LncTar algorithm [35] was used to explore any potential interaction between lincRNA_1317 and DENV-2 genome (accession no. NC_001474.2) by finding the normalized mfe joint structure of two RNA molecules based on base pairing.

**Identification of differentially expressed lincRNAs upon infection**

The *Ae. aegypti* genome was annotated with the final list of lincRNAs and used as reference for RNA-Seq analysis in CLC Genomic Workbench. To measure the lincRNA normalized expression value, RPKM (Reads Per Kilobase per Million reads) was assigned for each library [36]. To find the differential expression pattern of lincRNAs in response to DENV infection, data from DENV-2 (Jam1409) infected midgut and carcass tissues at 4 days post-infection (dpi) were compared with their corresponding control groups [30]. Baggerley’s test, a count based statistical analysis was done on the data. The samples were given weights depending on their total counts. Based on the test the weights are obtained by supposing a Beta distribution on the proportions in a group, and estimating these, along with the proportion of a binomial distribution. We selected 20 potential lincRNAs with more than 4-fold change for further analysis with RT-
Fig 1. The lincRNA identification pipeline flowchart.

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qPCR in DENV-2 (New Guinea C strain) infected *Ae. aegypti* cell line (Aa20) and screened their expression profile in *Wolbachia*-infected mosquitoes.

*Ae. aegypti* infected with the wMelPop-CLA strain of *Wolbachia* (+Wol) and without *Wolbachia* (-Wol, tetracycline-cured line) were stocks produced previously [37]. For the experiments in this work, 4-day-old female mosquitoes were used from which total RNA was extracted with 6–10 adult mosquitoes for each biological replicates.

**Expression analysis of *Ae. aegypti* lincRNAs**

Detection and validation of the relative abundance of selected lincRNAs was carried through lincRNAs’ specific primers using SYBR Green chemistry in real time PCR machine. Briefly, total RNA was extracted from cells using Qiazol reagent according to the manufacturer’s instructions (Qiagen). The TURBO DNA-free kit (Ambion, USA) was used to remove possible genomic DNA contamination in RNA samples. First strand cDNA was synthesized from 2.5 μg of RNA using a poly-dT primer and Superscript III reverse transcriptase (Life Technologies). qPCR primers were designed using primer design tool of NCBI [38]. Quantifast SYBR Green PCR Master Mix with ROX was used to quantify the relative expression of lincRNAs between different treatments. Three independent biological replicates were considered along with three technical replicates for each treatment. Reactions were performed in a Rotor-Gene thermal cycler (Qiagen) under the following conditions: 95°C for 5 min, and 40 cycles of 95°C for 10s and 60°C for 30s, followed by the melting curve (68°C to 95°C). Melting curves were analysed to examine the specificity of amplification. Relative expressions were calculated using the Rotor-Gene software and the mosquito *RPS17* as reference gene for normalization. Unpaired t-test was used to identify statistically significant differences.

**RNAi of selected lincRNAs and virus replication assay**

To check the functional importance of the identified novel lincRNAs, dsRNAs were synthesized to knockdown selected lincRNAs (2329, 1613 and 1317) to check their effect on DENV replication. Briefly, primers with added T7 promoter sequence (S1 Table) were used to generate 250–600 bp PCR products from selected lincRNAs. Megascript T7 kit (Ambion) was used according to the manufacturer’s instruction to generate respective dsRNAs. To induce efficient RNA silencing, *Ae. aegypti* Aa20 cells were double transfected with dsRNAs against selected lincRNAs. Aa20 cells were re-suspended and ~3×10⁵ cells were added to each well of a 12-well plate. Cells were allowed to settle for ~1 h, medium was removed and replaced with a transfection mixture consisting of 0.5 ml medium (1:1 Schneider medium and Mitsuhashi–Maramorisch with 10% FBS), 8 μl Cellfectin (Invitrogen), and 5 μg dsRNA either for selected lincRNAs or GFP as control. Cells were also treated with 3 μg dsRNA 72 h after the primary transfection to increase the silencing efficiency of selected lincRNAs. Six hours after the secondary transfection, cells were infected at 1 multiplicity of infection (MOI) with DENV-2 (New Guinea C strain). All the treatments were collected three days post-infection. RNA extraction and cDNA synthesis were carried out as above. qPCR was performed to confirm the knockdown and the effect of particular lincRNA knockdown on the genomic RNA of DENV-2. Each treatment was repeated three times. All data from three biological replicates were subjected to one-way ANOVA statistical analysis. Brown-Forsythe test was used to check the equality of group variances and Tukey’s multiple comparisons test was also used to examine significant statistical differences among treatments.
Results and Discussion

