Figure or Table # Please group Extended Data items by type, in sequential order. Total number of items (Figs. + Tables) must not exceed 10.	Figure/Table title One sentence only	Filename Whole original file name including extension. i.e.: Smith_ED_Fig1.jpg	Figure/Table Legend If you are citing a reference for the first time in these legends, please include all new references in the main text Methods References section, and carry on the numbering from the main References section of the paper. If your paper does not have a Methods section, include all new references at the end of the main Reference list.
Extended Data Fig. 1	Co-occurrence of 139 viral contig clusters identified in <i>A. aegypti</i> and <i>A. albopictus</i> mosquitoes.	Olmo_ExtFigure1.j pg	Heatmap represents the small RNA abundance for each of the 139 viral contigs in our 91 small RNA libraries from <i>A. aegypti</i> and <i>A. albopictus</i> mosquitoes. White indicates absence of small RNAs mapping to that contig. Contig clusters were defined using the dendrogram shown on the heatmap. Clusters that had a RdRp sequence were classified as a putative virus. Virus presence was considered if >50% of contigs belonging to a cluster were represented.
Extended Data Fig. 2	Phylogeny of viruses identified in <i>A. aegypti</i> mosquitoes.	Olmo_ExtFigure2.j pg	Phylogenetic trees were generated using the RdRp amino acid (aa) or nucleotide (nt) sequences and the substitution models as indicated: a , Aslam narnavirus (aa - LG + G); b , Nyamuk partiti-like virus (aa - BLOSUM 62); c , Orbis virgavirus (aa - BLOSUM62 + F); d , Bahianus rhabdovirus (aa - BLOSUM62); e , Lactea totivirus (nt - Tamura-Nei 93). Bootstrap confidence is shown close to each clade and values under 60% were omitted.
Extended Data Fig. 3	Virus-derived small RNA profiles in mosquitoes.	Olmo_ExtFigure3.j pg	Small RNA size distribution and 5' base preference is shown on the left while the density of small RNAs (coverage) is shown on the right for representative contig(s) of each of the 12 viruses identified in this study.

Extended Data Fig. 4	Burden of viruses	Olmo_ExtFigure4.j	a , Abundance of small RNA sequences in pooled
	in mosquitoes	pg	libraries from each location. Each dot represents the
	from different		small RNA abundance in a contig, and violin plots
	collection sites.		represent contig clusters (see Extended Data Fig. 1) at
			different locations with colors matching the mosquito
			species. Error bars represent the standard deviation of
			the mean for each location. The number of contigs
			analyzed per location is indicated above each graph. b ,
			Detection of representative contigs of newly detected
			viruses by RT-qPCR (black bars) in comparison to the
			detection of small RNAs (20-30 nt length) to the same
			given contig (gray bars). RT-qPCR detection is
			normalized against the endogenous constitutive gene
			<i>RpL32</i> . *, indicates detection by conventional RT-PCR.
			c , viral contig detection by conventional RT-PCR using
			independent sets of primers pairs. Conventional PCR
			and qPCR were repeated twice on the same samples.
			The expected size of viral contigs is shown. <i>ns</i>
			indicates a non-specific band. d , Sequence variation
			between viral contigs of Orbis virgavirus in RNA
			samples originated from Suriname along the region
			that is complementary to RT-qPCR primers. e , Ratio
			between relative RT-qPCR and small RNA abundance
			for each virus. The number of independent mosquito
			samples analyzed per virus is indicated above each
			graph. f , Combined incidence of DENV and ZIKV for
			each mosquito capture location in the previous,
			current, and subsequent years of collection

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			(represented by -1, 0, and +1, respectively). Data were
			obtained from public sources for each location.
Extended Data Fig. 5	Characterization	Olmo_ExtFigure5.j	a , geographic distribution of mosquitoes carrying HTV
	of HTV and PCLV	pg	and PCLV in the city of Caratinga, Brazil. Maps show
	infection in wild		the density of adult <i>A. aegypti</i> mosquitoes captured
	and laboratory		from July 2010 until August 2011 estimated from the
	mosquitoes.		number of mosquitoes captured in individual traps. All
			mosquitoes, HTV positive, PCLV positive and double
			positive individuals are shown. Virus detection was
			performed by RT-qPCR. Map source: OpenStreetMap.
			b , c tissue tropism of HTV and PCLV upon natural and
			artificial infections in <i>A. aegypti</i> mosquitoes. b , Scheme
			of mosquito dissection and tissues tested for virus
			infection by RT-qPCR. Pie charts show the prevalence
			of HTV and PCLV infection, assessed in tissues of
			naturally infected wild mosquitoes or laboratory
			mosquitoes injected with HTV and PCLV. Individual
			tissues were tested for virus presence upon dissection
			at 2-, 4-, 6- and 8-days post injection (<i>d.p.i.</i>) by RT-
			qPCR. c , Detection of HTV and PCLV in eggs by RT-
			PCR. Eggs were either rinsed with distilled water (no
			treatment group) or washed with bleach (2,5% active
			chlorine) prior to RNA extraction. The endogenous
			constitutive gene <i>RpL32</i> was used as amplification
			control. Results are representative of two independent
			experiments. d-e , HTV and PCLV do not grow in
			mammalian cell culture. VERO cells were exposed to
			mosquito extracts containing HTV (d) and PCLV (e),
			and supernatants were collected at 1-, 3- and 5-days

			post exposure. A spike containing 10 ⁵ pfu of vesicular
			stomatitis virus (VSV) was added prior RNA extraction
			and used to normalize the quantification of HTV and
			PCLV in the supernatant. No statistically significant
			difference was observed in HTV and PCLV levels at 1-,
			3- and 5-days post infection as determined by two-
			sided one-way ANOVA with Dunns' correction for
			multiple comparisons. Dots and error bars indicate the
			mean and the standard error of the mean,
			respectively. <i>n</i> indicates the number of independent
			tissue culture wells tested for each virus at each time
			point.
Extended Data Fig. 6	HTV and PCLV	Olmo_ExtFigure6.j	a-c , Strategy to evaluate the interference of HTV and
	facilitate systemic	pg	PCLV for ZIKV infection and replication in natural
	ZIKV infection in		populations of mosquitoes. (a) HTV/PCLV infected and
	mosquitoes.		virus-free wild mosquito populations were infected
			with ZIKV by intrathoracic injection. Viral loads and
			prevalence of infection were measured in the (b)
			midgut and (c) carcass of mosquitoes at 2-, 4- and 8-
			days post feeding. The prevalence of infection in each
			group is shown below plots. d-f , Laboratory
			mosquitoes (d) were infected artificially with HTV and
			PCLV and 7 days later were fed on ZIKV-infected mice.
			Viral loads and prevalence of infection were measured
			in the (e) midgut and (f) carcass of mosquitoes at 4- and
			8-days post injection. The prevalence of infection in
			each group is shown below the plots. g-i, Laboratory
			mosquitoes were infected artificially with HTV and
			PCLV or control (mock) and 7 days later infected with

			ZIKV by intrathoracic injection. Viral loads and prevalence of infection were measured in the (h) midgut and (i) carcass of mosquitoes at 2-, 4- and 8- days post injection. The prevalence of infection in each group is shown below plots. j-n , Wild mosquito populations naturally infected with HTV and PCLV were allowed to feed in mice infected with ZIKV or mock-infected controls. Viral loads of HTV and PCLV were measured in the midgut (k , l) and in the carcass (m , n) of mosquitoes at 4-, 8- and 14-days post feeding. o-p , HTV/PCLV infected or control mosquitoes were exposed to ZIKV-infected mice (o). Viral loads and prevalence of infection were measured in salivary glands (p) of mosquitoes at the indicated time points. Pie charts below each group indicate the prevalence of ZIKV infection. <i>d.p.f.</i> – days post feeding, <i>d.p.i.</i> – days post injection, <i>NS</i> – non-significant. In box plots of b , c , e , f , h , i , k , l , m , n , and p , boxes show the second and third interquartile ranges divided by the median while whiskers represent maximum and minimum values. Statistical significance was determined by two-tailed Mann–Whitney U-test. Numbers of infected samples over the total number tested are indicated above each column. Each dot represents an individual sample. Statistical significance of prevalence was determined by two-tailed Fisher's exact test.
Extended Data Fig. 7	Differential gene expression in wild	Olmo_ExtFigure7.j pg	a , Differential gene expression in the carcass of wild mosquitoes carrying HTV and PCLV or non-infected
	mosquitoes	P5	siblings during DENV infection at 4, 8 and 14 days post
	mosquitoes		sidnings during DENV infection at 4, 8 and 14 days post

	carrying HTV and		feeding. b , number of up or down regulated genes
	PCLV.		regarding the infection with HTV and PCLV at each
			time point as shown in a . Common genes across time
			points are shown. c , Immune genes regulated during
			infection with HTV and PCLV in comparison to virus-
			free siblings at different times after DENV infection. d -
			e , Intrathoracic injection of ZIKV in wild mosquitoes
			carrying HTV and PCLV or virus free siblings (d).
			Histone H4 levels were quantified in the midgut of
			mosquitoes at 2, 4, and 8 days post injection with ZIKV
			(e). Error bars represent mean and standard
			deviations of the mean, and statistical significance was
			determined by two-sided one-way ANOVA with
			Tukey's correction for multiple comparisons. f-g ,
			Artificial infection of laboratory mosquitoes with HTV
			and PCLV does not modulate levels of histone H4. g ,
			laboratory mosquitoes were artificially infected with
			HTV and PCLV and histone H4 levels were analyzed at
			different time points. In box plots of <i>e</i> and <i>g</i> , boxes
			show the second and third interquartile ranges
			divided by the median while whiskers represent
			maximum and minimum values. Statistics were
			performed using two-sided one-way ANOVA with
			Tukey's correction for multiple comparisons. Each dot
			represents an individual sample. <i>CPM</i> – counts per
			million, <i>d.p.i.</i> – days post infection, <i>NS</i> – non-
			significant.
Extended Data Fig. 8	Complexity of	Olmo_ExtFigure8.j	a , <i>Histone H4</i> gene copies in the <i>A. aegypti</i> genome
	histone genes in	pg	(Vectorbase version 52) were reannotated using
	8 M	1 FO	

the general of A	PLAST similarity soarch with further confirmation of
the genome of A.	BLAST similarity search with further confirmation of
aegypti.	RNA-seq reads mapping to each gene copy. Along with
	other <i>histone</i> genes currently annotated in Vectorbase,
	the number of copies in chromosomes or supercontigs
	are shown and the largest cluster of genes highlighted.
	b , Organization of the largest cluster of histone genes
	on chromosome 3 as indicated by the gray box. c ,
	Weblogo showing the conservation of the amino acid
	sequence of histone H4 open reading frames, which
	only varied at the positions 36 and 98, indicated by a
	circle and an asterisk, respectively. The number of
	amino acid changes in each position is indicated. d ,
	Histone H4 genes organized by nucleotide sequence
	similarity according to the dendrogram with the
	expression indicated by the heatmap in different A.
	<i>aegypti</i> tissues. Bootstrap values over 60 are shown.
	<i>Histone H4</i> genes positioned in the cluster at
	chromosome 3 are indicated by gray boxes and the
	presence of a polyadenylation signal is indicated. e ,
	Histone H4 gene expression in wild mosquito
	populations carrying HTV and PCLV or virus free
	siblings infected with DENV, quantified by RT-qPCR
	from cDNAs synthesized with random primers
	(hexamers) or anchored oligo dT ₂₂ . <i>d.p.i.</i> , days post
	injection. In <i>d</i> , SRA accession numbers in same order as
	shown in the heatmap: Female whole body (non-BF
	SRR1585314, SRR1585315, SRR1585316; 48 h post-BF
	SRR1532683, SRR1532684, SRR1532685, SRR1532693,
	SRR1532694, SRR1532695); Female brain (48 h post-BF
	SRR1166497; 96 h post-BF SRR1167481); Male (brain

