The *Juan* non-LTR retrotransposon in mosquitoes: genomic impact, vertical transmission and indications of recent and widespread activity

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Abstract

**Background:** In contrast to DNA-mediated transposable elements (TEs), retrotransposons, particularly non-long terminal repeat retrotransposons (non-LTRs), are generally considered to have a much lower propensity towards horizontal transfer. Detailed studies on site-specific non-LTR families have demonstrated strict vertical transmission. More studies are needed with non-site-specific non-LTR families to determine whether strict vertical transmission is a phenomenon related to site specificity or a more general characteristic of all non-LTRs. *Juan* is a Jockey clade non-LTR retrotransposon first discovered in mosquitoes that is widely distributed in the mosquito family *Culicidae*. Being a non-site-specific non-LTR, *Juan* offers an opportunity to further investigate the hypothesis that non-LTRs are genomic elements that are primarily vertically transmitted.

**Results:** Systematic analysis of the ~1.3 Gbp *Aedes aegypti* (*Ae. aegypti*) genome sequence suggests that *Juan-A* is the only *Juan*-type non-LTR in *Aedes aegypti*. *Juan-A* is highly reiterated and comprises approximately 3% of the genome. Using minimum cutoffs of 90% length and 70% nucleotide (nt) identity, 663 copies were found by BLAST using the published *Juan-A* sequence as the query. All 663 copies are at least 95% identical to *Juan-A*, while 378 of these copies are 99% identical to *Juan-A*, indicating that the *Juan-A* family has been transposing recently in evolutionary history. Using the 0.34 Kb 5' UTR as the query, over 2000 copies were identified that may contain internal promoters, leading to questions on the genomic impact of *Juan-A*. *Juan* sequences were obtained by PCR, library screening, and database searches for 18 mosquito species of six genera including *Aedes*, *Ochlerotatus*, *Psorophora*, *Culex*, *Deinocerites*, and *Wyeomyia*. Comparison of host and *Juan* phylogenies shows overall congruence with few exceptions.

**Conclusion:** *Juan-A* is a major genomic component in *Ae. aegypti* and it has been retrotransposing recently in evolutionary history. There are also indications that *Juan* has been recently active in a wide range of mosquito species. Furthermore, our research demonstrates that a Jockey clade non-LTR without target site-specificity has been sustained by vertical transmission in the mosquito family. These results strengthen the argument that non-LTRs tend to be genomic elements capable of persistence by vertical descent over a long evolutionary time.
Background

TEs, or mobile genetic elements, are integral components of the eukaryotic genomes. Because they have the ability to replicate and spread in the genome as primarily "selfish" genetic units [1], TEs tend to occupy significant portions of the genome [2]. Recent evidence suggests that the "selfish" property may have enabled TEs to provide the genome with potent agents to generate tremendous genetic and genomic plasticity [3]. TEs transpose through either RNA-mediated or DNA-mediated mechanisms [4]. DNA-mediated TEs generally transpose by a cut-and-paste process, directly from DNA to DNA. RNA-mediated TEs transpose by a replicative process that involves transcription, reverse transcription, and integration of cDNA molecules. TEs in this category include the long terminal repeat (LTR) retrotransposons, non-LTRs, or long interspersed nuclear elements (LINES), and short interspersed nuclear elements (SINEs).

It has been proposed in models of the lifecycle of DNA-mediated TEs [5-7] that most TEs will eventually become inactivated in a given species, which underscores the importance of horizontal transfer for TE survival, a mechanism that allows TEs to invade a naive genome. Horizontal transfers DNA-mediated TEs are well documented [8-10]. There are also cases of non-LTR horizontal transfer proposed [11-16], the most convincing case involving RTE clade elements [11,15,16]. RTE non-LTRs were first found in C. elegans and encode a single open-reading frame (ORF) containing reverse transcriptase and endonuclease activities [17]. In contrast, it has been argued that there is no reliable evidence of non-LTR horizontal transfer between eukaryotes in the last 600 million years according to age vs. divergence analysis [18,19]. Research involving arthropod R1 and R2 families, which are site-specific non-LTRs that insert into 28S ribosomal RNA genes, shows vertical inheritance of these elements since the origin of the Drosophila melanogaster species subgroup, approximately 17–20 million years ago (MYA) [20]. Even multiple lineages have been found to coexist in the rRNA loci and be maintained by vertical descent [21]. Other studies on R1 and R2 lineages concluded that they have been vertically transmitted since the inception of the Drosophila genus, approximately 60 MYA or longer [22,23]. The site-specificity of R1 and R2 may result in a bias toward vertical transmission as site-specificity could offer a "safe haven", protecting the genome from deleterious insertions elsewhere.

