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*Aedes albopictus*

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**ABSTRACT**

Since its genome details are publically available, the mosquito *Aedes albopictus* has become the central stage of attention for deciphering multiple biological and evolutionary aspects at the root of its success as an invasive species. Its genome of 1.967 Mb harbours an unusual high number of non-retroviral integrated RNA virus sequences (NIRVS). NIRVS are enriched in piRNA clusters and produce piRNAs, suggesting an antiviral effect. Here, we investigated the evolutionary history of NIRVS in geographically distant *Ae. albopictus* populations by comparing genetic variation as derived by neutral microsatellite loci and seven selected NIRVS. We found that the evolution of NIRVS was far to be neutral with variations both in their distribution and sequence polymorphism among *Ae. albopictus* populations. The Flaviviral elements AlbFlavi2 and AlbFlavi36 were more deeply investigated in their association with dissemination rates of dengue virus (DENV) and chikungunya virus (CHIKV) in *Ae. albopictus* at both population and individual levels. Our results show a complex association between NIRVS and DENV/CHIKV opening a new avenue for investigating the functional role of NIRVS as antiviral elements shaping vector competence of mosquitoes to arboviruses.

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**KEYWORDS** *Aedes albopictus*; arboviral diseases; genetic structure; vector competence; NIRVS

**Introduction**

In less than four decades, the Asian tiger mosquito *Aedes albopictus* (Skuse, 1894) has become a public health concern owing to its ability to transmit several human pathogenic viruses such as Chikungunya virus (CHIKV), Dengue viruses (DENV) and Zika virus (ZIKV) [1]. This vector is responsible of several recent arboviral outbreaks (reviewed by [2]) and astonished by the speed at which it has conquered all continents except Antarctica [3,4], thus becoming one of the world’s 100-most invasive species according to the Global Invasive Species Database. Originally native to tropical forests of South-East Asia, *Ae. albopictus* was confined for centuries to few regions in Asia. Starting in the eighteenth or nineteenth centuries, *Ae. albopictus* was introduced in the islands of the Indian Ocean by Asian immigrants. From the late 1970s, this species took advantage of the increase in global trade to invade most tropical and sub-tropical regions of the world [5]. Additionally, the capacity of its eggs to undergo photoperiodic diapause in winter, favoured *Ae. albopictus* colonization into temperate regions [6]. *Aedes albopictus* was first reported in Europe in 1979 [7], in North America in 1985 [8], in South America in 1986 [9] and in Africa in 1990 [10].

Migratory routes used by *Ae. albopictus* to expand from its Asian cradle can be defined using population genetic approaches [11–13]. Mosquito populations form groups of interbreeding individuals, which coexist in space and time. These genetic units are interconnected through gene flow. The current worldwide distribution of *Ae. albopictus* is a direct consequence of increasing human activities with rapid irradiation of populations [14–18]. Populations established locally presented a high genetic variability evidencing a mixture of individuals of distinct origins with disparate susceptibilities to arboviruses, which are reflected by different vector competences [17]. Vector competence refers to the ability of a mosquito population to become infected after an infectious blood meal, to support viral...
replication, dissemination and transmission to a new host in a subsequent blood-meal [19]. The level of vector competence depends on the tripartite interactions among the mosquito genotype, the virus genotype and environmental factors under GxGxE interactions [20].

In the early 2000s, non-retroviral integrated RNA virus sequences (NIRVS) were discovered in different metazoans, including mosquitoes [21,22]. Aedes mosquitoes can host NIRVS originated from different viruses related to arboviruses: mainly insect-specific viruses (ISVs) including insect-specific flaviviruses (ISFs) and other viruses belonging to the Mononegavirales order (such as rhabdoviruses) [22–25]. Additionally, most NIRVS correspond to fragmented viral open-reading frames, are flanked by transposable elements (TEs), enriched in PIWI-interacting RNA (piRNA) clusters and produce piRNAs [29]. piRNA clusters are genomic regions composed of fragmented sequences of TEs which are expressed as long primary single-stranded RNAs and processed into fragments of 24–30 nucleotides called piRNAs. In the model organism Drosophila melanogaster, piRNAs are primarily produced in germline cells and target TE transcripts based on sequence complementarity to protect from heritable lesions [26–29]. The landscape of TE fragments within piRNA clusters defines the regulatory properties of D. melanogaster strains to TE invasion [30]. The analogy between TE fragments and viral sequences in piRNA supports the hypothesis that viral sequences may contribute to mosquito susceptibility to subsequent viral infections. If viral integrations contribute in controlling virus replication, with consequences on vector competence, positive selection should be expected [31]. On the contrary, if NIRVS stand for fossil records, these sequences should reach fixation and evolve at a neutral rate [32,33].

In this study, we selected 19 Ae. albopictus populations that cover the geographical distribution of the species, where CHIKV and DENV were circulating, to study the evolutionary dynamics of seven selected NIRVS [24]. The occurrence of NIRVS in populations was compared to processes driving population genetic differentiation observed on neutral loci (i.e. microsatellites). We showed that (i) based on microsatellite marker polymorphism, populations of Ae. albopictus are distributed into two different genetic clusters, one of them divided into four subclusters, without any correlation between genetic and geographical distances, (ii) the distribution of NIRVS in geographic populations and their polymorphism are not related to genetic divergences within and between populations as depicted by microsatellite markers, suggesting that NIRVS are not evolving neutrally, and (iii) all NIRVS studied, except AlbFlav1 may have influence on vector competence to DENV and CHIKV, at the population level.

Materials and methods

Ethic statements

Mice were housed in the Institut Pasteur animal facilities accredited by the French Ministry of Agriculture for performing experiments on live rodents. Work on animals was performed in compliance with French and European regulations on care and protection of laboratory animals (EC Directive 2010/63, French Law 2013-118, February 6th, 2013). All experiments were approved by the Ethics Committee #89 and registered under the reference APAFIS#6573-201606142077987 v2.

Mosquitoes

To assess genetic variation within and among Ae. albopictus populations, 19 samples from different geographic locations were studied (Table 1). These populations ranging from 10 to 30 mosquitoes were selected in the native range of Ae. albopictus where arboviral outbreaks and epidemics occurred [17,34,35]. Only the 13 populations of 19–20 mosquitoes and the Foshan colony used as control were selected to characterize NIRVS polymorphism [36].

Mosquitoes were collected in the field as immature stages (larvae, pupae, eggs). Frozen mosquitoes (<13th generation) were used for population genetic and NIRVS analyses. For the pilot experiments for DENV and CHIKV infections, F1 mosquitoes from Tibati population (Cameroon) and F18 mosquitoes from Foshan laboratory strain were used. Larvae obtained after immersion in dechlorinated tap water of field-collected eggs were distributed in pans of 200 individuals. Immature stages were fed every two days with a yeast tablet dissolved in 1 L of dechlorinated tap water at 26 ± 1°C. Emerging adults were placed in cages and maintained at 28 ± 1°C with a light/dark cycle of 12/12 h at 80% relative humidity and supplied with a 10% sucrose solution. Females were exposed three times a week to anesthetized mice (OF1 mice; Charles River Laboratories, MA, USA) as a source of blood for producing eggs.