	SRR1167543); Female salivary glands (SRR2659965,
	SRR2659966); Female midgut (SRR5288077, SRR5288080,
	SRR5288082, SRR5288087, SRR5288093, SRR5288100);
	Female malp. Tubules (non-BF SRR3680433, SRR3680434);
	Female carcass (12 h post-BF SRR923823; 24 h post-BF
	SRR923830; 36 h post-BF SRR923835; 48 h post-BF
	SRR923841; 60 h post-BF SRR923847; 72 h post-BF
	SRR923736); Fem. Carcass (no ovaries) 24 h post-BF
	(SRR388683); Fem. Low reprod. Tract (0 h post-mating
	SRR3213863, SRR3213864; 6 h post-mating SRR3213865,
	SRR3213866; 24 h post-mating SRR3213867, SRR3213868);
	Male sperm (early SRR3554588; late SRR3554589); Male testis
	(SRR6311395, SRR6311396); Embryo (4-8 h SRR1578254,
	SRR1578255, SRR1578256); 1 day old female ovaries
	(SRR388680); Female ovaries (non-BF SRR1167515,
	SRR1167516, SRR1167517, SRR1167518, SRR1167519,
	SRR1167520; 24 h post-BF SRR388682; 96 h post-BF
	SRR1167538, SRR1167539). f-h , silencing of histone H4 by
	RNA interference in adult mosquitoes. f , strategy for
	dsRNA mediated gene silencing in adult mosquitoes. g-
	h , Histone H4 levels in the midgut of ISV free laboratory
	mosquitoes (g) or wild mosquitoes carrying HTV and
	PCLV (h) injected with dsRNA targeting GFP (dsGFP) as
	control or histone H4 (dsH4) at 4 days post feeding. I,
	AGO2 levels in the midgut of mosquitoes carrying HTV
	and PCLV injected with dsRNA targeting GFP (dsGFP)
	as control or Ago2 (dsAGO2). Each dot represents an
	individual sample. In box plots of <i>e</i> , <i>g</i> , <i>h</i> , and <i>i</i> , boxes
	show interquartile ranges divided by the median while
	whiskers represent maximum and minimum values.
	Statistical significance was determined using two-sided
I	

	one-way ANOVA with Tukey's correction for multiple comparisons.

Item	Present?	Filename Whole original file name including extension. i.e.: Smith_SI.pdf. The extension must be .pdf	A brief, numerical description of file contents. i.e.: Supplementary Figures 1-4, Supplementary Discussion, and Supplementary Tables 1-4.
Supplementary Information	No		
Reporting Summary	Yes	nr-reporting- summary_Olmo_et_al. pdf	
Peer Review Information	Choose an item.	OFFICE USE ONLY	

Туре	Number Each type of file (Table, Video, etc.) should be numbered from 1 onwards. Multiple files of the same type should be listed in sequence, i.e.: Supplementary Video 1, Supplementary Video 2, etc.	Filename Whole original file name including extension. i.e.: Smith_ Supplementary_Video_1.mov	Legend or Descriptive Caption Describe the contents of the file
Supplementary Table	Supplementary Table 1, Supplementary Table 2, Supplementary Table 3, Supplementary Table 4,	Olmo_et_al_SupplementaryTa bles.xlsx	Supplementary Table 1 - Overview of small RNA libraries. Metadata of small RNA libraries used in our

Supplementary Table 5, Supplementary Table 6	study (SRA deposit ID, species, mosquito capture location, number of mosquitoes per pool, RNA treatment, and sequencing method).
	Supplementary Table 2 - Small RNA assembly metrics. Detailed information of assembled contigs with length >50nt and length >199 nt per small RNA library.
	Supplementary Table 3 - Overview of contigs with similarity to viral sequences deposited in GenBank. Detailed
	information of BLASTn or BLASTp hits from contigs matching viral sequences (length >199 nt) per small RNA library.
	Supplementary Table 4 - Overview of CDHit clusters.

Number of contigs t compose each CDHit cluste	that er.
Supplementary Table 5 List of oligonucleotic used in this study.	
Supplementary Table 6 - Parameters used to mode DENV transmission.	

Parent Figure or Table	Filename Whole original file name including extension. i.e.: Smith_SourceData_Fig1.xls, or Smith_ Unmodified_Gels_Fig1.pdf	Data description i.e.: Unprocessed western Blots and/or gels, Statistical Source Data, etc.
Source Data Fig. 1	Olmo_Source_Data_Figure1.xlsx	Statistical Source Data
Source Data Fig. 2	Olmo_Source_Data_Figure2.xlsx	Statistical Source Data
Source Data Fig. 3	Olmo_Source_Data_Figure3.xlsx	Statistical Source Data
Source Data Fig. 4	Olmo_Source_Data_Figure4.xlsx	Statistical Source Data
Source Data Fig. 5	Olmo_Source_Data_Figure5.xlsx	Statistical Source Data
Source Data Extended	Olmo_Source_Data_ExtDataFigure1.xl	Statistical Source Data
Data Fig./Table 1	SX	
Source Data Extended	Olmo_Source_Data_ExtDataFigure2.zi	Source Data in fasta and Newick formats
Data Fig./Table 2		
Source Data Extended Data Fig./Table 3	Olmo_Source_Data_ExtDataFigure3.zi p	Source Data of coverage plots in BED formats and bar plots in Excel table.
Source Data Extended	Olmo_Source_Data_ExtDataFigure4.zi	Statistical Source Data, unprocessed gel image
Data Fig./Table 4	p	

Source Data Extended	Olmo_Source_Data_ExtDataFigure5.zi	Statistical Source Data, unprocessed gel image
Data Fig./Table 5	р	
Source Data Extended	Olmo_Source_Data_ExtDataFigure6.xl	Statistical Source Data
Data Fig./Table 6	SX	
Source Data Extended	Olmo_Source_Data_ExtDataFigure7.xl	Statistical Source Data
Data Fig./Table 7	SX	
Source Data Extended	Olmo_Source_Data_ExtDataFigure8.zi	Statistical Source Data, annotation of histone
Data Fig./Table 8	р	genes in gff3 format

⁸ ⁹ Mosquito vector competence for ¹⁰ dengue is modulated by insect ¹¹ specific viruses

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- 64

65 **ABSTRACT**

66 Aedes aegypti and Aedes albopictus mosquitoes are the main vectors for 67 dengue virus (DENV) and other arboviruses, including Zika virus (ZIKV). 68 Understanding the factors that affect transmission of arboviruses from 69 mosquitoes to humans is a priority, because it could inform public health and 70 targeted interventions. Reasoning that interactions among viruses in the vector 71 insect might affect transmission, we analysed the viromes of 815 urban Aedes 72 mosquitoes collected from 12 countries worldwide. Two mosquito-specific 73 viruses, Phasi Charoen-like virus (PCLV) and Humaita Tubiacanga virus (HTV) 74 were the most abundant in *A. aegypti* worldwide. Spatiotemporal analyses of 75 virus circulation in an endemic urban area revealed a 200% increase in chances 76 of having DENV in wild A. aegypti mosquitoes when both HTV and PCLV were 77 present. Using a mouse model in the laboratory, we showed that the presence 78 of HTV and PCLV increased the ability of mosquitoes to transmit DENV and ZIKV 79 to a vertebrate host. By transcriptomic analysis, we found that, in DENV infected 80 mosquitoes, HTV and PCLV block the downregulation of histone H4, which we 81 identify as an important pro-viral host factor in vivo.

82

83 MAIN

84 Dengue fever is the fastest growing vector-borne disease worldwide and causes an estimated 400 million new infections every year¹⁻⁴. In addition, over the past decades, 85 86 several other arboviruses, including ZIKV and chikungunya (CHKV), have emerged 87 and caused a substantial burden of disease. Increased transmission of arboviruses 88 has been underpinned by increased geographic reach of the main vector mosquitoes, 89 A. aegypti and A. albopictus ^{4,5}, mainly due to climate change, because warming 90 produces ideal conditions for mosquitoes. Vector abundance, assessed using cross-91 sectional surveys, has long been used as a proxy for infection risk, but the incidence 92 of arbovirus infection does not directly correlate with mosquito abundance⁶. We still 93 lack a complete understanding of the factors that affect rates of transmission to 94 humans.

95

96 Virologic surveillance of adult *Aedes* mosquitoes by metagenomic analysis can lead to 97 early identification of circulating arboviruses, and help raise preparedness to inform 98 public health measures that can curtail or even prevent outbreaks⁷. In addition to 99 arboviruses, these surveillance efforts have also identified an enormously diverse set 100 of insect-specific viruses (ISVs) in *Aedes* mosquitoes ^{8–12}. Although ISVs do not infect 101 vertebrates, they have been shown to affect the capacity of the mosquito to be infected,

maintain and transmit arboviruses, which together comprise vector competence and
 will therefore affect the incidence of infection in humans ^{7,13,14}.

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In order to carry out a comprehensive characterization of the viromes of mosquitoes that can harbour arboviruses, and inform risk assessment and public health strategies to mitigate arbovirus disease, we collected in the wild and performed metagenomic analysis of over 800 adult *Aedes* mosquitoes, and report our findings here.

109

110 **RESULTS**

111 Virome analysis of *Aedes* mosquitoes

112 Adult A. aegypti and A. albopictus mosquitoes were collected from the field in 12 113 different sites from six countries on four continents (Fig. 1a). In total, 815 adult 114 mosquitoes were pooled according to species, location and date of collection, resulting 115 in 91 samples derived from 69 A. aegypti and 22 A. albopictus. Details of the pools are 116 described in **Supplementary Table 1**. Whole mosquito samples were used to extract 117 RNA and construct small RNA libraries that were sequenced and analyzed using a 118 shotgun metagenomic strategy previously optimized to detect viruses (See ref.⁸). 119 Briefly, this strategy is based on the detection of virus-derived small RNAs that are 120 used to assemble longer contiguous sequences for further characterization. In total, 121 we identified putative viral 1448 contigs present in our mosquito samples (Fig. 1b,c). 122 Data curation (described in Fig. 2a and in the methods section) suggested that these 123 contigs represent at least 12 different viruses based on the phylogeny of polymerase 124 genes (Fig. 2b), including seven known viruses previously identified as ISVs. Out of 125 these, three remain unclassified while the other four belong to the Phenuiviridae, 126 Xinmoviridae, Bunyaviridae and Flaviviridae families (Fig. 2b). No known arboviruses 127 were detected in our metagenomic analysis. The five remaining viral polymerase 128 sequences showed low similarity to the closest known reference in Genbank. 129 Phylogenetic analyses confirmed that they are likely new viral species (Extended Data 130 Fig. 2), belonging to the Partitiviridae, Totiviridae, Rhabdoviridae, Narnaviridae and 131 Virgaridae families (Fig. 2b). These viruses were named according to their 132 classification (Fig. 2b). All new viruses were most closely related to known ISVs 133 (Extended Data Fig. 2), but their final classification requires biological 134 characterization.