Juan-A, a Jockey clade non site-specific non-LTR from Ae. aegypti has been reportedly involved in potential horizontal transfer between the non-sibling species Ae. albopictus and Ae. polynesiensis [24]. However, Crainey and colleagues [25] recently suggest that vertical transmission explains the evolutionary relationship between Juan elements in Ae. aegypti, Ae. albopictus, and Culex pipiens quinquefasciatus (herein referred to as C. quinquefasciatus). They also did not find evidence to support horizontal transfer of CR1 clade elements Q and T1 in mosquitoes, although an earlier report suggested horizontal transfer could conceivably explain the identities and distributions of CR1 families in diverse taxa [26]. Here we report a detailed evolutionary study of Juan in the mosquito family Culicidae. Sequences of full-length Juan elements have been reported from the yellow fever mosquito Ae. aegypti [24] and the house mosquito C. quinquefasciatus [27]. In this study, we have obtained sequences of Juan elements from 18 mosquito species of six genera. Our results support that non-LTRs are able to sustain their activity over long periods of evolutionary time relying primarily on vertical transmission while not excluding the possibility of rare horizontal transfer. Our whole-genome analysis suggests that Juan-A has been retrotransposing recently in evolutionary history, and it occupies approximately 3% of the Ae. aegypti genome. We have also discussed the potential evolutionary impacts of Juan-A in the Ae. aegypti genome.

Results

Juan in Aedes aegypti: abundance and recent activity

Juan-A contributes significantly to the genome size of Ae. aegypti, determined to be approximately 3% by RepeatMasker (see Methods). A highly variable copy number is found depending on what query region and identity criteria are used (Figure 1, Table 1). Juan-A appears to be the only Juan-type element in Ae. aegypti. After masking the genome sequence for Juan-A (80% nt identity) with RepeatMasker [28], tBLASTn [29] with Juan-A amino acid (aa) sequence was used to identify closely related families. The two closest families found, AaJockey64 and AaJockey66 have approximately 37% aa identity to Juan-A in the same region used for phylogenetic inference (Figure 1, Figure 2B).

Juan-A

| ORF1 | AP | RT |
|------|----|----|
| 939 bp |

5'UTR, 0.34 Kb

Full Length, 4.7 Kb

Figure 1

Structural organization of the Juan-A element of Aedes aegypti. ORF1 encodes a nucleic acid binding protein and ORF2 encodes both an apurinic/apyrimidinic (AP) endonuclease and reverse transcriptase (RT) domain. Arrows indicate the 939 bp region amplified by PCR that was used for phylogenetic inference. A canonical polyadenylation signal sequence is present in the 3' end of Juan-A (not shown). Regions used for copy number determination by database search in Table 1 are shown by horizontal lines.
Table 1: Copy numbers of Juan-A in Ae. aegypti determined by genomic analysis.

| Juan-A sequence region | 99%   | 98%   | 97%   | 95%   | 80%   | 70%   |
|------------------------|-------|-------|-------|-------|-------|-------|
| Full-length            | 378   | 637   | 662   | 663   | 663   | 663   |
| 5' UTR 0.34 Kb         | 957   | 1596  | 1920  | 2137  | 2274  | 2274  |
| 3' 0.34 Kb             | 180   | 867   | 1302  | 2886  | 4852  | 4853  |

* includes sequences having 90% length compared to a full-length Juan-A (see methods)