Microsatellite genotyping

Genomic DNA was extracted from individual mosquito using the Nucleospin Tissue kit (Macherey-Nagel, Hoerdt, France) according to manufacturer’s instructions. Briefly, mosquitoes were individually homogenized in 180 µL lysis buffer supplemented with 25 µL of Proteinase K. To bind total nucleic acids, homogenates were passed through columns. Silica membranes were further desalted and DNA was collected in 100 µL of elution buffer. Quality and quantity of DNA were then assessed using the
Table 1. Details on Aedes albopictus populations analyzed.

| Population name | Continent | Country | City | Generation | Year of collection | Mosquitoes |
|-----------------|-----------|---------|------|------------|--------------------|------------|
|                 |           |         |      |            |                    | females    | Males |
| 1 Alessandria   | Europe    | Italy   | Alessandria | F2 | 2012 | 10 | 10 |
| 2 Ucinj         | Europe    | Montenegro | Ucinj | F1 | 2013 | 10 | 10 |
| 3 Cagnes-sur-mer | Europe | France | Cagnes-sur-mer | F13 | 2000 | 10 | 0 |
| 4 Montsecret    | Europe    | France | Montsecret | F4 | 2002 | 0 | 10 |
| 5 Bar-sur-Loup  | Europe    | France | Bar-sur-Loup | F1 | 2011 | 10 | 0 |
| 6 Tirana        | Europe    | Albania | Tirana | F6 | 2016 | 15 | 15 |
| 7 Franceville   | Africa    | Gabon   | Franceville | F2 | 2015 | 10 | 10 |
| 8 Mfliou        | Africa    | Congo   | Mfliou (Brazzaville) | F3 | 2012 | 10 | 10 |
| 9 Bertoua       | Africa    | Cameroon | Bertoua | F5 | 2008 | 10 | 4 |
| 10 Saint-Denis  | Africa    | La Réunion | Saint-Denis | F2 | 1998/2006 | 10 | 10 |
| 11 Rabat        | Africa    | Morocco | Rabat | F1 | 2017 | 15 | 15 |
| 12 Vero Beach   | America   | USA     | VeroBeach | F5 | 2016 | 10 | 10 |
| 13 Rio          | America   | Brazil   | Rio de Janeiro | F1 | 2001 | 10 | 9 |
| 14 Jurujuba     | America   | Brazil   | Jurujuba | F1 | 2014 | 10 | 0 |
| 15 Manaus       | America   | Brazil   | Manaus | F1 | 2015 | 15 | 15 |
| 16 PMNI         | America   | Brazil   | PMNI (Nova Iguacu) | F1 | 2015 | 15 | 15 |
| 17 Binh Duong   | Asia      | Vietnam | Binh Duong (Ben Cat) Phu Hoa | F9 | 2014 | 10 | 10 |
| 18 Sarba        | Asia      | Lebanon | Sarba | F0 | 2011 | 10 | 10 |
| 19 Foshan       | Asia      | China   | Foshan | Lab colony | – | 10 | 10 |
| 20 Oahu         | America   | Hawaii  | Oahu | Lab colony | 1999 | 10 | 10 |

Note: Except the colony Foshan, all 19 populations were genetically characterized using 10 microsatellite markers, and 13 populations (in bold) were selected for studying NIRVS diversity.

Nanodrop 2000 Spectrophotometer (Thermo Scientific™, MA, USA) and a PCR was performed using histone h3 reference gene (NCBI: XM_019696438.1) as control. Eleven microsatellite loci were amplified using PCR specific primers flanking the repeated region [16]. PCR reaction mixtures in a final volume of 15 µL contained 50 ng genomic DNA, 1X PCR buffer, 1.5 mM MgCl2, 0.27 mM dNTPs (Invitrogen™, CA, USA), 1U Taq polymerase (Invitrogen™), and 10 µM of each primer (one was 5′ labelled with a fluorescent dye). PCR cycling conditions consisted in a step at 94°C for 5 min, followed by 29 cycles at 94°C for 30 s, 58°C for 30 s, and 72°C for 30 s with a final step at 72°C for 10 min. Aliquots of PCR products were visualized in 2% agarose gels stained with ethidium bromide under UV light. Each PCR product was then diluted 1:10 in ddH2O water and 2 µL of this dilution was added to 10 µL of a mixture of deionized formamide and GeneScan-500 ROX size standard (Applied Biosystems, CA, USA). Genotyping was processed in an ABI3730XL sequence analyser (Applied Biosystems) and data analyzed using GeneScan and Genemapper softwares.

**Genetic diversity of populations**

For statistical analyses, mosquitoes collected from each sampling site were assumed to represent local populations.

Amounts of heterozygosity at various levels of population structure were explored by using $F_{IS}$ and $F_{ST}$ values. The $F_{IS}$ inbreeding coefficient indicates the level of heterozygosity within each population. The $F_{ST}$ fixation index measures the reduction in heterozygosis due to random genetic drift between populations. The Wright’s $F$-statistics were computed in Genetix v.4.05.2. software [37] and tested using $10^4$ iterations according to Weir and Cockerham (1984). The number of alleles ($N_A$), allelic richness ($A_r$), expected ($H_e$), observed ($H_o$) heterozygosis, $F_{IS}$ by locus and $F_{ST}$ by populations were obtained. The significance of $F_{IS}$ and $F_{ST}$ were analyzed using FSTAT v.2.9.3.2 [38].

Departures from Hardy-Weinberg equilibrium, linkage disequilibrium between loci and molecular variance (AMOVA) over all populations were analyzed with Arlequin v3.5.2.2 to estimate intra- and inter-population variation [39]. The mean frequency of null alleles (a mutation in microsatellite flanking regions leading to an absence of amplification products) per population was also very low (i.e. 0.06) and ranged from 0.000 to 0.118, meaning that the selected microsatellite loci were successfully amplified and appropriate for population genetic analysis. Relationships between geographical and genetic $F_{ST}$ distances were tested between populations using Mantel test implemented in GenAlEx v 6.5 [40].

**Graphic representation of relatedness among populations**

Genetic relationships between populations were estimated by using PHYLIP 3.69, as previously described [41]. Cavalli-Sforza & Edwards’s (CSE) chord distance for each pair of populations was calculated (GENDIST module). The resulting distance matrix was used to create a phylogram based on Neighbour-Joining (NEIGHBOUR module). Node confidence was inferred via 100 bootstrap replicates (modules SEQ-BOOT, GENDIST, NEIGHBOUR and CONSENSE).

**Test of isolation by distance**

To test the hypothesis whether the geographical pattern of genetic differentiation is caused by isolation
by distance (IBD), we ran Mantel tests for pairwise matrices between geographical distances (kilometres) and genetic differentiation (measured as $F_{ST}/(1−F_{ST})$).

**Genetic structure of populations**

Genetic population structure was assessed with individual-based Bayesian clustering method implemented in the program STRUCTURE v.2.3.4 [42]. The likelihood of each possible number of genetic populations (K), ranging from 1 to 20, was calculated after 10 independent runs for each value of K, using a burn-in of 500,000 replications, 500,000 Markov Chain Monte Carlo steps, assuming an admixture model, with frequencies correlated between populations and without the use of sampling location as a prior. The most likely number of populations (K) was estimated by the ΔK method described by Evanno et al. [43] with Structure Harvester software (http://taylor0.biology.ucla.edu/structureHarvester/). The results were then graphically displayed with DISTRUCT 1.1 [44].

**NIRVS in natural populations**

Six NIRVS (AlbFlavi1, AlbFlavi2, AlbFlavi4, AlbFlavi10, AlbFlavi36, AlbFlavi41), plus CSA [21], were studied. They were chosen based on their unique occurrence in different regions of the Foshan genome, except for AlbFlavi41, known to be duplicated. Because of its length [21], CSA was characterized using two sets of primers (CSA-NS3 and CSA-JJL), and considered in analysis as two separated datasets (Supplementary Table 1). NIRVS were searched on the same mosquitoes as those used for microsatellite genotyping.