135

All 12 identified viruses, 7 known and 5 new, had RNA genomes, either single-stranded
(of positive and negative polarity) or double-stranded (Fig. 2b). The small RNA profile
observed for these viruses shows clear production of siRNAs (Extended Data Fig. 3),

139 which results from the activity of the RNAi pathway during active viral replication in the mosquito host ^{8,15–19}. Viruses detected in Aedes mosquitoes were strictly species-140 141 specific and often associated with specific locations (Fig. 2c). Out of the 12 identified 142 viruses, 10 were found in A. aegypti and two in A. albopictus suggesting a less diverse 143 virome in the latter even when accounting for a lower number of samples. Indeed, 144 looking at the diversity of the mosquito virome per country, a single virus species was 145 detected in each A. albopictus population while 4-6 different viruses were present in 146 A. aegypti (Fig. 2d). In addition, comparing different mosquito species that were 147 collected from the same sites in Caratinga, Montes Claros, Lope, Franceville and 148 Singapore, we observe that A. aegypti had higher virome diversity than A. albopictus 149 in 4 out of 5 cases (Fig. 2c).

150

151 Using the small RNAs mapping to each virus contig as a proxy for abundance, we 152 found that viral loads varied for the same virus in different locations and also between 153 different viruses (Extended Data Fig. 4a). Detection of these new viruses in the 154 original RNA samples from wild mosquitoes used for the metagenomic analysis was 155 confirmed by RT-gPCR and conventional PCR (Extended data Fig. 4b,c). These 156 results validated our small RNA sequencing strategy. Detection of Orbis virgavirus 157 (OVV) by RT-qPCR failed in samples from Suriname, but this was attributed to 158 polymorphisms in the primer annealing region (Extended data Fig. 4d). For each 159 virus, the viral load determined by small RNA abundance corresponded with gRT-PCR 160 detection. However, the relative quantification of viruses by gRT-PCR and small RNA 161 abundance was unique for each virus, with some being underestimated (PCLV) or 162 overestimated (Aslam narnavirus) (Extended data Fig. 4e).

163

164 **Biogeography of mosquito viruses**

165 Three of twelve locations had five or more viruses circulating in the local A. aegypti 166 population: Santos, Paramaribo and Singapore (Fig. 2c). Notably, these are all port 167 cities, which are likely to have a continuous influx of mosquitoes. Three different lines 168 of laboratory A. aegypti mosquitoes lacked any viruses according to our analysis (Fig. 169 2c). Most of the viruses we detected were present in mosquitoes at single sites but 170 five were present on at least two continents (Fig. 2d). In A. aegypti, two known ISVs, 171 PCLV and HTV, were present in more than half of the samples (Fig. 2c), with the 172 remaining eight viruses found in less than 20% of the samples. No viruses were found 173 with a prevalence higher than 20% in A. albopictus and only one was present in 174 multiple sites (Fig. 2c,d). HTV and PCLV in A. aegypti had the highest viral loads of all 175 viruses in either mosquito species (Extended Data Fig. 4) but were not present in any

176 samples of *A. albopictus*. Notably, HTV and PCLV were either absent, or present at 177 very low viral loads in A. aegypti mosquitoes collected in Africa (Fig. 2c), where 178 transmission of arboviruses, such as DENV and ZIKV is low (Extended data Fig. 4f)^{1.2}. 179 High loads of HTV and PCLV were observed in mosquitos sampled in areas with high 180 DENV/ZIKV incidence, namely Asia and South America (Extended data Fig. 4f)^{1.2}. 181 We hypothesized that there was a positive association between ISVs and arboviruses, 182 which was unexpected (competition between RNA viruses in the same host would be 183 more likely).

184

185 Circulation of ISVs and arboviruses in the wild

186 In order to examine the spatiotemporal dynamics of the two major resident viruses in 187 wild mosquitoes, HTV and PCLV, we chose to focus on one of the 12 sites used for 188 the metagenomic analysis using a collection of archived mosquito RNA samples, 515 189 A. aegypti and 24 A. albopictus, previously collected over a year, August 2010 to July 190 2011, in Caratinga city, southeast Brazil (Fig. 3a). This dataset was previously used to 191 assess DENV circulation in an endemic urban area²⁰. Based on our metagenomics 192 approach, we detected three viruses in Caratinga mosquitoes, OVV, HTV and PCLV 193 (Fig. 2c), which we confirmed using RT-qPCR. OVV, HTV and PCLV were detected in 194 wild A. aegypti but were absent from A. albopictus, even though both species were 195 often captured in the same traps (Fig. 3b). OVV was only detected in three individual 196 mosquitoes, and we focused our analyses on HTV and PCLV, which were present at 197 prevalence of 61% and 83%, respectively (Fig. 3b,c). Based on this survey of 198 individual mosquitoes by RT-qPCR, we confirmed that HTV and PCLV were highly 199 prevalent in natural mosquito populations during the whole period of collections and 200 independent of the location within the city (Fig. 3d and Extended Data Fig. 5a). 201 Moreover, we observed a strong positive association between the presence of HTV 202 and PCLV in mosquitoes (p<1E-10, chi-squared test), suggesting that co-infection 203 might be advantageous for these viruses.

204 HTV and PCLV are presumed to be ISVs although they are poorly 205 characterized to date 8.21.22. We detected HTV and PCLV in different tissues, including 206 the salivary glands, which suggested that they could be transmitted by a mosquito bite 207 (Extended Data Fig. 5b,c). To assess the possibility that HTV and PCLV could be 208 transmitted to humans, we analyzed human blood samples collected concomitantly 209 with mosquitoes in Caratinga city, southeast Brazil from February to July 2011. Human 210 blood samples and mosquitoes from Caratinga were previously analyzed by RT-qPCR 211 for the presence of DENV ²⁰ and these data were used for comparison. Plotting the 212 numbers of this previous analyses, we observe that DENV was detected in less than

213 5% of mosquitoes, but in more than 30% of human blood samples (Fig. 3c). We did 214 not detect HTV or PCLV in RNA extracted from human blood samples from Caratinga, 215 despite their high prevalence in mosquitoes, suggesting that these viruses are unable 216 to productively infect humans (Fig. 3c). Furthermore, HTV and PCLV do not grow in 217 mammalian cell lines, such as Vero cells, reinforcing the idea that they are ISVs 218 (Extended Data Fig. 5d,e). HTV and PCLV were detected in mosquito eggs, which 219 suggests that these viruses are maintained in mosquitos by vertical transmission 220 (Extended Data Fig. 5c).

221 Our data indicate that HTV and PCLV are not infectious to humans, but we 222 wanted to understand whether they may affect how arboviruses are transmitted. In the 223 dataset of mosquitoes from Caratinga city analyzed here, we observed a statistically 224 significant enrichment of HTV and PCLV in individuals that also harbored DENV (Fig. 225 **3E**). Statistical analyses revealed that both HTV alone (OR 2.59, [1.09,716] 95%CI) 226 and HTV/PCLV co-infections (OR 3.06, [1.29,8.46] 95% CI) are both associated with 227 the presence of DENV in mosquitoes, whereas PCLV alone had no statistically 228 significant association with DENV. However, due to positive association between HTV 229 and PCLV that we identified in mosquitoes from Caratinga (p < 1E - 10, chi-squared test, 230 described above), it is hard to dissect the contribution of each virus. We note that our 231 analysis is based on a small sample of mosquitoes, it and could be affected by 232 physical, ecological or environmental factors. However, in Caratinga, a town that is 233 only 3 km wide and 3 km long, ecological conditions are likely homogenous on the 234 same collection date. Population density was not found to be important in a previous 235 study from our group²³. In addition, our analyses found neither geographic nor temporal 236 patterns of virus distribution in Caratinga (Fig. 3d and Extended data Fig. 5a). Thus, 237 our observations using field samples strongly indicate a positive interaction between 238 two ISVs, HTV and PCLV, and the arbovirus DENV in mosquitoes.

239

240 HTV and PCLV increase arbovirus replication in mosquitoes

Since we had access to ovitraps from Rio de Janeiro, where HTV and PCLV were also found at high prevalence, we obtained a few hundred eggs from wild *A. aegypti* mosquitoes and reared them in the laboratory for 2 generations. From the F2 of labreared population, we pooled eggs from 5 individual females that were either free of any virus, or carried both HTV and PCLV to produce two separate mosquito lines (see Methods for details). We were unable to isolate lines carrying only HTV or PCLV, reinforcing the strong association observed in our wild sample cohort.

249 We exposed the two separate mosquito lines that were ISV free or co-infected 250 with HTV/PCLV to blood feeding on mice previously infected with DENV or ZIKV using 251 an infectious blood meal. We found that mosquitoes carrying HTV and PCLV had 252 similar prevalence and viral loads of DENV in the midgut at 4, 8 and 14 days post 253 feeding (d.p.f.) to that present in ISV-free mosquitoes (Fig. 4a,b). At 14 d.p.f., we 254 observed a trend towards higher viral load of DENV in the midgut of mosquitoes 255 carrying HTV/PCLV but this trend was not statistically significant (Fig. 4b). In the 256 carcass of mosquitoes, we observed a five-fold significant increase in DENV levels at 257 8 and 14 d.p.f. in individuals carrying HTV/PCLV compared to ISV free controls (Fig. 258 **4c**). Mosquitoes with HTV/PCLV also displayed higher susceptibility to ZIKV than ISV-259 free mosquitoes (Fig. 4d-f). ZIKV RNA levels were significantly increased at 4, 8 and 260 14 d.p.f. in the midgut of mosquitoes carrying HTV and PCLV compared to ISV free 261 controls (Fig. 4e). In the carcass, we observed a fivefold increase in dissemination at 262 4 d.p.f. and significantly 10-fold higher viral loads at 8 d.p.f. in the presence of HTV 263 and PCLV. Overall, our results demonstrate increased systemic DENV and ZIKV 264 infection in mosquitoes carrying HTV/PCLV compared to virus-free controls.

To further investigate the specific effect of HTV and PCLV during the systemic phase of infection, we directly injected ZIKV into the mosquito haemocele, which bypasses the stage of midgut infection (**Extended Data Fig. 6a**). Here, we also observed increased viral replication in the carcass of HTV/PCLV carrying wild mosquitoes compared to ISV-free controls (**Extended Data Fig. 6a-c**).