Figure 2
Phylogenetic comparison of Juan sequences and their hosts. A. Consensus tree of host phylogeny constructed with MrBayes (see methods) using nucleotide sequences of Vg-C, a single copy gene. Vg-C from An. gambiae is used to root the tree. Clade credibility values from 150,000 generations are given at each node. B. Consensus tree constructed with MrBayes (see methods) using nucleotide sequences of Vg-C, a single copy gene. Vg-C from An. gambiae is used to root the tree. Accessions are given beside sequence names. An asterisk indicates that the reading frame was intact. A bold capital "L" indicates that the sequence was obtained from a genomic library. The first three letters of a species name is used to label PCR and library sequences. Genus names are in bold beside brackets. Juan-A is from Ae. aegypti and Juan-C is from C. pipiens. Abbreviations: Ae. aegypti (aeg); Ae. albopictus (alb); Ae. simpsoni (sim); Ae. polynesiensis (pol); Ae. vexans (vex); C. molestus (mol); C. quinquefasciatus (qui); C. restuans (res); C. tarsalis (tar); D. cancer (can); O. atropalpus (atr); O. bahamensis (bahi); O. epactius (epa); O. taeniornithinclus (tae); O. triseriatus (tri); P. ciliata (cil); W. micelli (mic). Number indicates clone from PCR. Ae. aegypti and An. gambiae sequences from genomic database (e.g. Ag-Jock-1, AaJockeyEle2) can be found in the TEFam database [57]. See additional files 1 and 2 for alignments used for phylogenetic inference.
AaJockeyEle4 and AaJockeyEle6 are part of divergent Jockey clade families, not Juan-type elements (Figure 2B).

Analysis of Juan-A in Ae. aegypti reveals that the family has undergone recent amplification, evidenced by a high degree of homogeneity between copies (Table 1). We have chosen to look at groups of sequences having various identities to the Juan-A query to obtain a more comprehensive picture of Juan evolution in the genome. Using lower identity criteria should allow the identification of both older retrotransposed copies as well as copies from more divergent Juan-A-related sequences. There are 663 copies of full-length or nearly full-length Juan-A, defined as having at least 90% length and at least 70% nucleotide identity compared to the 4.7 kb published Juan-A sequence. All 663 copies are at least 95% identical to Juan-A, suggesting that there are no divergent subgroups among these nearly full-length Juan elements. Three hundred seventy-eight of the 663 copies are 99% identical to Juan-A, indicating that the Juan-A family has been transposing recently in evolutionary history. There is no appreciable difference in copy numbers between using 80% and 70% nt identity cutoffs (Table 1). As stated in the Methods, many gaps exist (36206 contigs) in the Ae. aegypti genome sequence assembly. We did not determine how many Juan-A copies were truncated by these gaps so it is possible that many more full-length copies exist in the genome. In addition, we used a gap parameter value of 50 bp for our BLAST processing program. Therefore, Juan sequences with insertions of over 50 bp are not counted as full-length sequences.

We also used 5’ and 3’ end regions as queries to get a general impression of Juan representation and activity in the genome. A higher number of 5’ ends were found than 3’ ends when looking at those copies that had greater than 97% identity to the query, a curious result. Using BLAST through NCBI, many Juan-A hits were found to EST sequences from full-length cDNA libraries (not shown). Several hits to the 3’ end of Juan-A were from sequencing reactions using oligo dT primers, indicating that these are from polyadenylated transcripts.

Juan appears to have been active throughout the mosquito family
Six species have 3 or more Juan sequences that share a high degree of intragenomic nt identity (Table 3). Values shown in Table 3 are the mean of comparisons of each sequence vs. the consensus generated from that group of sequences. Sequence identity of PCR clones ranges from approximately 97.1% in Ae. aegypti to 99.4% at the nucleotide level in C. quinquefasciatus. Four sequences from C. quinquefasciatus have over 99% identity. These do not appear to come from the same copy of Juan in the genome since a deletion is present in one sequence and substitutions can be found at various positions among the different sequences. PCR and library clones from 16 of 18 species yielded sequences that do not have frameshifts or stop codons within this analyzed portion of the ORF (Figure 2B). Altogether, these results indicate recent activity of Juan in both closely related and divergent species.