PCR primers flanking each NIRVS (Supplementary Table 1) were designed using PRIMER3 [45]. PCR reactions were performed in a final volume of 25 μL using 5 ng of DNA, PCR buffer 1X, 2.9 mM MgCl$_2$, 0.25 mM dNTP mix, 0.25 μM of each primer and 1.25 unit of Taq DNA polymerase (Invitrogen™). Amplifications were done using T100™ Thermal Cycler (Bio-Rad) according to the following cycle conditions: 95°C for 5 min, 35 cycles at 95°C for 15 s, 59–64°C for 90 s, 72°C for 90 s, and a final elongation step at 72°C for 10 min. PCR products were electrophoresed on 1.5% agarose gels stained with ethidium bromide and visualized under UV light. All negative samples were confirmed with a second-round PCR.

The number of each NIRVS per population (frequency) was represented using Graph Pad Prism version 6 (Graph Pad Software, CA, USA).

**NIRVS between populations**

NIRVS composition of each population was expressed in terms of relative abundance corresponding to the percentage of each NIRVS relative to the total number of tested mosquitoes. To assess variations of NIRVS abundance between populations, we calculated Bray Curtis dissimilarities, a metrics, widely used, including data standardization [46]. A dendrogram was generated by using this dissimilarity matrix and neighbour-joining method, to visualize NIRVS compositional differences across populations [46].

**Mantel tests between NIRVS and microsatellites**

Mantel tests implemented in GenAlEx v 6.5 [40], were used to evaluate the statistical significance of the correlation between two or more distance matrices, using permutation tests. The significance of associations between matrices of $\phi$’s t distance among populations as derived by microsatellite markers or NIRVS distribution was tested using the Mantel test with 999 permutations.

**NIRVS association with vector competence of Ae. albopictus at the population level**

Vector competence is assessed using several parameters [47]. The dissemination efficiency (DE) refers to the proportion of mosquitoes able to disseminate the virus beyond the midgut barrier after ingestion of infectious blood meal and active viral replication in midgut epithelial cells. Published data on DEs to DENV and CHIKV were retrieved for mosquito populations with close or identical geographical proximity with the populations analyzed (Supplementary Table 2). DEs were described using median and inter-quartile range (IQR). Logistic linear regression models were used to test association between the presence of NIRVS and DEs at the Ae. albopictus population level. P-values <0.05 were considered significant.

**Sequence polymorphism and evolutionary dynamics of AlbFlavi2 and AlbFlavi36**

**Sequence polymorphism of AlbFlavi2 and AlbFlavi36**

Amplification products for AlbFlavi2 and AlbFlavi36 were purified by using NucleoSpin® Gel and PCR Clean-up kit (Macherey-Nagel) according to the manufacturer’s instructions and sequenced by Eurofins Genomics (Cochin hospital platform, Paris, France). If the quality of chromatogram profiles assessed with Geneious® 10.1.3 did not meet the standard required, amplicons were subsequently cloned into pCRII®-TOPO® vector using TOPO® TA Cloning® Kit (Thermo Fisher Scientific) and transformed into One Shot® TOP10 Chemically Competent E. coli (Invitrogen™). Sequences were aligned in Geneious® 10.1.3 using MUSCLE algorithm [48], p-distance values, representing proportion of nucleotide sites at which two
sequences being compared are different, were calculated after alignments using MEGA 10.0.5.

**Phylogeny based on AlbFlavi2 and AlbFlavi36**

DNA sequences from AlbFlavi2 and AlbFlavi36 were aligned with MUSCLE algorithm [48]. Exogenous virus sequences were used as outgroup to determine the direction of character transformations. Phylogenetic trees were obtained by parsimony analysis implemented in the PAUP* software package (version 4.0), by using gap as 5th state or not (data not shown) and nearest neighbour interchange (NNI) for tree rearrangements [49].

**Pilot experiment assessing the association between AlbFlavi2/AlbFlavi36 and vector competence of Ae. albopictus at the individual level**

Logistic linear regression models were used to test association between the presence of AlbFlavi2 and AlbFlavi36 and viral dissemination in single mosquitoes from the Foshan colony and a field-collected African population (Tibati collected in Tibati, Cameroon in 2018). Foshan and Tibati mosquitoes were experimentally infected with DENV (DENV-2, Accession number: MK268692 [50]) and CHIKV (06.21, Accession number: AM258992 [51]) provided in blood meals as described in Amraoui et al. 2019 [52]. All surviving mosquitoes were examined individually at 14 days post-infection to define (i) the infectious status; presence of infectious particles in heads was estimated by focus fluorescent assay on C6/36 Aedes albopictus cells [52], and (ii) the presence of AlbFlavi2 and/or AlbFlavi36 by PCR on DNA extracted from bodies and thorax. Statistical analyses were conducted using the Stata software (StataCorp LP, Texas, and USA). P-values <0.05 were considered significant.

**Results**

**Characterization of populations using microsatellite data**

A total of 363 individuals (Table 1) were genotyped at 11 microsatellite markers. Ten of the 11 microsatellite loci, by showing a frequency upper to 0.95 for the most common allele, were considered polymorphic and as such, were included in the analysis. Only the A17 locus was monomorphic and then excluded from the analyses. No significant linkage disequilibrium between any pairs of loci was observed indicating that the 10 loci were statistically independent from each other. Each locus was tested for within-population deviations from HWE implementing Bonferroni correction for multiple testing: 20 deviations out of 190 combinations were found but did not cluster at any population or locus.

The loci displayed a mean PIC of 0.59 suggesting that they were sufficiently informative for assessing the degree of variability and structuring of mosquito populations (Table 2). The number of alleles, averaged over all loci, ranged from 4 to 19, with a mean value of 10.9 per locus, and a mean allelic richness at 2.44 (Table 2). Eight of the 10 loci presented a significant difference between observed and expected heterozygosity, with a mean $F_{IS}$ of 0.17 (p-value <0.001; Table 2), supporting an excess of homozygotes in Ae. albopictus populations.

**Population diversity**

The overall mean number of alleles per population was 3.7 varying between 2.0 and 5.0. Private alleles were rarely found, their mean frequency per population ranged from 0.00 to 0.079 (Table 3). Mean $F_{IS}$ value per population was −0.09, with values ranging from −0.0285 to 0.338 (Table 3). Estimation of the molecular variance within and among populations (AMOVA) revealed that most of the variation (89.6%) was detected within individuals whereas only 10.4% occurred among populations.