270

271 Our mosquito colonies established from individuals with and without ISVs were derived 272 from the same but highly heterogenous wild population. It is therefore possible that our 273 selection generated colonies composed of individuals that differed with regards to their 274 genetic backgrounds in addition to the presence of ISVs. To rule out a role for the 275 genetic background, we next performed experiments with laboratory mosquitoes that 276 are genetically more homogenous. Laboratory mosquitoes were artificially infected 277 with HTV and PCLV to test whether ISVs have a direct impact on the susceptibility to 278 arboviruses (Extended data Fig. 6d). Notably, HTV and PCLV loads and tissue 279 tropism during artificial injection were similar to naturally infected mosquitoes after 8 280 days post injection (Extended Data Fig. 5b). Artificially infected laboratory mosquitoes 281 had increased systemic ZIKV RNA levels at 8 d.p.f. compared to controls, similar to 282 what we observed for lines carrying HTV and PCLV derived from wild populations 283 (Extended data Fig. 6d-f). Increased systemic viral replication was also observed 284 when laboratory mosquitoes were artificially infected with HTV and PCLV before being 285 injected with ZIKV (Extended Data Fig. 6g-i). Although artificially infected lab

mosquitoes did not show increased ZIKV replication in the midgut, this can be
explained by the fact that naturally infected mosquitoes have more marked effects due
to presence of HTV and PCLV throughout development (Extended Data Fig. 5b).

289

Notably, in wild mosquitoes, ZIKV infection had a positive impact on PCLV levels in the midgut of infected mosquitoes (**Extended data Fig. 6j,k**), which suggests a mutual beneficial interaction between these viruses. As mentioned before, HTV was not detected in the midgut even in the presence of ZIKV (**Extended data Fig. 6I**). Neither HTV nor PCLV were consistently affected by ZIKV infection in the carcass, although we observed an increase in PCLV levels and a reduction in HTV levels in single time points (**Extended data Fig. 6m,n**).

297

298 Transmission of DENV and ZIKV is increased by ISVs

299 We tested whether increased ZIKV and DENV levels in mosquitoes carrying HTV and 300 PCLV led to increased amounts of arboviruses in mosquito salivary glands, and 301 increased transmission to a vertebrate host. Wild HTV/PCLV positive mosquitoes 302 showed faster kinetics and higher prevalence of ZIKV infection in salivary glands 303 compared to ISV-free controls (Fig. 4g and Extended Data Fig. 6o-p). We simulated 304 vectorial transmission in a susceptible animal model using mice deficient for type I and 305 type II interferon receptors^{24,25} (**Fig. 4h**). We opted to test ZIKV transmission because 306 the mouse model for this virus is more robust than for DENV. Mice were incubated with 307 ZIKV-infected mosquitoes at 6, 8 and 12 d.p.f. and viremia was analyzed in these 308 animals. No viremia was observed in mice bitten by mosquitoes at 6 d.p.f. (Fig. 4i). 309 Mosquitoes were able to efficiently transmit ZIKV to 5 out of 5 mice at 8 d.p.f. but only 310 in the presence of HTV and PCLV (Fig. 4i). At 12 d.p.f., mosquitoes with or without 311 HTV/PCLV were able to equally transmit ZIKV to 3 out of 3 mice (Fig. 4i). Thus, the 312 presence of ISVs is associated with shortening of the extrinsic incubation period (EIP) 313 of ZIKV, which is the time required for infected mosquitoes to become infectious to a 314 vertebrate host. While mosquitoes carrying HTV and PCLV were able to transmit ZIKV 315 between 7 and 8 days, ISV free individuals required between 9 and 12 days. Thus, the 316 presence of ISVs in mosquitoes could lead to shortening of the EIP between 1 and 5 317 days although our experiments did not allow us to pinpoint the exact difference. 318 Furthermore, mosquitoes carrying HTV and PCLV and analyzed at both 8 and 12 d.p.f. 319 had significantly higher ZIKV levels compared to ISV free controls (Fig. 4j). 320

To further elucidate the impact of HTV and PCLV infection on the EIP we applied a previously developed mathematical model ^{26,27}. Our modelling demonstrates that even

323 small changes in EIP could have a large impact on the number of human infections 324 (Fig. 4k). For example, shortening the EIP from 10 days to 8 days, within the range of 325 the difference we observed between mosquitoes with and without ISVs in the 326 laboratory, could lead to a 5-fold increase in the number of infections (Fig. 4k), which 327 can be explained by the short average life expectancy of Aedes mosquitoes in the wild. 328 As a consequence, arboviral prevalence in mosquitoes is also increased due to both 329 the increased availability of infected humans and faster viral kinetics inside mosquitoes 330 (Fig. 4I), providing a link between our field observations and laboratory experiments. 331 Although this model was parameterized for DENV transmission, the results should 332 broadly hold for other arboviral diseases transmitted by the same mosquito vector, 333 including ZIKV.

334

335 HTV and PCLV modulate histone H4 expression in mosquitoes

336 In order to probe the biological mechanisms by which ISVs affect systemic 337 dissemination of arboviruses in A. aegypti mosquitoes, we analyzed the transcriptome 338 of RNA harvested from entire mosquito carcasses at different times after DENV 339 infection (4, 8 and 14 d.p.f.). Overall, we found that the presence of HTV and PCLV 340 had little effect on the transcriptome of DENV infected mosquitoes (Extended data 341 Fig. 7a). Only 100/ 10000 genes analysed were significantly up- or down regulated 342 and less than 10 were common between time points (Extended data Fig. 7b). Of 343 interest, genes associated with known antiviral pathways, such as Toll, IMD, Jak-344 STAT, autophagy and RNA interference did not show any consistent differences in 345 expression (Extended data Fig. 7c). Next, we compared the transcriptome of DENV 346 infected and non-infected individuals from groups of mosquitoes carrying HTV and 347 PCLV, or virus-free controls, using Gene Set Enrichment analysis (GSEA)²⁸ (Fig. 5a). 348 We focused our analysis on the carcass of mosquitoes infected by DENV at 8 and 14 349 d.p.f. where the presence of HTV and PCLV had the strongest effect (Fig. 5a). This 350 analysis identified 7 biological pathways that were significantly affected both by the 351 presence of HTV / PCLV and DENV infection in at least one time point (Fig. 5b). 352 Notably, all pathways were downregulated during DENV infection and upregulated by 353 the presence of HTV and PCLV, as indicated by the enrichment score (NES) (Fig. 5b). 354 Out of these, four pathways were significantly affected in at least 6 out of 8 355 comparisons: nucleosome, nucleosome assembly, DNA templated transcription 356 initiation and protein heterodimerization activity (Fig. 5b). Analysis of genes 357 responsible for the significant enrichment, showed that they were almost the same for 358 these 4 biological pathways (Fig. 5c). Indeed, histones represented the majority of 359 genes differentially regulated by DENV infection and the presence of HTV and PCLV,

360 with histone H4 topping the list (Fig. 5c). Thus, we used RT-qPCR to analyze histone 361 H4 expression and validate our observations in independent experiments using ZIKV. 362 Histone H4 expression was significantly downregulated by ZIKV infection in the 363 carcass of infected mosquitoes in a time-dependent manner (Fig. 5d,e), which was 364 prevented by the presence of HTV and PCLV (Fig. 5e). Furthermore, levels of histone 365 H4 were significantly higher in the presence of HTV and PCLV at every time point 366 tested when compared to ISV free controls (Fig. 5e). We also observed upregulation 367 of histone H4 in the midgut, but only at 4 d.p.i. (Extended Data Fig. 7d,e). Importantly, 368 differential expression of histone H4 between mosquitoes with or without HTV/PCLV 369 was only observed in the presence of DENV and ZIKV infections (Fig. 5f,g). We 370 showed that artificial infection of laboratory mosquitoes with HTV and PCLV alone did 371 not significantly affect histone H4 expression (Extended Data Fig. 7f,g).

372

373 Histone genes are highly conserved, often found in multiple copies that lack 374 polyadenylation signals ²⁹. Yet, there are non-canonical histone genes that possess 375 polyadenylation signals. The genome of A. aegypti encodes at least 299 histone genes 376 in the assembled chromosomes (chr.) and another 135 copies present in extra 377 Supercontigs (Extended Data Fig. 8a). In comparison, humans encode only about 80 378 histone genes in total²⁹, despite having a larger genome. Most histone genes (267 out 379 of 299) were found in a single cluster on chr. 3 (Extended Data Fig. 8a,b). With 380 regards to histone H4 we identified 66 genes in total and 59 in the cluster on chr. 3 381 (Extended Data Fig. 8a). Histone H4 genes showed high similarity, with almost 100% 382 aminoacid conservation but some sequence variation at the nucleotide level, 383 especially in the copies outside of the chr.3 cluster (Extended Data Fig. 8c,d). Histone 384 H4 genes that have clear polyadenylation signals were the most detected in our 385 dataset and in the available transcriptome of mosquito tissues since they all were 386 prepared using polyA selection (Extended Data Fig. 8d). Interestingly, the effect of 387 HTV and PCLV on histone H4 RNA levels was not significant when we analyzed 388 polyadenylated copies, which represented less than 10% of all histone H4 expression 389 (Extended Data Fig. 8e). This suggests HTV and PCLV may primarily affect non-390 polyadenylated histones that are coordinately synthesized with DNA replication during 391 the S-phase of cell division and are stable after incorporation into chromatin ³⁰.

392

393 Silencing of histone H4 and DENV replication in mosquitoes

We applied dsRNA-mediated gene silencing to knock down histone H4 expression in adult mosquitos prior to infection with DENV or ZIKV (**Fig. 5h** and **Extended Data Fig.** Mosquitos silenced for histone H4 before infection had significantly reduced

397 DENV levels in the midgut at 4 and 8 d.p.i., although the difference was not significant 398 at the later time point (Fig. 5i). As a result, we observed slower kinetics of infection in 399 the carcass with lower prevalence at 4 d.p.i. and reduced viral loads at later time points 400 (Fig. 5j). These results indicate that histone H4 is an important pro-viral host factor 401 during DENV infection. As observed for the effect of HTV and PCLV, histone H4 402 affected the kinetics of DENV infection but was not essential for viral replication. 403 Together, our data suggest that two highly prevalent ISVs, HTV and PCLV, affect 404 mosquito vector competence for DENV and ZIKV by preventing the downregulation of 405 histone H4, a novel proviral host factor. To test this hypothesis, we directly targeted 406 histone H4 in mosquitoes carrying HTV and PCLV by using dsRNA mediated gene 407 silencing. We did not observe any changes in histone H4 expression in mosquitoes 408 injected with cognate dsRNA (Extended Data Fig. 8h). This is in stark contrast to the 409 efficient silencing triggered by the same dsRNA sequence in mosquitoes that did not 410 carry HTV and PCLV (Extended Data Fig. 8g). Notably, another dsRNA targeting the 411 AGO2 nuclease that is central to the RNA interference pathways was able to trigger 412 efficient silencing in the same mosquitoes carrying HTV and PCLV (Extended Data 413 Fig. 8i). These results again point to a specific effect of HTV and PCLV on histone H4 414 expression in mosquitoes.

415

416 **DISCUSSION**

We report positive interactions between ISVs and arboviruses in mosquitoes in the wild and in the laboratory. Previously ISVs have mainly been reported to interfere with arbovirus replication in mosquitoes (superinfection exclusion)^{7,31,32}. As HTV and PCLV are the most abundant ISVs that we detected in wild *A. aegypti* mosquitoes, it is feasible that they can have a substantial impact on the global transmission of DENV and ZIKV.