Identification and characterisation of *Ae. aegypti* lincRNAs

In total, 3,482 putative lincRNAs in 1,114 *Ae. aegypti* genome scaffolds were identified (S2 Table). The *Ae. aegypti* lincRNA genes displayed a slightly lower GC content (mean: 40.1%) in comparison to 47.8% in their protein-coding gene sequences (Fig 2A). The lower GC content or AT enrichment is a typical characteristic of lincRNAs and our findings are congruent with predicted lincRNAs in other species [14,39,40]. The majority of *Ae. aegypti* predicted lincRNAs are smaller than 3000 bases and their length distribution is represented in Fig 2B. These mosquito lincRNA candidates are notably shorter in length than protein-coding genes.

![Graphs showing the distribution of GC content, sequence length, gene length, and lincRNA distribution among different *Ae. aegypti* genome scaffolds.](image)

**Fig 2. Aedes aegypti lincRNA characterization.** A) Comparison of the GC content in protein-coding genes and the putative lincRNA genes. B) Sequence length distribution of *Ae. aegypti* lincRNA candidates. C) Comparison of gene length in protein-coding genes and putative lincRNA genes. D) lincRNA distribution among different *Ae. aegypti* genome scaffolds. The majority of scaffolds (~77%) only contain 1–4 lincRNAs, while only 23 *Ae. aegypti* genome scaffolds contain more than 10 lincRNAs (~2%).

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demonstrating another well-known characteristic of lincRNA transcripts (Fig 2C) [41,42]. The majority of *Ae. aegypti* genome scaffolds contain less than five lincRNA loci (~80%), however, 23 of scaffolds (2%) were enriched with more than 10 lincRNAs (Fig 2D). The detailed information of these scaffolds, which contain the highest number of lincRNAs are summarized in Table 1.

We examined all the identified lincRNAs to determine their association with *Ae. aegypti* miRNA precursors and piRNA clusters. We found that the pre-miRNA sequences of aae-miR-2940 and aae-miR-285 are located in lincRNAs 1431 and 3299, respectively. We could not detect any other pre-miRNA sequences identified in *Ae. aegypti* in the lincRNAs. Also, lincRNA 1978 and 792 are originated from two previously reported piRNA clusters [43] located at supercontig 1.478 and 1.98, respectively.

LincRNAs demonstrate low evolutionary sequence conservation even among closely related species [10,14]. We used the BLAST algorithm bit score to identify the level of similarity among *Ae. aegypti* lincRNA sequences with other closely related insect genomes such as *Aedes albopictus*, *Culex quinquefasciatus* and *Anopheles gambiae* (Fig 3A). As expected, most of the identified lincRNAs showed high level of similarity with *Ae. albopictus* genome sequence and probably are genus specific. The E-value cut off $10^{-50}$ was applied to our screening with the BLAST algorithm to identify the conserved sequences. Although the *Ae. aegypti* lincRNAs shared high level of sequence similarity with the genome of *Ae. albopictus*, only 62 and 7 lincRNAs had sequence similarity with *Cx. quinquefasciatus* and *An. gambiae*, respectively (Fig 3B). They were mostly limited to a single short region with high conservation.

Table 1. Distribution of potential *Ae. aegypti* lincRNAs in different genome scaffolds with more than 10 lincRNAs and their comparison with the number of protein-coding genes.