**Population genetic structure**

The overall differentiation across all 19 populations was high ($F_{ST} = 0.239$) with $F_{ST}$ being highly significant (p-value <0.001), suggesting a significant genetic structure. To identify genetic clusters among tested individuals, we implemented 25 independent simulations in the software Structure according to the method of Evanno; the uppermost level of structuring in the model was observed at K = 2 (AK = 2293.4; Supplementary Figure 1(A)). The best assignment of individuals made by Structure led to two clusters: (i) cluster 1, which includes populations from Brazil (i.e. PMNI and Manaus), Northern Africa (i.e. Rabat) and a population from Albania, and (ii) cluster 2, which includes the remaining populations (Figure 1 and

| Table 2. Genetic diversity at each microsatellite locus for all mosquito populations. |
|---------------------------------------------------------------|
| Loci | $N_A$ | $N$ | Allelic richness | PIC | $H_o$ | $H_e$ | $F_{IS}$ |
|------|-------|-----|------------------|-----|-------|-------|----------|
| A1   | 4     | 288 | 0.44             | 0.16| 0.47  | 0.59  |          |
| A2   | 10    | 360 | 2.43             | 0.55| 0.63  | 0.62  | −0.15    |
| A3   | 18    | 341 | 3.14             | 0.74| 0.62  | 0.78  | 0.08     |
| A5   | 19    | 334 | 2.90             | 0.75| 0.52  | 0.78  | 0.13     |
| A6   | 11    | 330 | 2.59             | 0.68| 0.54  | 0.68  | −0.04    |
| A9   | 9     | 326 | 2.73             | 0.65| 0.59  | 0.70  | 0.01     |
| A11  | 9     | 323 | 2.81             | 0.71| 0.53  | 0.75  | 0.15     |
| A14  | 9     | 351 | 1.71             | 0.41| 0.17  | 0.40  | 0.38     |
| A15  | 6     | 347 | 1.25             | 0.13| 0.04  | 0.15  | 0.57     |
| A16  | 14    | 279 | 0.83             | 0.66| 0.84  | 0.04  |          |
| Mean | 10.9  | 327.9 | 2.44             | 0.59| 0.45  | 0.62  | 0.17     |

Notes: $N_A$, number of alleles; $N$, number of individuals examined at a locus; PIC, polymorphism information content; $H_o$, observed heterozygosity; $H_e$, expected heterozygosity; $F_{IS}$, the inbreeding coefficient. Values in bold are significant at the 0.1% after Bonferroni correction.
Table 3. Analysis of genetic variability of different geographical populations of Aedes albopictus.

| Populations          | Country  | Continent | N  | nA  | nA/N | np  | np/N | Hs  | Hs/M | Afs  | Fis  |
|----------------------|----------|-----------|----|-----|-------|-----|------|-----|------|------|------|
| Vero Beach           | USA      | America   | 20 | 4.4 | 0.22  | 2   | 0.10 | 0.07| 0.048| 0.33 | 0.08 | 0.12***|
| Oahu                 | Hawaii   |           | 20 | 2.6 | 0.13  | 0   | 0.00 | 0.00| 0.25 | 0.34 | 0.08 | 0.30***|
| Manaus               | Brazil   |           | 30 | 3.4 | 0.11  | 3   | 0.10 | 0.07| 0.32 | 0.47 | 0.12 | 0.34***|
| PMNI                 |          |           | 30 | 3.6 | 0.12  | 1   | 0.03 | 0.03| 0.39 | 0.51 | 0.11 | 0.25***|
| Rio                  |          |           | 19 | 3.4 | 0.18  | 0   | 0.00 | 0.00| 0.42 | 0.50 | 0.08 | 0.18***|
| Jurujuba             |          |           | 10 | 3.3 | 0.33  | 0   | 0.00 | 0.00| 0.47 | 0.42 | 0.00 | 0.06   |
| Montsecret           | France   | Europe    | 10 | 2.8 | 0.28  | 0.3 | 0.10 | 0.06| 0.35 | 0.38 | 0.01 | 0.014  |
| Cagnes-sur-Mer       | France   | Europe    | 10 | 3.8 | 0.38  | 2   | 0.20 | 0.08| 0.50 | 0.52 | 0.05 | 0.13*  |
| Alessandria          | Italy    |           | 20 | 4.1 | 0.21  | 2   | 0.10 | 0.05| 0.43 | 0.50 | 0.06 | 0.18***|
| Ulcinj               | Montenegro|         | 20 | 3.1 | 0.16  | 0   | 0.00 | 0.00| 0.40 | 0.39 | 0.03 | 0.02   |
| Cagnes-sur-Mer       | France   | Europe    | 10 | 3.8 | 0.38  | 2   | 0.20 | 0.08| 0.50 | 0.52 | 0.05 | 0.13***|
| Rabat                | Morocco  |           | 30 | 3.2 | 0.11  | 0   | 0.00 | 0.00| 0.42 | 0.47 | 0.06 | 0.11***|
| Bar-sur-Loup         | France   | Europe    | 10 | 2.8 | 0.13  | 3   | 0.10 | 0.07| 0.35 | 0.35 | 0.04 | 0.08   |
| Bertoua              | Cameroon |           | 14 | 4.3 | 0.31  | 1   | 0.07 | 0.07| 0.46 | 0.52 | 0.08 | 0.16***|
| Franceville          | Gabon    | Africa    | 20 | 4.8 | 0.24  | 1   | 0.05 | 0.03| 0.44 | 0.36 | 0.08 | 0.23***|
| Mfilou               | Congo    |           | 20 | 4.0 | 0.20  | 0   | 0.00 | 0.00| 0.60 | 0.55 | 0.02 | 0.06   |
| Saint-Denis          | La Réunion|        | 20 | 5.1 | 0.26  | 6   | 0.30 | 0.05| 0.52 | 0.55 | 0.07 | 0.08   |
| Sarba                | Lebanon  | Asia      | 10 | 3.5 | 0.35  | 4   | 0.40 | 0.06| 0.55 | 0.44 | 0.00 | 0.18   |
| Binh Duong           | Vietnam  |           | 20 | 5.0 | 0.25  | 3   | 0.15 | 0.08| 0.32 | 0.41 | 0.05 | 0.11***|
| Mean                 |          |           | 19.1| 3.7 | 0.022 | 1.5 | 0.09 | 0.04| 0.45 | 0.47 | 0.06 | 0.09   |

Notes: N, population size; nA, mean number of alleles; nA/N, mean number of alleles/individual; np, number of private alleles; np/N, mean number of private alleles/individual; Afs, mean frequency of private alleles; Hs, mean observed heterozygosity; Hs/M, mean expected heterozygosity; Afs, mean frequency of null alleles; Fis, the inbreeding coefficient.

**p-value ≤ 0.01; ***p-value ≤ 0.001.

Supplementary Table 1(B)). Although Structure identified only two clusters, their genetic differentiation was significantly high ($F_{ST} = 0.12$, $p$-value ≤ 0.001). Additionally, these two genetic clusters were found to deviate significantly ($p$-value ≤ 0.01) from Hardy-Weinberg proportions, with $F_{IS}$ values of 0.23 and 0.36, respectively after implementing 10^4 permutations on allele frequencies in the software GENETIX. This result suggested a Wahlund effect which indicated a genetic substructure. As a consequence, we re-analyzed the two clusters separately and detected substructures in Cluster 2 (Figure 1 and Supplementary Figure 1 (C)). Sub-structuring of Cluster 2 emphasized differentiation among populations from South America (Juru-juba and Rio), Europe (i.e. all French and Italian populations), Africa (Bertoua and Mfilou) and Asia. On the contrary, populations from Florida (Vero Beach), Middle East (Sarba), La Réunion Island (Saint Denis) and Gabon (Franceville) appeared genetically mixed suggesting that these sites may represent cross-roads in Aedes albopictus colonization out of Asia (Figure 1).

Spatial and genetic data

When considering the genetic differentiation according to geographical distances using Mantel test, only 21% of the genetic variability was explained by the geographical distance ($p$-value = 0.03).