423 We also showed that HTV and PCLV increase histone H4 expression during 424 DENV infection and that histone H4 is a pro-viral host factor for the replication of DENV 425 in mosquitoes. We propose that ISVs increase DENV infection through upregulation of 426 histone H4. However, we were unable to establish a direct connection between the 427 regulation of histone H4 expression and the increase in vector competence. Regarding 428 the role of histone H4 as a putative proviral factor, it is worth mentioning that C protein 429 from flaviviruses interacts with histones and is capable of interfering with nucleosome 430 assembly ³³. Recent work further suggests that the C protein of Yellow fever virus and 431 possibly other flaviviruses mimics the tail of histone H4 and regulates gene expression 432 to favour infection ³⁴. Thus, downregulation of histone H4 may be part of a coordinated 433 host response to limit the ability of flaviviruses to control gene expression, which could

be counteracted by HTV and PCLV. We observed no major changes in gene expression in the presence of HTV and PCLV, which suggests that the proviral role of histone H4 is not achieved through major regulation of gene expression. Rather, replication-dependent histones seem to be preferentially regulated by HTV and PCLV, which could point to a mechanism involving cell division. Notably, replicationdependent histones are targeted by RNA interference ³⁵, which could provide a connection with the role of this pathway in the antiviral defense of mosquitoes ^{16,36}.

Although previous studies have reported interactions between ISVs and arboviruses, most were performed in cell lines^{37–40}. One exception is Nhumirim virus (NHUV) and Cell fusing agent virus (CFAV), which were shown to interfere with replication of arboviruses in the same family as West Nile virus, DENV and ZIKV ^{31,41,42}. Interestingly, in cell lines, PCLV either inhibited or did not affect the replication of ZIKV^{21,22}. Also since we could not test the presence of PCLV alone in adult mosquitoes, we cannot rule out that HTV has the predominant proviral effect.

Further work will be needed to understand how HTV and PCLV regulate histone H4 during DENV and ZIKV infection, and whether this mechanism affects other arboviruses such as CHIKV. Understanding how ISVs affect vector competence could reveal alternative strategies for controlling arbovirus transmission.

453 **METHODS**

454

455 **Ethics statement**

456 All procedures involving vertebrate animals were approved by the ethical review 457 committee of the Universidade Federal de Minas Gerais (CEUA 337/2016 and 458 118/2022 to J.T.M.). Mosquito were collected in Gabon under the research 459 AR0013/17/MESRS/CENAREST/CG/CST/CSAR authorization delivered by 460 CENASREST. Unlinked anonymous testing of human blood samples was approved 461 by the ethics committee in research (COEP) of Universidade Federal de Minas Gerais 462 (number 415/04 to EGK).

463

464 Human blood samples

Human samples were previously reported²⁰. Forty-four blood samples were collected from patients that sought medical attention in the city of Caratinga between February and July of 2011 by professional nurses as part of a city surveillance plan. Blood samples were mixed with EDTA as an anticoagulant and stored at 4 °C. Serum was obtained from blood samples and inactivated at 56 °C for 30 min. Total RNA extraction from human blood samples was performed using Trizol LS Reagent (Invitrogen).

471

472 Data reporting

473 No statistical methods were used to predetermine sample size. Experiments were not
474 randomized and investigators were not blinded to allocation during experiments and
475 outcome assessment.

476

477 Mosquito collection in the field

Locations of mosquito collections are described in the **Supplementary Table 1**. Field traps were used to collect adult mosquitoes that were further identified using morphological characteristics. Whole mosquitoes were ground in TRIzol (Invitrogen) and kept refrigerated prior to RNA extraction. Collection and processing of individual mosquitoes from Caratinga for spatiotemporal analysis of virus circulation were previously reported²⁰. RNA samples of these mosquitoes were re-analyzed in this work.

485

486 **RNA extraction**

RNA was isolated using TRIzol reagent (Invitrogen) following the manufacturer
protocol with minor modifications. Briefly, individual mosquito samples or tissues were
collected in 1,5 mL tubes, 3-5 glass beads (1 mm diameter) and ice-cold TRIzol were

added before being homogenized in a Mini-BeadBeater-16 (Biospec[®]) for 90 seconds.
Glycogen (Ambion) was added (10µg per sample) to facilitate pellet visualization upon
RNA precipitation. RNAs were resuspended in RNAse-free water (Ambion) and stored
at -80°C.

494

495 Small RNA library construction

496 Different strategies for library construction were implemented and are indicated in 497 Supplementary Table 1. The strategy was determined according to RNA quality 498 evaluated by the 2100 Bioanalyzer system (Agilent). Libraries were built using total 499 RNA or size selected small RNAs (18-30 nt), depending on quality and yield of the 500 sample. In the case of low RNA yield, especially when the source was a single 501 mosquito, total RNA was directly used as input for library preparation. For samples 502 with more than 20 ug of RNA available, small RNAs were selected by size (18–30 nt) 503 on a denaturing PAGE. For samples with more than 20 ug of total RNA that displayed 504 a degradation profile (i.e., lack of sharp ribosomal RNA peaks), total RNA was 505 subjected to oxidation using sodium periodate ^{43,44}, prior to size selection. Oxidized 506 and non-oxidized size selected RNAs (18-30 nt) were used as input for library 507 construction. In all cases, libraries were prepared utilizing the TruSeg[®] Small RNA 508 Library Prep Kit (Illumina®) or NEBNext® Multiplex Small RNA Library Prep Set for 509 Illumina[®] (New England BioLabs inc.) following protocols recommended by the 510 manufacturers. Libraries were pooled and sequenced at the GenomEast sequencing 511 platform at the Institut de Génétique et de Biologie Moléculaire et Cellulaire in 512 Strasbourg, France.

513

514 Small RNA-based metagenomics for virus identification

515 After sequencing, raw sequenced reads from small RNA libraries were submitted to 516 adaptor trimming using cutadapt ⁴⁵ v1.12, discarding sequences with low Phred quality 517 (< 20), ambiguous nucleotides and/or with length shorter than 15 nt. Remaining 518 sequences were mapped to reference sequences of A. aegypti (AaeL5)⁴⁶ or A. 519 albopictus ⁴⁷ using Bowtie ⁴⁸ v1.1.2 allowing no mismatches. Size profiles of small 520 RNAs matching reference sequences and 5' nt frequency were calculated using in-521 house Perl v5.16.3, BioPerl library v1.6.924 and R v3.3.1 scripts. Plots were made in 522 R using ggplot2 v2.2.0 package. Sequences that did not present similarities with 523 bacteria or the host genomes were used for contig assembly and subsequent 524 analyses. Assembly was performed essentially as previously described ⁸ with the 525 following changes: (1) We replaced Velvet ⁴⁹ assembler by SPAdes ⁵⁰ on the second 526 round of contig assembly. (2) Assembled contigs ranging from 50 to 199 nt were

527 characterized solely based on sequence similarity search against Viral RefSeq 528 Database ⁵¹. (3) Contigs greater than 200nt were characterized based on sequence 529 similarity against the NCBI NT and NR databases and submitted to pattern-based 530 strategies. (4) For manual curation of putative viral contigs, top 5 BLAST ⁵² hits were 531 analyzed to rule out similarity to other organisms, ORF organization and small RNA 532 size profile (distribution and coverage) were analyzed to differentiate between viruses. 533 Contigs containing truncated ORFs and small RNA profiles without the presence of 534 symmetric small RNA peaks at 21 nt were considered to be EVEs as described ¹⁵. (5) 535 Manually curated viral contigs were grouped using CD-HIT ⁵³ requiring 90% of 536 coverage with 90% of identity to remove redundancy. Representative contigs were 537 used for co-occurrence analysis based on small RNA abundance on each of the small 538 RNA libraries available. Contigs grouped into a single cluster (Hierarchical clustering 539 based on Pearson correlation) were then used as trusted on SPAdes for a re-assembly 540 step using all the libraries in which that viral sequence was found. In total, we 541 assembled 7260 contigs larger than 200 nt (Fig. 1b, assembly metrics in 542 Supplementary Table 2). 1448/ 7240 contigs were identified as putative viral 543 sequences using sequence similarity searches against non-redundant nucleotide and 544 protein databases (NT and NR, respectively) at GenBank (Fig. 1b, Supplementary 545 **Table 3**). Although the number of contigs assembled per library varied greatly, we 546 observed high abundance and diversity of viral contigs in most samples (Fig. 1c). 547 Comparing results from the two mosquito species, the percentage of viral contigs was 548 substantially smaller in A. albopictus libraries compared with A. aegypti (Fig. 1c). In 549 addition, we noted more variation in the number of assembled contigs, and larger 550 proportions of unknown contigs, in libraries from A. albopictus, probably because this 551 species is less studied compared to A. aegypti (Fig. 1c). Most animal genomes contain 552 integrated viral sequences known as endogenous viral elements (EVEs) that are transcribed and generate small RNAs 54-56. In order to discriminate sequences of 553 554 viruses from EVEs, we took advantage of the small RNA profile associated with ORF 555 analysis and contig size (Fig. 2a) ¹⁵. This filter identified 446 putative EVE sequences 556 that were removed from the initial viral contigs (Fig. 2a). The remaining viral contigs, 557 representing putative viruses, were grouped into 158 unique clusters (based on 558 sequence similarity) (Supplementary Table 4). In 19 clusters, parts of contigs had 559 significant similarity to two different viruses. Our results suggested that these were 560 misassemblies and thus they were removed from further analyses (Fig. 2a). Contigs 561 representing the remaining clusters were used to evaluate co-occurrence in the 91 562 small RNA libraries from A. aegypti and A. albopictus (Extended Data Fig. 1). The 563 occurrence of contigs in each library was indicated by the normalized number of small

564 RNA reads mapped to each reference. Across the small RNA libraries, contigs that 565 consistently co-occurred and shared similar expression profiles were considered 566 probable fragments from the same viral genome (Fig. 2a). This analysis yielded a total 567 of 12 clusters of co-occurring contigs and 3 single contigs that had no additional 568 partners. Contigs from each cluster were further analyzed based on the closest 569 reference sequence to determine the putative organization of fragments along the viral 570 genome. This analysis showed that clusters #2 and #3 contained contigs belonging to 571 the same virus, PCLV, and were considered together for further analyses (Extended 572 Data Fig. 1). To further classify the clusters and single contigs, we focused on the 573 ones that represented sequences encoding viral polymerases. We were able to identify 574 clear polymerase sequences in each of the 11 clusters and in one out the 3 single 575 contigs.

576

577 Phylogenetic analyses

578 Assembled viral contigs were submitted to analysis of conserved domains to identify 579 NCBI RdRp-related regions using Conserved Domain Search 580 (https://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi). For each putative virus, the 581 largest RdRp segment was used to identify virus relatives at NCBI sequence 582 databases (NT and NR) using sequence-similarity searches through BLAST tool. 583 Multiple alignments were performed using the MAFFT online tool ⁵⁷ available at 584 https://www.ebi.ac.uk/Tools/msa/mafft/. For putative new viruses identified at protein 585 level (Narnaviridae, Partitiviridae, Rhabdoviridae, Totivirdae and Virgaviridae), amino 586 acid sequences were selected, and phylogenetic analyses were carried out on 587 CIPRES Portal version 3.3 (https://www.phylo.org/portal2) ⁵⁸. The best-fit model of 588 protein evolution was selected in ModelTest-NG ⁵⁹ for each viral species, using 589 Maximum Likelihood (ML) method. For the virus from *Totiviridae* family, an additional 590 strategy was also applied using nucleotide sequences where the best-fit model was 591 defined using MEGA-X tool ⁶⁰, and the tree was constructed using ML method. For all 592 phylogenetic trees, clade robustness was assessed using bootstrap method (1000 61 593 pseudoreplicates) and trees were edited using iTOL version 5.7 594 (https://itol.embl.de/).