Negative selection has been acting on Juan
The rates of synonymous (dS) and nonsynonymous (dN) codon substitution have been commonly used as a measure of selection pressure. A value of dS/dN close to 1 is taken to indicate neutral selection as would be expected for a pseudogene. Values below and above 1 indicate positive and negative selection. Vitellogenin-C (Vg-C), a single copy yolk protein-encoding gene [30] was used as a

Table 2: Species from which Juan sequences were obtained by PCR or library screening.

| Genus        | Species                                   |
|--------------|-------------------------------------------|
| Aedes        | aegypti, albopictus*, polyomensis, simpsani, vexans |
| Ochlerotatus | atrapalpus, bahamensis, epactus, toeniorphus, triseriatus ciliato |
| Psorophora   |                                          |
| Culex        | molestus, nigripalpus, quinquefasciatus*, restuans, tarsalis cancer |
| Deinocerites |                                          |
| Wyeomyia     | michelli                                  |

Note: An asterisk indicates that sequences were obtained by both PCR and library screening.
Furthermore, there may complicate phylogenetic resolution here. The sequence is degenerate (note long branch length) and therefore may not affect the interpretation of the known to be a relatively fast-evolving gene [30]. We have also included Vg-C sequences from species for which vertical transmission has been previously constructed using Vg-C [30]. We have screened an Ae. albopictus genomic library and found grouped with Ae. polynesiensis sequences as expected according to known mosquito phylogeny. Another case involves C. quinquefasciatus, for which we also have sequences from both PCR and a genomic library. The two library sequences group with C. molestus and C. pipiens (Juan-C), as expected according to host phylogeny. However, the PCR sequences group most closely with C. nigripalpus. O. atropalpus (atr2, Figure 2B) and O. exactus (epa6, Figure 2B) sequences are nearly identical with over 99% nucleotide identity, but they come from species that are in the same species complex where introgression may exist.

**Discussion**

**Genomic impacts of Juan-A in Ae. aegypti**

Juan contributes approximately 3% to the Ae. aegypti genome sequence while the entire TE complement is estimated to be 47% (Ae. aegypti genome consortium, unpublished). With its significant contribution to genome size and the presence of hundreds of highly homogeneous full-length or near full-length copies, a natural question concerns the genomic impact of Juan. TEs can cause chromosomal inversions by providing sites for ectopic homologous recombination and by other mechanisms [31]. It might be thought that the hundreds of highly homogeneous copies might contribute to genomic instability.

Most non-LTR families usually consist of a large majority of 5’ truncated copies, which has been attributed to incomplete reverse transcription, template switching, or other mechanisms [32-35]. However, when using higher stringency for copy number determination (representing more recently amplified elements), there is a higher copy number of 5’ ends of Juan-A sequences than 3’ ends (Table 1). This could be a result of selection for 5’ ends, selection against 3’ ends, or possibly a distribution bias of 3’ end insertion into regions that are underrepresented in the genome sequence. Full-length non-LTRs have been shown to contain their own self-sufficient internal pol II promoter in the 5’ UTR [36-40]. It is interesting that so many 5’ UTRs of Juan-A are present in the genome. These 5’ UTRs, if functional as internal promoters, may produce a transcriptional burden. It is interesting to note that our

| Table 3: Juan sequences from several species of four genera have a high degree of sequence identity. |
|------------------------|-----------------|---|
| Species               | Nucleotide Identity | # Sequences compared |
| Ae. aegypti           | 99.0%            | 768 |
| Ae. simpsoni          | 97.1 +/- 0.3%    | 4  |
| C. molestus           | 98.5 +/- 0.2%    | 3  |
| C. quinquefasciatus   | 99.4 +/- 0.2%    | 4  |
| O. taeniorhinchus     | 91.1 +/- 0.1%    | 3  |
| W. michelli           | 97.5 +/- 0.7%    | 3  |

Note: only sequences in the same lineage are compared.
reporter assays have not demonstrated promoter activity of the Juan-A 5’UTR in cell lines from three mosquito species, while 5’UTRs of mosquito non-LTRs from 3 non-LTR clades have proven active in all 3 lines (not shown). Perhaps Juan-A is dependent on upstream promoter elements for transcription, as upstream sequences have been found to greatly influence the activity of the human L1 promoter activity [41]. Past analysis of Juan-C transcripts from cell culture showed that all transcripts analyzed were transcribed from upstream of the Juan element [42]. With its recent amplification and recent activity, the study of Juan may offer a good opportunity to increase our understanding of the competing forces of non-LTR activity and host regulation.

Juan evolution
To address the topic of vertical transmission and to analyze the distribution and evolution of Juan in Culicidae, a detailed phylogeny of the host species was needed. Most phylogenetic inferences of mosquitoes based on molecular data have been focused on the Anophèles genus due to its medical importance. More comprehensive analyses have been performed using the white gene [43] and Vg-C [30]. The Vg-C sequences available to us offered the most comprehensive phylogeny with many species from the Aedes and Culex genera, where Juan was discovered.