| Scaffold | Length (kbp) | Number of known genes | Number of lincRNAs | Length range (bp) | Average size of lincRNA |
|----------|--------------|-----------------------|--------------------|-------------------|------------------------|
| supercont1.1 | 5,856,339 | 124 | 20 | 261–7780 | 1,869 |
| supercont1.16 | 4,402,401 | 62 | 19 | 264–7065 | 1,761 |
| supercont1.35 | 3,598,302 | 34 | 17 | 420–6101 | 1,850 |
| supercont1.3 | 5,167,134 | 61 | 16 | 297–5683 | 1,897 |
| supercont1.19 | 4,221,289 | 48 | 14 | 339–6537 | 1,776 |
| supercont1.70 | 2,929,944 | 29 | 14 | 509–6314 | 2,039 |
| supercont1.37 | 3,744,586 | 35 | 13 | 497–8297 | 3,585 |
| supercont1.28 | 3,768,427 | 45 | 13 | 235–7876 | 2,229 |
| supercont1.29 | 3,855,786 | 42 | 13 | 301–5559 | 2,376 |
| supercont1.78 | 2,909,025 | 17 | 13 | 211–5561 | 1,497 |
| supercont1.18 | 4,266,046 | 76 | 12 | 400–7336 | 2,087 |
| supercont1.6 | 5,075,626 | 93 | 12 | 547–3182 | 1,380 |
| supercont1.225 | 1,723,990 | 19 | 12 | 489–4846 | 1,475 |
| supercont1.38 | 3,498,553 | 45 | 12 | 248–8797 | 2,080 |
| supercont1.46 | 3,321,798 | 42 | 12 | 248–7447 | 2,210 |
| supercont1.5 | 5,058,281 | 60 | 12 | 323–4291 | 2,168 |
| supercont1.120 | 2,427,180 | 38 | 11 | 805–5190 | 2,116 |
| supercont1.244 | 1,610,334 | 30 | 11 | 503–6058 | 2,512 |
| supercont1.49 | 3,164,279 | 51 | 11 | 324–5492 | 1,988 |
| supercont1.107 | 2,543,601 | 42 | 11 | 416–8896 | 2,876 |
| supercont1.22 | 4,100,794 | 51 | 11 | 409–4234 | 1,441 |
| supercont1.44 | 3,232,429 | 47 | 11 | 271–8994 | 2,175 |
| supercont1.92 | 2,802,290 | 34 | 11 | 242–4424 | 1,667 |

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Aedes aegypti lincRNAs change upon microbial challenge

Following the identification of *Ae. aegypti* lincRNAs, we analyzed their transcript levels in DENV-2 infected mosquitoes. To produce the lincRNA profile of infected and non-infected mosquitoes, we re-analyzed previously published RNA-Seq data from *Ae. aegypti* midgut and carcass samples at 4 dpi [30] (Fig 4). 248 and 203 lincRNAs with fold changes above four were identified in the RNA-Seq libraries of midgut and carcass, respectively (S3 Table). The majority of differentially expressed lincRNAs were considerably overexpressed in both samples. The abundance of only 32% of *Ae. aegypti* lincRNA candidates decreased in response to DENV-2 infection in the mosquito carcass sample. Thirty lincRNAs were differentially expressed in both examined samples. The transcription levels of 72 lincRNAs increased after infection while their expression could not be detected in the non-infected midgut tissue sample.

We selected 20 candidates of those differentially expressed lincRNAs from RNA-Seq analysis data for further investigation. The relative expression of lincRNA candidates were examined by reverse transcription quantitative polymerase chain reaction (RT-qPCR) upon DENV-2 infection in Aa20 mosquito cells. Only significantly overexpressed lincRNAs after DENV-2 infection are represented in Fig 5. Although we used Aa20 cells for the lincRNA expression assays, the expression patterns of almost all the examined lincRNAs (5 out of 6) were consistent with the RNA-Seq data (adult mosquito carcass sample). We used a poly-dT primer to produce cDNA, which also confirmed that all of those identified transcripts have poly-A tails and therefore are true transcripts. Based on these results, significant increase in the transcription levels of a selected number of *Ae. aegypti* lincRNAs suggests their possible involvement in host-pathogen interaction but further investigations are required to confirm their roles in antiviral/immune responses.