595

596 Mosquitoes

597 Wild *A. aegypti* mosquitoes used in cage experiments were F2 to F5 generations 598 derived from eggs collected in the Rio de Janeiro city (Urca neighborhood) in Brazil 599 and were kindly provided by Dr. Rafael M. de Freitas from Fiocruz-RJ and Dr. Luciano 600 A. Moreira from Fiocruz-MG. The laboratory Red-eye strain ⁶² was kindly provided by

Prof. Pedro C. Oliveira from the Universidade Federal do Rio de Janeiro - UFRJ. *A. aegypti* mosquitoes were maintained in a climatic chamber at 28°C and 70-80%
relative humidity, in a 14:10 hour light:dark photoperiod, and 10% sucrose solution *ad libitum.* Mosquito cages contained individuals that emerged in a 48h-interval.

605

606 Generation of HTV+/PCLV+ and HTV-/PCLV- mosquito lines

607 Mosquito lines persistently infected with PCLV and HTV or non-infected counterparts 608 were generated from F2 generations of wild mosquitoes. Three days after a blood 609 meal, F2 mated females were individually isolated in tubes containing a filter paper 610 and 0.5 cm of water and were allowed to lay eggs for 24 hours. Individual females were 611 collected, and the total RNA was isolated using TRIzol (Invitrogen) following the 612 standard protocol. Detection of HTV and PCLV was performed by RT-qPCR using the 613 primers described in **Supplementary Table 5**. Eggs corresponding to 5 female 614 mosquitoes infected with HTV, PCLV or both viruses were pooled prior to hatching. 615 Pools of eggs from 5 females negative for both viruses were pooled similarly to create 616 virus free lines. Subsequent detection of HTV and PCLV was performed to confirm 617 correct identification of lines. Lines generated from females carrying a single virus 618 (HTV or PCLV) were tested and found to carry both viruses. Therefore, only virus free 619 and double infected lines were expanded for two more generations for experiments 620 described in this work.

621

622 Artificial infection of naïve laboratory A. aegypti mosquitoes with HTV and PCLV

623 Extracts of naturally infected A. aegypti mosquitoes were used as source for HTV and 624 PCLV since we were not able to produce these viruses in cell culture. Viral stocks were 625 produced from pools of 15 A. aegypti naturally infected with HTV and PCLV or non-626 infected mosquitoes (virus-free controls), that were grinded using pestles in 1200 µL 627 of L-15 Leibovitz medium (Gibco) supplemented with 10% fetal bovine serum (FBS). 628 Samples were centrifuged at 3000 × g for 15 minutes at 4°C for clarification. 629 Supernatants were collected and passed through a 0.22 µm filter, aliquoted, and stored 630 at -80°C prior use. Infection with HTV and PCLV or mock control was performed by 631 microinjecting 69 nL of the extract into naïve laboratory mosquitoes (A. aegypti RedEye 632 strain) using a Nanoject II microinjector (Drummond).

633

634 Infection of Vero cells with HTV and PCLV

Filtered *A. aegypti* extracts (500 μL) containing HTV and PCLV (obtained as described
 above) were transferred into T-25 flasks containing 90% confluent Vero cells in non supplemented DMEM medium. After one hour of viral adsorption, 4.5 mL of DMEM

638 medium supplemented with penicillin/streptomycin and 10% of FBS were added to 639 cells, that were incubated at 37°C and 5% CO₂. 100 μ L aliquots of the supernatant 640 were collected during each passage on days 1, 3 and 5 after exposure to HTV and 641 PCLV. Virus in the supernatant was assessed by RT-qPCR. Vesicular stomatitis virus 642 (VSV) was added as a spike immediately before RNA extraction to be used as an 643 internal control.

644

645 **DENV and ZIKV propagation**

Viral isolates of DENV1 (MV09) and ZIKV (PE243/2015)⁶³ were propagated in C6/36 646 647 A. albopictus cells or Vero cells respectively. For DENV1 propagation, C6/36 cells were 648 maintained on L15 medium supplemented with 5% FBS (fetal bovine serum) and 1x 649 Antibiotic-Antimycotic (Gibco) as described²⁰. Cells were seeded to 70% confluence 650 and infected at a multiplicity of infection (MOI) of 0.01 and maintained for 6 to 9 days 651 at 28°C. For ZIKV propagation, a similar procedure was followed using Vero cells that 652 were maintained in DMEM medium supplemented with 10% FBS (fetal bovine serum) 653 and 1x Antibiotic-Antimycotic (Gibco). Vero cells were seeded to a confluence of 70-654 80% infected with ZIKV at a MOI of 0.01 and maintained for 6 days in culture. For both 655 viruses the supernatant was collected and clarified by centrifugation to generate virus 656 stocks that were kept at -80°C prior to use. Mock-infected supernatants used as 657 controls were prepared under same procedure without virus infection. Titration of 658 DENV1 was performed in BHK-21 cells while ZIKV was titrated in Vero cells, both using 659 the plaque assay method to determine viral titer.

660

661 **DENV and ZIKV infection in mosquitoes**

662 Natural infection in mosquitoes was performed using mice deficient for interferon-I and 663 interferon-II receptors as described ¹⁹. Briefly, infection in AG129 mice was established 664 by intraperitoneal injection of approximately 10⁶ PFU of DENV1 or 10⁶ PFU of ZIKV. 665 Infected mice were anaesthetized 3 days post infection (peak of viraemia) using 666 ketamine/xylazine (80/8 mg per kg) and placed on top of the netting-covered 667 containers with adult mosquito females. Mosquitoes were allowed to feed on mice for 668 30 minutes to 1 hour, alternating on the same animal between cages every 10 min if 669 two groups were to be compared. Viremia of each mouse was tested by RT-qPCR 670 immediately after feeding and the experiment would be discarded if a discrepancy of 671 more than 10x in viral load was observed. For infections by membrane feeding, 5-6 672 day old adult females were starved for 24h and fed with a mixture of blood and virus 673 supernatant containing 10⁷ PFU/mL of DENV serotype 1 utilizing a glass artificial 674 feeding system covered with pig intestine membrane as previously described ¹⁹. After

675 blood feeding, fully engorged females were selected and kept in standard rearing 676 conditions until collection at different time points. Mosquitoes infected by injection were 677 anesthetized with cold at 4°C and kept on ice during the whole procedure. Virus stocks 678 were diluted with L15 medium (Gibco) and injections were carried out using the 679 microinjector Nanoject II (Drummond) with a volume of 69 nL. Each individual mosquito 680 was injected with 16 PFU of DENV or ZIKV. Mosquitoes were harvested at different 681 days post injection for dissection and RNA extraction. Tissues (midguts, salivary 682 glands, and ovaries) were dissected in ice-cold 1x phosphate buffered saline (PBS) 683 containing 0.01% Triton X-100 (Sigma). Remnants of mosquito tissues were 684 considered as carcass, as illustrated in figure schemes. Tissues or mosquitoes were ground in TRIzol (Invitrogen) using glass beads as previously described ¹⁹ and kept 685 686 frozen at -80°C until RNA extraction as described above.

687

688 Mouse model for vectorial transmission of arboviruses

689 Groups of mosquitoes naturally infected with HTV and PCLV or PCLV/HTV-free 690 siblings were fed on the same ZIKV-infected AG129 mouse as described above. One 691 mouse was used per time point of transmission that was evaluated. Engorged females 692 were kept after feeding and, three days later, allowed to lay eggs for 48 hours in dark 693 cups containing 1 cm of water and a paper sheet attached to their walls. At 6-, 8-, and 694 12-days post feeding, 10 to 12 mosquito females of each group were allowed to feed 695 on naïve AG129 mice for up to 3 hours. Mice were bled 3 days after mosquito biting 696 and viral RNA levels were quantified by RT-qPCR. Viral loads were also quantified by 697 RT-gPCR in engorged mosquitoes from each group after feeding on a naïve mouse.

698

699 dsRNA mediated gene silencing

700 RNA transcription was performed using T7 Megascript kit (Ambion) following the 701 manufacturer's instructions. Briefly, template DNA containing T7 promoter sequences 702 in both 5' and 3' extremities was obtained by RT-PCR for dsAGO2 and dsH4, or by 703 PCR amplification from plasmid pDSAG (Addgene #62289) for dsGFP. Primer 704 sequences are provided in Supplementary Table 5. Adult 4-day-old females were 705 intrathoracically injected with 69 nl of a dsRNA solution (7.2 μ g μ l-1) diluted in 706 annealing buffer (20 mM Tris-HCl pH 7.5, 100 mM NaCl) using a nano-injector 707 Nanoject II (Drummond Scientific Company). Mosquitoes were allowed to recover for 708 48 h before further experiments. Once recovered, dsRNA-injected mosquitoes were 709 allowed to feed or were microinjected with virus following the procedures described 710 above.

712 **RT-qPCR and RT-PCR**

713 Total RNA extracted from individual mosquitoes or individual tissues were reverse 714 transcribed using M-MLV reverse transcriptase (RT) using random primers (hexamers) 715 for initiation. Negative controls were prepared following the same protocol without 716 adding reverse transcriptase. cDNA was subjected to polymerase chain reaction (RT-717 PCR) using the kit GoTag Hot Start Green PCR Master Mix (Promega) or quantitative 718 polymerase chain reaction (RT-qPCR) utilizing the kit Power SYBR[®] Green Master Mix 719 (Applied Biosystems) following manufacturer instructions. Results were expressed 720 using the 2- Δ Ct method relative to the endogenous control *rpL32*. Primer sequences 721 are listed in **Supplementary Table 5**. In experiments designed to differentiate the 722 expression of Histone H4 polyA transcripts versus non-polyA, cDNAs were 723 synthesized starting from the same amount of total RNA in two independent reactions 724 either using random hexamers or anchored oligo dT₂₂ as reverse transcription 725 initiators. Subsequent quantifications by RT-qPCR were performed as described 726 above.

727

728 Mathematical model

729 To investigate the effect of variations in the extrinsic incubation period (EIP) on the 730 cumulative incidence of DENV infections, we used a previously developed spatially 731 explicit, individual-based meta-population model ²⁸. Briefly, humans and mosquitoes 732 are defined with a unique state representing their current epidemiological status and, 733 in case of humans, their infection history. Human individuals are considered to be 734 either susceptible, exposed, infectious or recovered with respect to each serotype, 735 allowing up to four sequential infections. For mosquitoes, only the susceptible, 736 exposed, and infectious states of the epidemiologically relevant adult life-stage are 737 considered. The sizes of the respective populations are kept constant with deaths 738 being replaced by births. For simplicity, rather than accounting for seasonality through 739 changes in mosquito densities or temperature-dependent variations in EIP, this is done 740 here by varying daily mosquito biting rates and given as

741
$$a_{\nu}(t) = a_0(\beta + (1-\beta)\sin(t\pi/364)^4),$$

where *t* denotes time days and assuming a 364-day year. Both human and vector mortality rates are age-dependent, *i.e.*, the per capita risk of death is assumed to increase with age, which prevents individuals living beyond biologically reasonable ages. For computational efficiency, individuals' life expectancies are assigned at birth. Spatial structure is included by subdividing humans and mosquitoes into a spatially 747 organised sets of non-overlapping communities, where individuals mix 748 homogeneously. Mosquitoes preferentially bite humans in their own community and 749 adjacent communities but can also bite individuals of non-adjacent communities with 750 low probability to account for (daily) human mobility and associated long-distant 751 transmission events. Parameter values are listed in **Supplementary Table 6** and were 752 chosen to capture the qualitative dynamics of DENV in a high-transmission setting with 753 four co-circulating serotypes (DENV1-4).