The Jockey clade is comprised of highly divergent families which have been found in several insect species [18,25,44,45]. Representatives of Juan have been reported in mosquitoes and in Drosophila [45]. However, those elements are distant relatives of the Juan-A and Juan-C elements (Figure 2B), which we are investigating in this study. We have focused on Juan-A and Juan-C (Juan sensu stricto) because use of paralogous sequences can lead to erroneous conclusions of phylogenetic relationships. Results from Crainey and colleagues (2005) are consistent with vertical transmission but they also included many paralogous sequences from three mosquito genera. As mentioned above, we focus on Juan sensu stricto and survey many mosquito species to investigate the question of Juan evolution. It is important to note that our results indicate that JuanDm [45] is not actually a Juan element, strictly speaking, since it groups with three divergent Ae. aegypti Jockey elements, having 99% support (Figure 2B). This underscores the importance of including many divergent representatives while performing phylogenetic inference.

Regarding the cases of potential horizontal transfer, there are alternative explanations. For the Ae. albopictus (alb 3, 6, 9, Figure 2B) and O. epactius (epa 6, Figure 2B) sequences, the first suspicion is genomic DNA contamination of the PCR reaction. The Ae. albopictus sequences obtained from a genomic library were found grouped with Ae. polynesiensis, as expected. It should be noted that the library was constructed from the Nepal strain and PCR was performed on the Oahu strain. Bensaadi-Merchermek, Salvador, and Mouches (1994) reported the absence of Juan-A from Ae. albopictus Oahu strain (1971). If our PCR results can be corroborated using other methods, this would suggest the horizontal transfer of Juan-A to this strain of Ae. albopictus. However, Juan-A was also reported absent from strains of Ae. polynesiensis and O. triseriatus [42], both species of which we were able to obtain PCR products that grouped phylogenetically as expected, supporting vertical transmission of these elements. For C. quinquefasciatus, sequences obtained from library screening correspond with the host phylogeny, being grouped in the C. pipiens species complex. In contrast, sequences obtained from PCR are found outside this group and placed closely with C. nigricalpus with approximately 94% nucleotide identity to nig5 (Figure 2B). Although possible, the nucleotide identities between the C. quinquefasciatus sequences and the C. nigricalpus sequence are not close enough to suspect genomic DNA contamination of the PCR. Another possibility is that different sublineages of Juan could have been sampled by PCR versus library screening. For example, there are two sublineages represented in O. taeniorhinchus. The amplification of Juan sequences from contaminating genomic DNA cannot be ruled out, especially when using degenerate primers with low stringency PCR conditions. This seems unlikely in the case of C. quinquefasciatus, because these multiple sequences form their own homogeneous group with high nucleotide identity. If they resulted from contaminating genomic DNA, then they would be expected to group with sequences of the contaminating species. In summary, there is evidence for multiple Juan lineages, which could explain some of the observed phylogenetic incongruence. However, further analysis is required to determine whether the phylogeny of the suspect sequences is due to horizontal transfer, genomic DNA contamination, or sampling of different sublineages.

Conclusion
It has been proposed that the horizontal transfer of non-LTRs are rare events and few reported cases have strong supporting evidence without alternative explanations [18,19]. In contrast, there are many cases documented for the horizontal transfer of DNA-mediated TEs. Without excluding the possibility of horizontal transfers, we find that Juan family members do mirror their host’s phylogeny closely, supporting the vertical transmission of these elements. Our results suggest the Juan family was able to sustain its activity in the mosquito family over long periods of evolutionary time. Estimates of the time since Aedes and Culex divergence would suggest that Juan has been maintained for at least 22–52 million years [46]. Furthermore, the presence of multiple Culicinae lineages approxi-
approximately 120 MYA has been proposed [47], suggesting that Juan may have persisted for at least this time. Detailed studies involving the site-specific non-LTRs R1 and R2 in Drosophila showed that they are vertically transmitted and are maintained in their respective genomes [20,22,23]. It may be argued that vertical transmission of R1 and R2 over a long evolutionary time could be unique to site-specific non-LTRs. This study, which was performed in a different insect group using a non-LTR that does not exhibit site-specificity, strengthens the hypothesis that non-LTRs are able sustain their activity without the need of horizontal transfer. It will be interesting to see if other non-LTRs behave in a similar fashion, especially those from other clades and divergent taxa that have not been studied in detail.