NIRVS analysis

We further chose to study NIRVS distribution in 13 Aedes albopictus populations selected previously for assessing genetic diversity based on epidemiological data of arboviral outbreaks and Aedes albopictus widespread out if its native range [17,34,35]; we analyzed 10 females and 10 males per population (Table 1). NIRVS distribution was not homogenous across populations (Figure 2, Supplementary Figure 2). AlbFlavi1 (Figure 2(A)) and CSA-JJL (Figure 2(H)) were the most widespread NIRVS, being detected in 85–100% of tested mosquitoes, respectively. On the opposite, AlbFlavi10 was the rarest NIRVS, being found in 17% of the tested mosquitoes. Despite displaying the highest presence of AlbFlavi2 and AlbFlavi4 (present in most populations), Brazilian populations showed the lowest number of NIRVS, with the absence of AlbFlavi36 and AlbFlavi10 and the lower frequency of AlbFlavi41 compared to the other Aedes albopictus populations (Figure 2).

Interestingly, when targeting the CSA locus, no amplification was obtained for Brazilian mosquitoes using the NS3-CSA primer set (Supplementary Table 1; Figure 2(G)), which was not consistent with the results obtained with the JIL-CSA primer set that amplified the same NIRVS but targeting a different region (Figure 2(H)). This suggests then that recombination events occurred at this genomic position for Manaus, Rio and PMNI mosquitoes.

Comparison between microsatellite data and NIRVS abundance profiles

To define whether NIRVS evolve under selective pressures or randomly, we compared (i) the relatedness between populations based on microsatellite polymorphism (Figure 3(A)) with (ii) the similarity between populations based on NIRVS abundance profiles (Figure 3(B)). At the population level, the Neighbour-Joining clustering analyses applied to genetic frequencies (Figures 1 and 3(A), Supplementary
Figure 1(B) mainly revealed two clusters of populations: one with Tirana, Rabat, Manaus and PMNI populations, and the second with the remaining populations (i.e. Binh Duong, Oahu, Vero Beach, Rio, Franceville, Alessandria, Ulcinj and Mflou). Regarding NIRVS contents, two major clusters distantly related from each other were also obtained (Figure 3(B)), one including only Brazilian populations (i.e. Manaus, PMNI and Rio) sharing a low abundance of NIRVS, and the other including geographically distant populations. Therefore, populations from Vero Beach and Rio shared closely related genetic relationships and harboured clearly different abundance profiles of NIRVS (Figure 3(A,B)). Conversely, genetic distinct populations from Vero Beach, Rabat and Alessandria shared the same contents of NIRVS. In short, closely related populations can have different NIRVS contents and geographic and/or genetic distant populations can contain similar NIRVS abundance profiles. Thus, we observed significantly different relatedness between populations according to the marker used, that may indicate that random genetic drift was not the main force shaping these NIRVS distribution. In addition, no correlation was identified between the two matrices \( R^2 = 0.0006 \), showing that NIRVS composition of populations was not related to the genetic structure caused by random genetic drift (Supplementary Figure 2).

### Relationships between NIRVS landscape and vector competence (from published data)

Our results showed that NIRVS were not neutrally distributed across populations supporting the hypothesis of a biological function of NIRVS, such as antiviral functions, as it has been previously suggested. To analyze whether NIRVS contribute to the control of arboviral replication, we used logistic regression models to evaluate potential association between NIRVS distribution frequencies among populations and dissemination efficiencies (DEs) of DENV and CHIKV in corresponding or geographically close populations selected from the literature (Supplementary Table 2). On this basis, we classified our tested populations depending on their frequencies of each NIRVS using the median. Several associations between NIRVS and DENV DEs were found. Whereas high AlbFlavi2 and CSA-JJL frequencies were significantly associated with high DEs, high AlbFlavi10, AlbFlavi36, AlbFlavi41 and CSA-NS3 frequencies were however correlated to low DENV DEs (Table 4). Opposite associations to CHIKV DEs were also observed but with fewer NIRVS. Indeed, high frequencies of AlbFlavi4 were associated with high CHIKV DEs while high distribution of AlbFlavi36 and CSA-NS3 among populations was correlated to low CHIKV DEs. Together, these results indicated...
that the presence of NIRVS was associated with changes in DEs of arboviruses in *Ae. albopictus* at the population level.

Focus on *AlbFlavi2 and AlbFlavi36*

Besides its association to high DENV DEs, we further focused on *AlbFlavi2* located in a piRNA cluster and...
presenting a large geographical distribution encompassing all continents. Moreover, AlbFlavi36 located in an intergenic region was not found in populations from South America and presented an opposite pattern of association to DENV DEs; it was used for comparison to AlbFlavi2. Therefore, sequence polymorphism, evolution and potential association to vector competence to arboviruses were assessed in Ae. albopictus at the individual level.

### Table 4. Association between NIRVS and arboviral dissemination efficiencies in Ae. albopictus populations (logistic regression models).

| NIRVS | N | OR (95% CI) | p-value |
|-------|---|-------------|---------|
| DENV  |    |             |         |
| AlbFlavi1 | 707 | 0.82 (0.61–1.11) | NS |
| AlbFlavi2 | 3.07 (2.3–4.2) | *** |
| AlbFlavi4 | 1.46 (1.0–2.1) | NS |
| AlbFlavi10 | 0.46 (0.3–0.6) | *** |
| AlbFlavi36 | 0.46 (0.3–0.6) | *** |
| AlbFlavi1 | 0.43 (0.29–0.62) | *** |
| CSA-NS3 | 0.68 (0.5–0.9) | ** |
| CSA-JIL | 3.87 (1.0–14) | *** |
| CSA-JI | 0.56 (0.4–1.0) | NS |
| CSA-JH | 0.56 (0.3–1.0) | NS |
| CSA-JI | 0.68 (0.3–1.7) | NS |

Notes: Dissemination efficiencies (DE) data were assessed from the same or geographically close Ae. albopictus populations (see Supplementary Table 2). Populations were characterized as high or low frequencies for each NIRVS by using the median and were analyzed with DE data by logistic regression models to find any association. Odds ratio (OR) >1 and <1 indicated positive and negative association respectively. 95% CI: 95% confidence intervals; N: sample size.

Sequence polymorphism and evolution of AlbFlavi2 and AlbFlavi36

AlbFlavi2 and AlbFlavi36 of individual mosquitoes were successfully amplified in 10 out of the 13 Ae. albopictus populations studied (Figure 2). Sequences of AlbFlavi2 and AlbFlavi36 amplified fragments were aligned with the closely related exogenous viral sequences: the Kamiti River virus (KRV) for AlbFlavi2 and the Aedes Flavivirus (AeFV) for AlbFlavi36.

The p-distance calculated between AlbFlavi2 sequences revealed a moderate diversity, with values ranging from 0% to 3.9% (data not shown). All AlbFlavi2 sequences underwent deletion events that were sometimes shared by several individuals from different populations or observed in a specific Ae. albopictus population (Figure 4; Supplementary AlbFlavi36).

The p-distance values calculated between AlbFlavi36 sequences were low and varied only from 0% to 0.9% (data not shown). Therefore, AlbFlavi36 sequences mapping to an intergenic region [24] appeared more similar between individuals than AlbFlavi2 sequences (which are integrated in piRNA cluster 27 [53]).
We further performed phylogenetic analysis to describe the evolutionary history of AlbFlavi2 and AlbFlavi36 (Figure 4; Supplementary Figure 4). As expected, AlbFlavi2 displayed higher divergence than AlbFlavi36. The resulted trees based on AlbFlavi2 supported four major clusters sharing sequences with the Aedes albopictus individuals. Phylogram of AlbFlavi2 sequences based on parsimony with gaps considered as 5th nucleotides. Each node was found in 98–100% of the trees obtained through NNI rearrangements. Significant bootstrap values were indicated at nodes. hmwb: high molecular weight band Values; in brackets: alignment coordinates of deletion. The same result was obtained by parsimony without gap as 5th nucleotide, except for the sequence cluster of mosquitoes from Morocco.