754 The model exhibits pronounced demographic and epidemiological stochasticities that 755 arise from the fully probabilistic nature of state transitions and result in significant inter-756 annual oscillations in both disease incidence and relative prevalence of the four co-757 circulating serotypes. To investigate the effect of shortening, or lengthening the 758 extrinsic incubation period, we ran the model for a period of 100 years and recorded 759 the total annual number of infections for the last 50 years, discarding the transient 760 dynamics. Due to the stochastic nature of the model, we averaged the 50-year 761 cumulative incidence over five model runs for each value of the EIP. Mosquito and 762 human prevalence was equally evaluated over a 50-year time period but taken as the 763 proportion of infected individuals on a single day during the seasonal peak, resulting 764 in 50 individual data points.

765

766 **Poly-A selection, RNA library construction, and transcriptomic analysis**

767 RNA samples from individual mosquitoes were pooled and RNA quality was verified 768 using the 2100 Bioanalyzer system (Agilent). mRNA libraries were constructed using 769 the kits NEBNext[®] Poly(A) mRNA Magnetic Isolation Module and NEBNext[®] Ultra[™] II 770 Directional RNA Library Prep Kit for Illumina® (New England BioLabs inc.) following 771 the manufacturer protocol. Libraries were pooled and sequenced at the GenomEast 772 sequencing platform at the Institut de Génétique et de Biologie Moléculaire et Cellulaire 773 in Strasbourg, France. Sequenced reads with an average quality score above phred 774 25 had adaptors removed using Trimmomatic v0.39 and were further mapped to the 775 decoyed transcriptome of A. aegypti (Vectorbase release 48) using Salmon v1.3.0^{64,65}. 776 Quasi-mapping quantifications were imported into R v3.6.3 and data normalization was 777 performed using the packages EdgeR v3.28.1 and TMM ^{66,67}. Differentially expressed 778 genes were inferred using the exactTest function assuming a natural dispersion of 40% 779 in gene expression, whose input was used in the function decideTestsDGE. Fold 780 change plots were created using the package ggplot2 v3.3.6 and Euler diagrams 781 generated with the package venneuler v1.1-3. Heatmaps were generated using the R

packages tydiverse v1.3.1 and gplots v3.1.3. Ranked lists of gene expression for each comparison was used as input for Gene Set Enrichment Analysis (GSEA) ²⁸ using the R package fgsea v1.12.0 ⁶⁸ and in-house developed gene-sets comprising Gene Ontology annotation, pathways, and genes of interest. Sets with adjusted *p*-value < 0.1 were considered in our analysis.

787

788 Analysis of Histone H4 genes in *A. aegypti*

789 Alignment of putative histone H4 genes was performed using T-Coffee webserver ⁶⁹ 790 with the variant M-Coffee that allows to combine multiple outputs from different aligners 791 (MAFFT, Clustal and Muscle). The model (TN93 + I) for the phylogenetic reconstruction was defined using SMS within PhyMI server ^{70,71}. Finally, the Maximum-792 793 likelihood tree was constructed with PhyMI requiring 1000 bootstrap replicates and 794 edited using iTOL server ⁶¹. The *Histone H4* polyadenilation signature was defined 795 according to the presence of a canonical polyadenylation signal (AAUAAA) as 796 previously described ³⁵. The histone H4 expression heatmap shown in **Extended Data** 797 Fig. 8d was produced in R v4.0.2 using the package gplots v3.1.3 and show transcripts 798 TPM counts and normalized by Z-score (row) and sorted according to the phylogenetic 799 tree. Libraries were obtained at the Sequence Read Archive (SRA/NCBI) and 800 accession numbers are shown.

801

802 Statistical analyses

803 Evaluation of statistical significance was performed using GraphPad Prism 9.0 or R-804 cran v3.3.1 software unless stated otherwise. Viral loads of RT-qPCR positive-only 805 individual mosquito/tissues were log transformed and subjected to Mann-Whitney U 806 test. Pairwise comparisons of virus infection prevalence was evaluated by Fisher's 807 exact test or sequentially by a generalized linear model considering the interaction 808 between time points and their prevalence, followed by ANOVA type II using the car 809 package v3.1-0 on R. Presence and absence of DENV in mosquitoes was modelled with univariate and multivariate zero-inflated binomial model (ZIB) 72,73, since 95% of 810 811 the collections are zeroes. The covariate or covariates (in case of multivariate model) 812 was/were the same for the logit and logistic parts of the model. In particular, we 813 considered PCLV, HTV and their interaction. Model selection was carried out by AIC 814 and BIC comparison ⁷⁴. The Vuong test was conducted a priori to test if the zero inflated 815 binomial model was statistically significant and better (in terms of AIC and BIC) than a 816 non-zero inflated model. Data were analysed using R and the 'pscl' package v1.5.5⁷⁵. 817 Finally, we tested the presence of spatial autocorrelation in the two viruses - via

818 variogram analyses ²⁰ - but no significant autocorrelations have been found (results

819 not reported).

821 **Data availability**

- Small RNA and transcriptome libraries from this study have been deposited in the
 Sequence Read Archive (SRA) at NCBI (project accession PRJNA722589).
 Sequences spanning the RdRP region from newly discovered viruses were deposited
 in GenBank under accession MZ556103-MZ556111. Accession numbers for small
 RNA libraries are provided in **Supplementary Table 1**.
- 827

828 Code availability

All scripts used in this work were deposited in GitHub and can be accessed on https://github.com/ericgdp/sRNA-virome code version 1.0.

831

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838

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865

866 Contributions

- 867 R.P.O., E.R.G.R.A., J.-L.I. and J.T.M conceived the project. Methodology: R.P.O.,
- 868 Y.M.H.T., E.R.G.R.A., J.P.P.A., J.N.A., L.S. and J.T.M. designed the experiments and
- 869 performed computational analysis. R.P.O., Y.M.H.T., I.J.S.F., F.V.F., A.G.A.F.,
- 870 S.C.G.A., A.T.S.S., K.P.R.S., A.P.P.V., A.B. performed experiments. C.H.T., M.D.,
- 871 A.G., C.P., J.O.N., T.M.V., C.J.M.K., M.A.W., A.L.C.C., M.T.P., M.C.P.P., M.L.N.,
- V.A.S., R.N.M., M.A.Z.B., B.P.D., E.G.K., E.M. provided samples and reagents. M.R.
- performed the mathematical modelling. R.P.O., Y.M.H.T., E.R.G.R.A., J.-L.I. and
- J.T.M. wrote the manuscript. All authors read and contributed to manuscript editing.
- 875

876 **Competing interests**

- 877 Authors declare that they have no competing interests.
- 878

879 Figure Legends

880

Fig. 1 The virome of *A. aegypti* and *A. albopictus* mosquitoes.

882 a, World map indicating sites of mosquito collection. Pie charts show the proportion of 883 A. aegypti and A. albopictus in red and blue, respectively, at each collection site. Adult 884 mosquitoes were captured either using traps or human baits. Laboratory strains of 885 mosquitoes analyzed in this work are indicated at the bottom. b, Overview of our 886 analysis pipeline. Captured mosquitoes were morphologically identified by species and 887 stored in a biobank of RNA samples, that were pooled to prepare small RNA libraries 888 for high-throughput sequencing. Using our metagenomic pipeline, assembled contigs 889 were classified into viral and non-viral sequences based on similarity against reference 890 databases. Sequences that lack any similarity to known references are indicated as 891 unknown. c, Individual results from our metagenomic analysis for each of the 91 small 892 RNA libraries in this study. Bars indicate the total number of contigs and the proportion 893 of viral, non-viral and unknown contigs per library.

894

Fig. 2 A highly diverse and distinct collection of viruses in *A. aegypti* and *A. albopictus.*

897 a, Overview of the strategy for manual curation of viral contigs to confirm the origin and 898 remove sequences potentially derived from EVEs. Curation consisted of BLAST 899 search for similar viral sequences, inspection of predicted ORF structure including 900 continuity and extension throughout each contig, and evaluation of the small RNA 901 profile for the identification of signatures of siRNA production (symmetrical 902 accumulation of RNAs with 20-23 nt length that mapped to each strand with no 5' base 903 preference) or piRNA production (accumulation of 24-29 nt length RNAs with 5' U 904 preference); and overall contig coverage by small RNAs. Remaining viral contigs were 905 clustered by sequence similarity and co-ocurrence to identify groups of contigs that 906 belong to same viruses. Re-assembly was performed within these groups and resulting 907 contigs were analyzed for the presence of domains. Potential polymerases identified 908 were used to classify viruses based on sequence similarity and phylogeny. b, Host, 909 virus genomic organization, family, and closest reference on GenBank identified by 910 BLAST similarity searches for each of the 12 viruses identified in our datasets. New 911 viral species are indicated in red while previously known viruses are in black. 912 Sequence similarity and accession number according to the closest viral sequence at 913 the nucleotide (nt) or protein (aa) levels are indicated. c, Viral load shown as a heatmap 914 for each of the twelve viruses in mosquito populations from each collection site or 915 laboratory strains. In the heatmap, white indicates absence of a virus and NA indicates 916 absence of samples from a given location. Prevalence of each virus is shown on the 917 right as number of samples with detectable virus over the total. Number of individuals 918 per pool and number of species per collection site are described on the 919 Supplementary Table 1. d, Pie charts represent the overall burden of virus and viral 920 diversity for A. aegypti and A. albopictus populations in each collection site across the 921 word. X indicates that no viral contigs have been identified.

922

923 Fig. 3 Concurrent detection of HTV, PCLV and DENV.

a, Location of the study site, the city of Caratinga in the Southeast of Brazil. b,
 Prevalence of individual and co-infections by OVV, HTV and PCLV tested by RT qPCR. c, Prevalence of DENV, HTV and PCLV separately in individual *A. aegypti* mosquitoes and human blood samples accessed by RT-qPCR. d, Monthly prevalence
 of HTV and PCLV separately in individual *A. aegypti* mosquitoes. e, Prevalence of HTV

and PCLV co-infection in DENV infected (DENV+) and DENV non-infected (DENV-)
mosquitoes. Statistical significance was determined by two-tailed Fisher's exact test.
f, Likelihood of DENV infection in mosquitoes carrying HTV or PCLV and HTV shown
by odds ratio.