Methods

PCR amplification of genomic DNA and cloning

Degenerate primers GDFNAKH (forward) and FKNM-KAPG (reverse) (Sigma Genosys) were designed according to conserved amino acid sequence including 939 bp found in an alignment of ORF2 of the Juan element from Juan-A of Ae. aegypti and Juan-C of C. pipiens (Figure 1). In contrast to the commonly used RT region, we chose to use this less conserved region to increase resolution between sequences from closely related species. Genomic DNA was isolated from several individuals of a given species using the DNAzol Genomic DNA Isolation Reagent (Molecular Research Center). PCR was performed on genomic DNA from a total of 30 species of mosquitoes from 10 genera. The calculated Tms of the forward and reverse primers were 54.2°C and 62.7°C. Each 20 ul PCR reaction consisted of approximately 3 ng of genomic DNA, 1U of Taq polymerase (Takara), 1.5 mM MgCl2, and 0.2 mM each dNTP. PCR was performed by denaturation at 95 °C for 90s and 30 cycles of 95 °C for 30s, 48 °C for 50s, and 72 °C for 90s. Amplified products were size-separated on a 0.7% agarose gel and purified using the Sephaglass BandPrep Kit (Amersham Pharmacia Biotech). These products were cloned into the pCR 2.1 TOPO vector using the TOPO TA Cloning Kit version K2 (Invitrogen). Plasmids were purified using the Wizard Plus Minipreps DNA Purification System (Promega).

For construction of mosquito (host) phylogeny, we used a 987 bp region (excluding intron sequence) of Vg-C, a single copy yolk protein-encoding gene [30]. This region was amplified from Ae. simpsoni by nested PCR in our lab to add this species to the mosquito phylogeny. The following describes methods according to Isoe's work [30]. Degenerate primers were designed to amplify a 1.1 kb region that is specific for the Vg-C ortholog that includes the second intron. Primers Vg-C-specific forward (5’-(A/G)A(T/C)(A/G)TNA(A/G)CA(T/C)CCNA(A/G)G-3’), Vg-C-specific reverse (5’-TC(A/G)TT(T/C)TG(T/C)TT(A/G)TA(T/C)TG(A/G)TCC-3’), and Aeles universal reverse (5’-C(A/G)T(A/G)CC(A/G)CANTCNCCCAT-3’) were used in nested PCR. The first PCR used the Vg-C-specific forward and reverse primers for 1 cycle at 94 °C for 3 minutes, 32 cycles at 94 °C for 1 minute, 50 °C for 1.5 minute, and 1 extension cycle at 72 °C for 10 minutes. The second PCR used the Vg-C-specific and Aeles universal reverse primers with the same conditions except that the annealing temperature was increased to 54 °C. PCR products for Ae. simpsoni were cloned and sequenced as described above. Cloned inserts were sequenced in our laboratory using a GENE READIR DNA sequencer (LI-COR) with fluorescent-labeled T7 and m13r primers, or by DNA sequencing services (Ambiclon Express and VBI-Blacksburg, VA). H2O was used as a no-template negative control for PCR.

Genome and sequence analysis

Genome analysis was performed on the contig version of the Aeles aegypti genome sequence, which consists of 36206 contigs comprising 1310.1 Mb, having 7.6 x coverage (Broad Institute). BLAST and other programs (TEpost, FromTEpost) developed in our lab [44] were used to extract and filter sequences from BLAST output. Genome contribution by Juan-A was estimated using RepeatMasker [28] using 70% identity cutoff with full-length Juan-A as query. The Wisconsin Package GCG version 10.2-UNIX (Genetics Computer Group) was used for analysis of cloned and sequenced PCR products. Alignments were produced with ClustalX 1.81 [48]. To obtain ds/dN values, substitution analysis was performed using the SNAP program on the web [49,50]. Only sequences that had intact sequence regions were used for substitution analysis. Mean values are calculated based on all pair-wise comparisons from that group.