We also examined the impact of an endosymbiotic bacterium, *Wolbachia*, on some selected *Ae. aegypti* lincRNAs, which showed significant changes in response to DENV-2 infection. This gram-negative bacterium is transmitted maternally and potentially infects more than 40% of all insect species, manipulating its hosts using different strategies [44,45]. A fascinating
aspect of Wolbachia infection is limiting replication of vector-borne pathogens in mosquitoes [45,46]. However, the mechanism(s) behind virus blocking is largely unknown. Here, we found

![Volcano plot of differentially expressed Aedes aegypti lincRNAs in DENV-2 infected samples (midgut and carcass) compared with their corresponding controls. Dots with red color represent lincRNAs with more than 4-fold changes due to DENV-2 infection.](image)

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![DENV infection leads to changes in the abundance of Aedes aegypti lincRNAs.](image)

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![Bar charts showing the relative transcript levels of selected numbers of Aedes aegypti lincRNAs.](image)

Fig 5. DENV infection leads to changes in the abundance of Aedes aegypti lincRNAs. The relative transcript levels of selected numbers of Aedes aegypti lincRNAs were measured by RT-qPCR analysis of Aa20 cells infected with 1 MOI of DENV-2 for three days. Three biological replicates were used for each treatment with three technical replicates each. *, p < 0.05; **, p < 0.01; ****, p < 0.0001.
that the transcript levels of several lincRNA genes significantly increased in Wolbachia-infected *Ae. aegypti* mosquitoes (Fig 6), which may lead to differential regulation of cellular protein-coding genes. Our previous studies showed that *Wolbachia* could manipulate host small ncRNAs such as miRNAs and piRNAs [47]. An overall induction of small ncRNAs between 18 and 28 nucleotides was also observed in *Ae. aegypti* cell line infected with wMelPop-CLA strain of *Wolbachia* [48]. It was assumed that the upregulation of small ncRNAs in infected cells may result in an enhanced immune response and activated RNAi pathway. However, the role of these modifications in the host lincRNA gene expression profile, and potentially in anti-viral responses, is unknown and may lead to the discovery of lincRNAs that could be utilized for inhibition of virus replication in mosquitoes.

A recent study on mouse bone marrow-derived macrophage (BMDM) model reported a significant upregulation in 72 lincRNAs after treatment with the synthetic bacterial lipoprotein Pam3CSK4, which acts through Toll-like receptor [49]. In another study, differential expression of approximately 500 annotated mouse IncRNAs was reported during infection with severe acute respiratory syndrome coronavirus [50]. Recently, it has been shown that honeybee lincRNAs are also differentially expressed during infection with various viruses such as sacbrood virus (SBV) and deformed wing virus (DWV), but the biological significance of these lincRNAs is completely unknown [51]. Although exploring the in vivo functions of immune-related lincRNAs is one exciting area for future studies, the differential expression of some
lincRNAs could simply be byproducts of mRNA biogenesis or changes in global transcriptional profile due to microbial challenges [52,53]. Struhl (2007) believed that the transcriptional machinery is not perfect producing RNAs that serve no purpose or have no significant role in infection [54]. On the other hand, there are several examples which have shown that lincRNAs could be potentially important factors in host antimicrobial responses, and may represent a new class of signaling molecules involved in innate immunity or provide a new layer in gene regulation. For instance, two interferon (INF) induced lncRNAs, which were upregulated by influenza and vesicular stomatitis viruses, regulate the expression of the antiviral factor tetherin in human HuH7 cells [55].

RNAi of selected lincRNAs and enhancement of DENV-2 replication

To confirm the role of DENV-induced lincRNAs in viral replication, we used RNAi-mediated silencing of two selected lincRNAs (lincRNA_1317 and 1613) using dsRNA in Aa20 cells followed by DENV-2 infection. Only RNAi-mediated silencing of lincRNA_1317 led to enhancement of DENV-2 replication (Fig 7A). Silencing of the lincRNA was confirmed by RT-qPCR (Fig 7B). Interestingly, expression of Aedes aegypti lincRNA_1317 increased substantially following the progression of infection (Fig 7C) suggesting that this lincRNA might be involved in antiviral response. This idea is consistent with the finding that lincRNA_1317 was also highly overexpressed (2.33 fold) in Wolbachia-infected mosquitoes as compared with non-infected mosquitoes (Fig 6).