Figure 4. Divergence of AlbFlavi2 among Aedes albopictus individuals. Phylogram of AlbFlavi2 sequences based on parsimony with gaps considered as 5th nucleotides. Each node was found in 98–100% of the trees obtained through NNI rearrangements. Significant bootstrap values were indicated at nodes. hmwb: high molecular weight band Values; in brackets: alignment coordinates of deletion. The same result was obtained by parsimony without gap as 5th nucleotide, except for the sequence cluster of mosquitoes from Morocco.

We further performed phylogenetic analysis to describe the evolutionary history of AlbFlavi2 and AlbFlavi36 (Figure 4; Supplementary Figure 4). As expected, AlbFlavi2 displayed higher divergence than AlbFlavi36. The resulted trees based on AlbFlavi2 supported four major clusters sharing sequences with the
same deletion events (Figure 4). One of them, defined by a 11 bp deletion was subdivided into two subclusters of sequences sharing respectively a 248 bp and a 46 bp deletion. Overall, these clusters did not show any strict relationship with geographical origin of populations.

AlbFlavi2 from Brazilian populations (PMNI, Manaus and Rio) appeared clearly polyphyletic. Three clusters represented only sequences from Brazil: an older cluster of sequences sharing a 405 bp insertion and two other clusters of sequences sharing respectively a 46 bp and a 6–65 bp deletions. Only AlbFlavi2 from Rabat population formed a monophyletic group, a unique cluster of sequences characterized by a 248 bp deletion.

One cluster contained most of the sequences from Tirana (Albania), Vero Beach (USA), Alessandria (Italy) and Binh Duong (Vietnam). However, some sequences from Binh Duong, Vero Beach and Alessandria populations, were also isolated or found in other clusters. Moreover, half of the AlbFlavi2 sequences from Foshan formed a unique cluster whereas the other half shared the heterogeneous cluster of sequences described above.

The polymorphism of AlbFlavi36 was low in *Ae. albopictus* populations and all the sequences showed close relationships with each other, without any significant bootstrap values (Supplementary Figure 4). However, sequences from African mosquitoes (Congo, Gabon and Morocco) in branching deeply in the tree may be more ancient than sequences of Asian (Binh Duong, Foshan) and European mosquitoes.

Collectively, phylogenetic studies revealed different histories of NIRVS in mosquito populations and particularly, complex history of Albflavi2 evolving by both mutation and deletion events.

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**Figure 5.** Pilot analysis showing the association between frequencies of AlbFlavi2/AlbFlavi36 and DENV/CHIKV dissemination efficiencies (DE) in *Aedes albopictus* populations. The Foshan colony and the Tibati population (Cameroon, generation F1) were used for the analysis. (A) Presence/absence of AlbFlavi2 and DEs to DENV and CHIKV obtained for the Foshan colony. (B) Presence/absence of AlbFlavi36 and DEs to DENV and CHIKV obtained for the Foshan colony. (C) Presence/absence of AlbFlavi2 and DEs to DENV and CHIKV obtained for the Tibati population. (D) Presence/absence of AlbFlavi36 and DEs to DENV and CHIKV obtained for the Tibati population. DEs were obtained for both viruses at 14 days post-infection. In total, 191 and 122 individuals were examined for presence of AlbFlavi2/AlbFlavi36 after infection DENV and CHIKV, respectively. Interactions of populations and frequencies of AlbFlavi2/AlbFlavi36 with DEs were tested using logistic regression models.
Association between AlbFlavi2/AlbFlavi36 and vector competence in *Ae. albopictus* at the individual level

To further investigate the biological role of AlbFlavi2 and AlbFlavi36, we extended our analyses of association between the presence of these two NIRVS and viral dissemination from a population level to an individual level. To do so, mosquitoes of the Foshan colony strain and the field-collected Tibati population (Cameroon) were infected with DENV-2 and CHIKV. At 14 days post-infection, mosquito dissemination status and the presence of both AlbFlavi2 and AlbFlavi36 were determined for 313 mosquitoes (Figure 5). When comparing DENV dissemination, Foshan better disseminated than Tibati (respectively 89.8% and 42.2%; p-value <10^{-4}; data not shown). However, the presence of AlbFlavi2 or AlbFlavi36 in mosquitoes from Foshan or Tibati was not significantly associated to DENV dissemination (p-values > 0.209; Figure 5(A–D)). Moreover, the same results were obtained regarding the association between AlbFlavi2/AlbFlavi36 presence and CHIKV dissemination (p-values > 0.3; Figure 5(A–D)). In all, using both laboratory colony and field-collected mosquitoes, no association was found between any of the two NIRVS (AlbFlavi2 and AlbFlavi36) and DENV/CHIKV dissemination.

Discussion

The presence of sequences with similarities to flaviviruses in the genome of *Aedes* spp. mosquitoes and their enrichment in piRNA clusters support the hypothesis that viral integrations, at least some, could have a biological role [24,25,54]. Here we demonstrated that the NIRVS studied are not neutrally distributed among *Ae. albopictus* populations and all of them (except AlbFlavi1) were significantly associated to vector competence to DENV and CHIKV.

NIRVS are not neutral markers

NIRVS are endogenous viral sequences located in protein-coding gene exons, intergenic regions, and PIWI-interacting RNA (piRNA) clusters [55]. Contrary to the other categories, piRNA genes are evolving rapidly under positive selection to generate a high diversity of piRNAs [56–63]. Approximately 30 sequences of flavivirus ORFs (primarily NS1 and NS5) were detected in the Foshan colony [24] including the early-detected Flavivirus-like sequences [21,64]. It has been suggested that the variable number and frequency of NIRVS across *Ae. albopictus* populations contribute to variations of genome size among mosquitoes [36]. Our study targeted seven NIRVS (AlbFlavi1, AlbFlavi2, AlbFlavi4, AlbFlavi10, AlbFlavi36, AlbFlavi41, CSA), for which their evolution under selection pressures or simply by genetic drift remained unknown.

Contrary to NIRVS deriving from flaviviruses in *Ae. aegypti* (unpublished data), the presence of the tested NIRVS was highly variable among worldwide populations, with the lowest number being detected in mosquitoes from Brazil. This suggests that these NIRVS have not reached fixation in any of the *Ae. albopictus* populations tested (with potential exception for AlbFlavi1 and a fragment of CSA). Interestingly, Brazilian populations (Manaus, Rio, and PMNI) displayed the lowest number of NIRVS studied with a total absence of AlbFlavi10, AlbFlavi36 and CSA-NS3. Considering the ancient origin of NIRVS, estimated between 6.5 thousands to 2.5 million years ago [65], we hypothesized that they might have been less exposed to ISFs in the past, leading to fewer NIRVS in their genomes or that populations may have acquired NIRVS in the past which have been lost over time.

We showed that the microsatellite polymorphism of populations did not match with the NIRVS abundance profiles. Therefore, because microsatellites are sequences that neutrally evolve in the genome, we speculate that the evolution of NIRVS is far from neutral and NIRVS could provide benefits to the host. NIRVS may have been produced for specific purposes in the host rather than being the consequences of random endogenization of exogenous viral fragments.