Phylogenetic inference

Phylogenetic inference was performed using MrBayes version 3.1.2 [51,52]. Sequences were aligned using ClustalX version 1.83 [53] using the following parameters: pairwise alignment gap opening = 10, gap extension = 0.1; multiple alignment gap opening = 10, gap extension = 0.2. Nucleotide Vg-C sequence data (see above) were used for the host phylogeny. The Modeltest server (version 3.7) [54,55] was used to determine the best nucleotide evolutionary model (General Time Reversible (GTR) allowing for variable substitution rates among sites) according to an Akaike Information Criteria (AIC) score. The model was implemented with MrBayes, running 150,000 generations, concluding with an average standard deviation of split frequencies below 0.01 (as suggested in the MrBayes manual), evidence of convergence of two independent tree searches.
Conceptually translated nucleotide sequences and sequences form Genbank (accessions) were used for non-LTR phylogeny. Sequences were aligned as described above. MrBayes was used to explore 9 fixed-rate amino acid evolutionary models, finding Jones [56] to have the highest score. Two hundred thousand generations were run resulting in an average standard deviation of split frequencies below 0.01. For all consensus trees displayed, clade credibility values are given at each node representing samplings of 1 of every 100th generation, while discarding the first 25% of all generations (the "burnin" period). Another analysis performed for 1,000,000 generations produced the same tree topology. See additional files 1 and 2 for alignments used for phylogenetic inference.

Juan-A copy number determination in the Ae. aegypti genome sequence

Different regions of the Juan-A sequence were used to determine Juan copy number in the Ae. aegypti genome by database search using BLAST (Figure 1, Table 1). The Juan-A 3′ UTR is approximately 240 bp. For copy number determination, we used 0.34 Kb of the 3′ end as the query to be consistent with the use of the 0.34 Kb 5′ UTR. Hits were counted which had sequence identities greater than or equal to 70%, 80%, 95%, 97%, 98%, or 99% compared to the query. In each case, a hit had to have at least 90% length of the query sequence. The full-length published Juan-A sequence is 4727 bp [24].

Sequence identity comparisons

In Table 3, values shown for all species except Ae. aegypti are means plus one standard deviation from pair-wise comparisons of nucleotide sequences obtained by PCR. Only sequences from the same lineages are compared. Comparisons were made between sequences and the consensus derived from the number of sequences in column 3. Ae. aegypti sequences were obtained by database search using a query that spans the same sequence amplified by PCR (see Figure 1). The number 768 is higher than what is shown in row 1 of Table 1 because the query here is the segment used for PCR.

Library screening

Amplified genomic libraries for Ae. albopictus, Ae. polynesiensis, C. tarsalis, and C. quinquefasciatus made using the Zap Express or Dash II kits (Stratagene) were screened using Digoxegenin-labeled (Roche Diagnostics) ssDNA probes generated from asymmetric PCR reactions. Two probes used for screening libraries of the Aedes or Culex genus were made from cloned PCR products amplified from Ae. aegypti and C. tarsalis using degenerate primers described above. The average insert size for the genomic libraries was 7 kb for Aedes and Culex libraries. Approximately 15,000 – 50,000 plaques were plated on NZY Agar plates and lifts were performed with nylon membranes (Osmonics). The membranes were blocked with prehybridization solution, containing 5 × SSC, 0.1% N-lauroylsarcosine, 0.02% SDS, and 2% nonfat milk for 2 hours at 55.0°C in a rotating hybridization incubator. Hybridization was performed with about 20 ng/ml of Digoxegenin-labeled probe in prehybridization solution for 6 hours to overnight at 55.0°C in a rotating hybridization incubator. Stringency washes were done using 0.5 × SSC, 0.1% SDS. Membranes were incubated with an anti-Digoxegenin antibody conjugated to alkaline phosphatase, and then developed with substrates BCIP and NBT for colorimetric detection. The copy number of Juan was calculated using known values of haploid genome size, average insert size of the library, and the ratio of positives to total number of plaques.

Authors’ contributions

ZT and JKB designed the study and drafted the manuscript. JKB performed sequence analysis and phylogenetic inference. All authors read and approved the final manuscript.

Additional material

Click here for file [http://www.biomedcentral.com/content/supplementary/1471-2148-7-112-S2.txt] Additional file 2
alignment used for phylogenetic inference (Figure 2B). Nexus format generated with Clustal
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