While there are no reports on the involvement of lncRNAs in host-pathogen interactions in insects, time-dependent over-expression of host lincRNAs in response to viral infection has been observed in humans. A recent study showed more than 80% of host cell lincRNAs were upregulated upon an adenovirus infection of human primary lung fibroblast cells [56]. Zhang et al. (2013) reported alterations of expression of cellular lncRNAs in HIV-1-infected T cells. Among differentially expressed lncRNAs, NEAT1 expression notably increased in infected cells. When NEAT1 was silenced, virus production was enhanced by increasing the nucleus-to-cytoplasm export of HIV-1 transcripts containing Rev-dependent instability element [57]. A significant induction in this lincRNA expression in response to influenza virus and herpes simplex virus infection has also been shown [58].

To further investigate the potential role of lincRNA_1317 in mosquito-pathogen interaction, we determined its association with host endogenous small RNAs and its possible direct interaction with DENV. Although this lincRNA is not located in any of the known piRNA clusters, the majority of mappable small RNA reads to its sequence are in the range of 26–29 nt (S1 Fig). However, there was no difference in the mapping pattern and mapped read length distribution when reads from DENV-infected and non-infected small RNA libraries were mapped to lincRNA_1317 (S1 Fig). It has been shown that piRNA-like small RNAs have a large impact on lincRNA transcriptome [57], but our knowledge about the function of piRNA-mediated lncRNAs is still limited. Recently, it has been reported that piRNAs derived from transposons and pseudogenes facilitate the degradation of lncRNAs in mouse late spermatocytes [57].

Next, we hypothesized that Aedes aegypti lincRNA_1317 response to microbial challenge could be due to cross-regulation between miRNAs and the lincRNA. Aedes aegypti miRNA recognition elements on lincRNA_1317 were identified by calculating the normalized minimum free energy (mfe) of hybridization for each Aedes aegypti miRNA and lincRNA_1317 using RNAhybrid core script. Binding site enrichment was detected for a few miRNAs with more than two recognition elements (Table 2). For instance, more than four recognition sites were predicted for miR-278-5p and miR-252-3p on lincRNA_1317. We also identified some hot spots for miRNA recognition sites on lincRNA_1317, which may allow multiple miRNAs to bind to the
same regions (S2 Fig). miRNAs can reduce lincRNA stability by targeting their transcripts similar to mRNAs. Also, lincRNAs with multiple recognition sites may actually be competitive inhibitors of miRNA function and stopping them from binding to their genuine targets by sequestering them [21]. Although the mfe for some of those miRNA-lincRNA recognition sites suggests high probability of a binding event, further experimental investigations are required to validate this interface.

We also used LncTar algorithm to predict any direct interaction between lincRNA_1317 and DENV-2 genome. One potential interaction was predicted in the region 1–3370 of lincRNA_1317 and the region of 3210–6579 of DENV-2 genome with mfe of -61.73 (normalized dG -0.0184). This tool has accuracy rate of 80% [35], but does not consider the tertiary
The involvement of lincRNA_1317 in host response to viral infection might be through its interactions with regulatory proteins that are involved in epigenetic changes by directly interacting with chromatin modifying enzymes or DNA binding proteins such as transcription factors. This interaction has been shown in several examples in mammalian systems, including host-virus interactions in which lncRNAs mediate antiviral responses by controlling the expression of immune-related genes (reviewed in [58]).

Although our knowledge of the biological function of this class of ncRNAs in mosquitoes is still limited, the results generated from this study will facilitate forthcoming explorations of lincRNA functions in insects. Clearly, further research is required to provide concrete experimental evidence to support the role of lincRNA_1317 or any other Ae. aegypti lincRNAs in host-pathogen interaction. With advances in technology, the mosquito lincRNA-protein interactions can be identified using high-throughput sequencing of immunoprecipitated RNA after cross-linking (CLIP-Seq). Further, functional studies could be carried out to characterize immune-related lincRNAs. The involvement of lincRNAs in pathways associated with responses to viral infection and cellular stress makes them interesting candidates as potential targets for manipulation to inhibit virus replication or control vector populations.

### Supporting Information

S1 Fig. Length distribution of small RNA reads mapped to lincRNA_1317.

(TIF)
S2 Fig. The miRNA recognition hot spot sites on lincRNA_1317.
(TIF)

S1 Table. List of primers used in this study.
(DOCX)

S2 Table. Identified lincRNA candidates in Ae. aegypti and their genome coordinates.
(XLSX)

S3 Table. Differentially expressed lincRNAs in response to DENV-2 infection in midgut and carcass.
(XLSX)

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