The focus on NIRVS from piRNA cluster (AlbFlavi2) and intergenic region (AlbFlavi36) revealed different sequence polymorphism and evolutionary histories despite their relatively wide distribution among *Ae. albopictus* populations. Whereas AlbFlavi36 appeared monophyletic with highly conserved sequences among populations, the phylogenetic analyses showed a particularly complex history for AlbFlavi2 evolving by both mutation and deletion events. We postulate that owing to its high variability, AlbFlavi2 may act similarly to TE fragments and piRNA genes, and evolve according to exposures to related exogenous virus burden.

Association between NIRVS and vector competence to arboviruses

Similar to some endogenous retroviruses (ERVs) that have an effect on viruses from different subfamilies and genera [66], we questioned whether NIRVS deriving from ISF sequences affect the dissemination of different arboviruses and contribute to the regulation of vector competence of *Ae. albopictus* mosquitoes. Because NIRVS located in piRNA clusters, such as AlbFlavi2, produce piRNAs, are differentially distributed among populations and appeared older integrations than NIRVS located in codons [65], they were proposed to function as novel mosquito antiviral immune factors. However, contradictory results were obtained using both population- and individual-level analysis.
At the population level, many NIRVS were present in Ae. albopictus populations. The frequency of some of them appears correlated (positively or negatively) to DENV dissemination. Whereas some NIRVS were associated with high viral dissemination (i.e. AlbFlavi2 and CSA-JJL), others were significantly related to low viral dissemination (AlbFlavi10, AlbFlavi36, AlbFlavi41 and CSA-NS3), suggesting different functions of NIRVS deriving from ISFs. Moreover, fewer NIRVS were positively (i.e. AlbFlavi4) and negatively (i.e. AlbFlavi36 and CSA-NS3) correlated to CHIKV dissemination. These results are consistent with the contradictory results obtained in studies assessing the impact of exogenous ISFs on arbovirus fitness in mosquitoes. Indeed, whereas some ISFs have shown to repress arboviral replication [67], others have been proved to facilitate infection [68]. Further experiments should be performed to confirm these results, as only 20 mosquitoes per population were tested for NIRVS distribution.

At the level of individuals, our pilot experiment based on mosquitoes from both laboratory colony and field did not show any significant association between dissemination of DENV and CHIKV, and NIRVS, AlbFlavi2 or AlbFlavi36. while this does not seem surprising for the alphavirus CHIKV considering that NIRVS examined are homologous to flavivirus sequences, it is still questionable for DENV. Since it appears that they do not evolve neutrally and were preserved from purifying selection, NIRVS such as CSA has demonstrated to produce a transcript of 4671 nt long [21,69]; its functional role should be investigated. Lastly, other components should be considered such as the virome [70] in addition to the anatomical barriers in the mosquito such as the salivary glands for viral transmission [19].

To conclude, our results clearly show that NIRVS in Ae. albopictus follow processes different from that of neutral genes such as microsatellites and most NIRVS are far from reaching fixation. Flaviviral integrations are differentially distributed among Ae. albopictus populations and are here suggested to be associated with the vector competence to arboviruses in mosquitoes that remain to be elucidated. Finally, this study opens the way to new perspectives on evolution and biological functions of NIRVS, in part on vector competence.

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Author contributions

ABF and XDL designed the experiments; VH performed the research; GG did the microsatellite genotyping; CD did the phylogenetic reconstruction; MS and MM did the analysis of genetic structure; YD did the statistical analysis; LM and PSY provided a technical help; MB edited the paper; VH, CD, ABF wrote the paper.

Disclosure statement

No potential conflict of interest was reported by the authors.

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References

[1] Paupy C, Delatte H, Bagny L, et al. Aedes albopictus, an arbovirus vector: from the darkness to the light. Microbes Infect. 2009;11:1177–1185. doi:10.1016/j.micinf.2009.05.005.
[2] Bonizzoni M, Gasperi G, Chen X, et al. The invasive mosquito species Aedes albopictus: current knowledge and future perspectives. Trends Parasitol. 2013;29:460–468. doi:10.1016/j.pt.2013.07.003.
[3] Caminade C, Medlock JM, Ducheyne E, et al. Suitability of European climate for the Asian tiger mosquito Aedes albopictus: recent trends and future scenarios. J R Soc Interface. 2012;9:2708–2717. doi:10.1098/rsif.2012.0138.
[4] Benedict MQ, Levine RS, Hawley WA, et al. Spread of the tiger: global risk of invasion by the mosquito Aedes albopictus. Vector Borne Zoonotic Dis. 2007;7:76–85. doi:10.1089/vbz.2006.0562.
[5] Loumibos LP. Invasions by insect vectors of human disease. Annu Rev Entomol. 2002;47:233–266. doi:10.1146/annurev.ento.47.091201.145206.
[6] Hawley WA. The biology of Aedes albopictus. J Am Mosq Control Assoc Suppl. 1988;1:1–39.
[7] Adhami J, Reiter P. Introduction and establishment of Aedes (Stegomyia) albopictus skuse (Diptera: Culicidae) in Albania. J Am Mosq Control Assoc. 1998;14:340–343.
[8] Sprenger D, Wuithiranyagool T. The discovery and distribution of Aedes albopictus in Harris County, Texas. J Am Mosq Control Assoc. 1986;2:217–219.
[9] Consoli RAGB OR. Principais mosquitos de importância sanitária no Brasil. Rio de Janeiro. Editora FIOCRUZ. 1994;228.
[10] Cornet AJ, Hunt RH. *Aedes albopictus* in Africa? First evidence of live vectors in imported tires in Cape Town. J Am Mosq Control Assoc. 1991;7:107–108.

[11] Goubert C, Minard G, Vieira C, et al. Population genetics of the Asian tiger mosquito *Aedes albopictus*, an invasive vector of human diseases. Heredity (Edinb). 2016;117:125–134. doi:10.1038/hdy.2016.35.

[12] Jackson H, Strubbe D, Tollington S, et al. Ancestral origins and invasion pathways in a globally invasive bird correlate with climate and influences from bird trade. Mol Ecol. 2015;24:4269–4285. doi:10.1111/mec.13307.

[13] Powell JR, Tabachnick WJ. History of domestication and spread of Aedes aegypti—a review. Mem Inst Oswaldo Cruz. 2013;108(Suppl 1):11–17. doi:10.1590/0070-247120130395.

[14] Maynard AJ, Ambrose L, Cooper RD, et al. Tiger on the prowl: invasion history and spatio-temporal genetic structure of the Asian tiger mosquito *Aedes albopictus* (Skuse 1894) in the Indo-Pacific. PLoS Negl Trop Dis. 2017;11:e0005546. doi:10.1371/journal.pntd.0005546.

[15] Kotsakiozi P, Richardson JB, Pichler V, et al. Molecular markers for analyses of intraspecific genetic diversity in the Asian tiger mosquito, *Aedes albopictus*. Parasit Vectors. 2011;4:188. doi:10.1186/1756-3305-4-188.

[16] Manni M, Gomulski LM, Aketarawong N, et al. Comparative genomics shows that viral integrations in vector genomes correlate with climate and influence from bird trade. Mol Ecol. 2012;21:381–390. doi:10.1111/j.1365-294X.2011.05332.X.

[17] Manni M, Guglielmino CR, Scolari F, et al. Sequences of piRNAs in the arboviral vector, *Aedes albopictus*. Mol Biochem. 2016;54:333–344. doi:10.1007/s11033-016-3334-7.

[18] Musso D, Rodriguez-Morales AJ, Levi JE, et al. ORF10, a novel endogenous virus-like protein in *Aedes aegypti*. PLoS Negl Trop Dis. 2017;11:e0005749. doi:10.1371/journal.pntd.0005749.

[19] Kramer LD, Ebel GD. Advances in virus research. Adv Virus Res. 2003;60:187–232.

[20] Zouache K, Fontaine A, Vega-Rúa A, et al. Three-way interactions between mosquito populations, viral strain and temperature underlying chikungunya virus transmission potential. Proc Biol Sci. 2014;380:281. doi:10.1098/rspb.2014.1078.

[21] Crochu S, et al. Sequences of flavivirus-related RNA viruses persist in DNA form integrated in the genome of *Aedes* spp. mosquitoes. J Gen Virol. 2004;85:1971–1980. doi:10.1099/vir.0.79850-0.

[22] Katzourakis A, Gifford RJ. Endogenous viral elements in animal genomes. PLoS Genet. 2010;6:e1001191. doi:10.1371/journal.pgen.1001191.

[23] Fort P, Albertini A, Van-Hua A, et al. Fossil rhodovibrio sequences integrated into arthropod genomes: ontogeny, evolution, and potential functionality. Mol Biol Evol. 2012;29:381–390. doi:10.1093/molbev/msr226.

[24] Palatini U, Miesen P, Carballar-Lejarazu R, et al. Comparative genomics shows that viral integrations are abundant and express piRNAs in the arboviral vectors *Aedes aegypti* and *Aedes albopictus*. BMC Genomics. 2017;18:312. doi:10.1186/s12864-017-3903-3.

[25] Whitfield ZJ, Dolan PT, Kunimori M, et al. The diversity, structure, and function of heritable adaptive immunity sequences in the *Aedes aegypti* genome. Curr Biol. 2017;27:3511–3519. doi:10.1016/j.cub.2017.09.067. e3517.

[26] Brennecke J, Aravin AA, Stark A, et al. Discrete small RNA-generating loci as master regulators of transposon activity in *Drosophila*. Cell. 2007;128:1089–1103. doi:10.1016/j.cell.2007.01.043.

[27] Pelisson A, Sarot E, Payen-Groschene G, et al. A novel repeat-associated small interfering RNA-mediated silencing pathway downregulates complementary sense gypsy transcripts in somatic cells of the *Drosophila* ovary. J Virol. 2007;81:1951–1960. doi:10.1128/JVI.01980-06.

[28] Saito K, Nishida KM, Mori T, et al. Specific association of Pwii with rasiRNAs derived from retrotransposon and heterochromatic regions in the *Drosophila* genome. Genes Dev. 2006;20:2214–2222. doi:10.1101/gad.1454806.

[29] Vagin VV, Sigova A, Li C, et al. A distinct small RNA pathway silences selfish genetic elements in the germ-line. Science. 2006;313:320–324. doi:10.1126/science.1129333.

[30] Zanni V, Eymery A, Coiffet M, et al. Distribution, evolution, and diversity of retrotransposons at the flmenco locus reflect the regulatory properties of piRNA clusters. Proc Natl Acad Sci U S A. 2013;110:19842–19847. doi:10.1073/pnas.1313677110.

[31] Frank JA, Frischotte C. Co-option of endogenous viral sequences for host cell function.Curr Opin Virol. 2017;25:81–89. doi:10.1016/j.coviro.2017.07.021.

[32] Aswad A, Katzourakis A. Paleovirology and virally derived immunity. Trends Ecol Evol. 2012;27:627–636. doi:10.1016/j.tree.2012.07.007.

[33] Katzourakis A. Paleovirology: inferring viral evolution from host genome sequence data. Philos Trans R Soc Lond B Biol Sci. 2013;368:20120493. doi:10.1098/rstb.2012.0493.

[34] Mosso D, Rodriguez-Morales AJ, Levi JE, et al. Unexpected outbreaks of arbovirus infections: lessons learned from the Pacific and tropical America. Lancet Infect Dis. 2018;18:e355–e361. doi:10.1016/S1473-3099(18)30269-X.

[35] Fritzell C, Rouset D, Adde A, et al. Current challenges and implications for dengue, chikungunya and Zika seroprevalence studies worldwide: A scoping review. PLoS Negl Trop Dis. 2018;12:e0006533. doi:10.1371/journal.pntd.0006533.

[36] Chen XG, Jiang X, Gu J, et al. Genome sequence of the arbovirus vector, *Aedes albopictus*. Mem Inst Oswaldo Cruz. 2015;110:19842–19847. doi:10.1590/M130195. doi:10.1037/134. doi:10.1590/0005-332. doi:10.1101/gad.1454806.

[37] Genetix 4.05 Logiciel Sous Windows TM Pour la Génétique des Populations. 1996–2004.

[38] FSTAT, a program to estimate and test gene diversities and fixation indices (version 2.9.3). 2001.

[39] Excel/IF, L. Ischer HE. Arlequin suite ver 3.5: a new series of programs to perform population genetics analyses under Linux and Windows. Mol Ecol Resour. 2010;10:564–567. doi:10.1111/j.1755-0998.2010.02847.x.

[40] Peakall R, Smouse PE. Genalex 6.5: genetic analysis in programs to perform population genetics analyses under Linux and Windows. Mol Ecol Resour. 2010;10:564–567. doi:10.1111/j.1755-0998.2010.02847.x.

[41] Vazeille M, Zouache K, Vega-Rúa A, et al. Importance of mosquito “quasispecies” in selecting an epidemic arthropod-borne virus. Sci Rep. 2016;6:29564. doi:10.1038/srep29564.
[42] Pritchard JK, Stephens M, Donnelly P. Inference of population structure using multilocus genotype data. Genetics. 2000;155:945–959.

[43] Evanno G, Regnaut S, Goudet J. Detecting the number of clusters of individuals using the software STRUCTURE: a simulation study. Mol Ecol. 2005;14:2611–2620. doi:10.1111/j.1365-294X.2005.02553.x.

[44] Rosenberg N. DISTUCT: a program for the graphical display of population structure. Mol Ecol Notes. 2004;4:137–138. doi:10.1046/j.1471-8286.2003.00566.x.

[45] Rozen S, Skaletsky H. Primer3 on the WWW for general users and for biologist programmers. Methods Mol Biol. 2000;132:365–386.

[46] Ricotta C, Podani J. On some properties of the Bray-Curtis dissimilarity and their ecological meaning. Ecol Complex. 2017;31:201–205.

[47] Vega-Rua A, Zouache K, Girod R, et al. High level of retroviral restriction gene Fv1: inhibition of non-retroviral elements in genomes of Aedes aegypti and Aedes albopictus from ten American countries as a crucial factor in the spread of chikungunya virus. J Virol. 2014;88:6294–6306. doi:10.1128/JVI.00370-14.

[48] Edgar RC. MUSCLE: a multiple sequence alignment method with reduced time and space complexity. BMC Bioinformatics. 2004;5:113. doi:10.1186/1471-2105-5-113.

[49] PAUP*. Phylogenetic analysis using parsimony (and other methods). Version 4. Sunderland (MA), 1998.

[50] Vazeille-Falcoz M, Mousson L, Rodhain F, et al. Variation in oral susceptibility to dengue type 2 virus from the islands of Tahiti and Moorea, French Polynesia. Am J Trop Med Hyg. 1999;60:292–299.

[51] Schuenecker J, Iteman I, Michault A, et al. Genome variation in oral susceptibility to dengue type 2 virus discovery: significance for the arbovirus community. Viruses. 2015;7:4911–4928. doi:10.3390/v7092851.