933

934 Fig. 4 Effects of insect specific viruses on DENV and ZIKV transmission.

935 a-g, Strategy to evaluate the interference of HTV and PCLV on DENV (a-c) or ZIKV 936 (d-g) infection in wild mosquito populations. HTV/PCLV infected or virus free 937 mosquitoes were exposed to DENV (a) or ZIKV (d) on a blood meal. Viral loads and 938 prevalence of infection were measured in the midgut (b,e), carcass (c,f) and salivary 939 glands (g) of mosquitoes at the indicated days post feeding (d.p.f.). Pie charts below 940 each group indicate the prevalence of DENV or ZIKV infection at each time point. h, 941 Strategy to compare the ability of mosquitoes with or without HTV and PCLV to 942 transmit ZIKV to the susceptible AG129 mouse model. i, Viremia in mice was 943 determined 3 days after exposure to mosquito bites, comparing animals that were 944 bitten by mosquitoes carrying or not HTV and PCLV at 6, 8 and 12 days post oral 945 infection. ND - not detected. Pie charts below each group indicate the prevalence of 946 infection. (i) ZIKV RNA levels in mosquitoes used in the transmission experiment in i. 947 NS – non-significant. k-I Spatially explicit, individual-based meta-population model 948 showing the effect of EIP duration on the number of human infections (k) and virus 949 prevalence in mosquitoes (I) taken as a proportion of infected individuals on a single 950 day during the seasonal peak over a 50 year period (*n*=50). In box plots of **b**, **c**, **e**, **f**, **j**, 951 and *I*, boxes show the second and third interguartile ranges divided by the median 952 while whiskers represent maximum and minimum values. Statistical significance was 953 determined by two-tailed Mann–Whitney U-test. In k, lines represent the averaged 954 cumulative incidence over five model runs for each value of the EIP, while shadows 955 depict confidence intervals. Numbers of infected midguts (**b**,**e**), carcasses (**c**,**f**), or 956 whole mosquitoes (j) over the total number tested are indicated above each column. 957 Each dot represents an individual sample. Statistical significance of the prevalence of 958 infection was determined in **b**, **c**, **e**, **f**, **i**, and **j** by two-tailed Fisher's exact test or, in **g**, 959 using a binomial generalized linear model followed by ANOVA type II testing for time 960 points as factor.

961

Fig. 5 HTV and PCLV regulate the expression of proviral histone H4 during DENV infection.

964 a, Overall strategy to identify biological pathways associated with HTV and PCLV 965 interaction with DENV. HTV/PCLV infected and virus free wild mosquito populations 966 were allowed to feed on DENV-infected mice. The transcriptome of DENV infected and 967 non-infected mosquitoes from HTV/PCLV and virus free groups were analyzed 968 separately at 8- and 14-days post feeding. Gene Set Enrichment Analysis (GSEA) was 969 performed for each comparison. b. Biological processes significantly enriched 970 (adjusted *p*-value < 0.01) in the comparisons of DENV-infected versus non-infected 971 mosquitoes and PCLV/HTV infected versus virus free controls are shown. c, Overlap 972 of leading-edge genes belonging to the 4 biological processes enriched in at least 6 973 out of 8 comparisons. Size of each circle represents the number of leading-edge 974 genes. Histogram shows that histone genes represent the majority in the leading edge 975 of significantly enriched processes. d-e, Histone H4 levels in the carcass of mosquitoes 976 at 2, 4, and 8 d.p.i. with ZIKV in wild mosquitoes carrying HTV/PCLV or virus free 977 controls (e). f-g. Differential expression of histone H4 levels in the carcass of 978 mosquitoes in the presence of HTV and PCLV is only observed in DENV infected 979 individuals. h-j, Silencing of histone H4 mRNA in adult mosquitoes exposed to DENV. 980 DENV infection was analyzed at 4 and 8 d.p.f. in the midgut (i) and carcass (j) of 981 silenced (dsH4) and control (dsGFP) mosquitoes. Pie charts below each group

982 indicate the prevalence of DENV or ZIKV infection. d.p.f. - days post feeding. d.p.i. -983 days post injection, NS - non-significant. In box plots of e, g, h, and j, boxes show the 984 second and third interquartile ranges divided by the median while whiskers represent 985 maximum and minimum values. Numbers of infected samples over the total number 986 tested are indicated above each column. Statistical significance was determined by 987 two-tailed one-way ANOVA with Tukey's correction for multiple comparisons (e,g) or 988 by two-tailed Mann-Whitney U-test (h,i). Each dot represents an individual sample. 989 Statistical significance of prevalence of infection was determined by two-tailed Fisher's 990 exact test.

992 R e	eferences
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994	1. Weaver, S. C., Charlier, C., Vasilakis, N. & Lecuit, M. Zika, Chikungunya, and
995	Other Emerging Vector-Borne Viral Diseases. Annu Rev Med 69, 395–408 (2018).

- Bhatt, S. *et al.* The global distribution and burden of dengue. *Nature* 496, 504–507
 (2013).
- 3. Messina, J. P. *et al.* Global spread of dengue virus types: mapping the 70 year
 history. *Trends Microbiol* 22, 138–146 (2014).
- 4. Franklinos, L. H. V., Jones, K. E., Redding, D. W. & Abubakar, I. The effect of
 global change on mosquito-borne disease. *Lancet Infect Dis* 19, e302–e312
 (2019).
- 1003 5. Kraemer, M. U. G. *et al.* Past and future spread of the arbovirus vectors Aedes
 1004 aegypti and Aedes albopictus. *Nat Microbiol* 4, 854–863 (2019).
- 1005 6. Cromwell, E. A. *et al.* The relationship between entomological indicators of Aedes
 1006 aegypti abundance and dengue virus infection. *PLoS Negl Trop Dis* **11**, e0005429
 1007 (2017).
- 1008 7. de Almeida, J. P., Aguiar, E. R., Armache, J. N., Olmo, R. P. & Marques, J. T. The
 1009 virome of vector mosquitoes. *Curr Opin Virol* 49, 7–12 (2021).
- 1010 8. Aguiar, E. R. G. R. et al. Sequence-independent characterization of viruses based
- 1011 on the pattern of viral small RNAs produced by the host. *Nucleic Acids Res* 43,
 1012 6191–6206 (2015).
- 1013 9. Boyles, S. M. *et al.* Under-the-Radar Dengue Virus Infections in Natural
 1014 Populations of Aedes aegypti Mosquitoes. *mSphere* 5, (2020).
- 1015 10. Ramos-Nino, M. E. et al. High prevalence of Phasi Charoen-like virus from wild-
- 1016 caught Aedes aegypti in Grenada, W.I. as revealed by metagenomic analysis.
- 1017 *PLoS One* **15**, e0227998 (2020).

- 1018 11. Shi, C. *et al.* Stable distinct core eukaryotic viromes in different mosquito species
 1019 from Guadeloupe, using single mosquito viral metagenomics. *Microbiome* 7, 121
 1020 (2019).
- 1021 12. Zakrzewski, M. *et al.* Mapping the virome in wild-caught Aedes aegypti from Cairns
 1022 and Bangkok. *Sci Rep* 8, 4690 (2018).
- 1023 13. Patterson, E. I., Villinger, J., Muthoni, J. N., Dobel-Ober, L. & Hughes, G. L.
- 1024 Exploiting insect-specific viruses as a novel strategy to control vector-borne 1025 disease. *Curr Opin Insect Sci* **39**, 50–56 (2020).
- 1026 14. Vasilakis, N. & Tesh, R. B. Insect-specific viruses and their potential impact on
 1027 arbovirus transmission. *Curr Opin Virol* **15**, 69–74 (2015).
- 1028 15. Aguiar, E. R. G. R., Olmo, R. P. & Marques, J. T. Virus-derived small RNAs:
 1029 molecular footprints of host-pathogen interactions. *Wiley Interdiscip Rev RNA* 7,
 1030 824–837 (2016).
- 1031 16. Morazzani, E. M., Wiley, M. R., Murreddu, M. G., Adelman, Z. N. & Myles, K. M.
- Production of virus-derived ping-pong-dependent piRNA-like small RNAs in the
 mosquito soma. *PLoS Pathog* 8, e1002470 (2012).
- 1034 17. Myles, K. M., Wiley, M. R., Morazzani, E. M. & Adelman, Z. N. Alphavirus-derived
- small RNAs modulate pathogenesis in disease vector mosquitoes. *Proc Natl Acad Sci U S A* 105, 19938–19943 (2008).
- 1037 18. Frangeul, L., Blanc, H., Saleh, M.-C. & Suzuki, Y. Differential Small RNA
 1038 Responses against Co-Infecting Insect-Specific Viruses in Aedes albopictus
 1039 Mosquitoes. *Viruses* 12, (2020).
- 1040 19. Olmo, R. P. *et al.* Control of dengue virus in the midgut of Aedes aegypti by ectopic
- 1041 expression of the dsRNA-binding protein Loqs2. *Nat Microbiol* 3, 1385–13931042 (2018).
- 1043 20. Sedda, L. *et al.* The spatial and temporal scales of local dengue virus transmission
- 1044 in natural settings: a retrospective analysis. *Parasit Vectors* **11**, 79 (2018).

- 1045 21. Schultz, M. J., Frydman, H. M. & Connor, J. H. Dual Insect specific virus infection
 1046 limits Arbovirus replication in Aedes mosquito cells. *Virology* **518**, 406–413 (2018).
- 1047 22. Fredericks, A. C. et al. Aedes aegypti (Aag2)-derived clonal mosquito cell lines
- reveal the effects of pre-existing persistent infection with the insect-specific
 bunyavirus Phasi Charoen-like virus on arbovirus replication. *PLoS Negl Trop Dis* **10**50 **13**, e0007346 (2019).
- 1051 23. Sedda, L., Taylor, B. M., Eiras, A. E., Marques, J. T. & Dillon, R. J. Using the
 1052 intrinsic growth rate of the mosquito population improves spatio-temporal dengue
 1053 risk estimation. *Acta Trop* 208, 105519 (2020).
- 105424. Lin, J.-J. *et al.* Aggressive organ penetration and high vector transmissibility of1055epidemic dengue virus-2 Cosmopolitan genotype in a transmission mouse model.
- 1056 PLoS Pathog **17**, e1009480 (2021).
- 1057 25. Sun, P. *et al.* A mosquito salivary protein promotes flavivirus transmission by
 1058 activation of autophagy. *Nat Commun* **11**, 260 (2020).
- 1059 26. Lourenço, J. & Recker, M. Natural, persistent oscillations in a spatial multi-strain
 1060 disease system with application to dengue. *PLoS Comput Biol* 9, e1003308 (2013).
- 1061 27. Lourenço, J. *et al.* Epidemiological and ecological determinants of Zika virus
 1062 transmission in an urban setting. *Elife* 6, e29820 (2017).
- 1063 28. Subramanian, A. *et al.* Gene set enrichment analysis: a knowledge-based
- approach for interpreting genome-wide expression profiles. *Proc Natl Acad Sci U*S A 102, 15545–15550 (2005).
- 1066 29. Flaus, A., Downs, J. A. & Owen-Hughes, T. Histone isoforms and the oncohistone
 1067 code. *Curr Opin Genet Dev* 67, 61–66 (2021).
- 30. Lyons, S. M. *et al.* A subset of replication-dependent histone mRNAs are
 expressed as polyadenylated RNAs in terminally differentiated tissues. *Nucleic Acids Res* 44, 9190–9205 (2016).

- 1071 31. Baidaliuk, A. *et al.* Cell-Fusing Agent Virus Reduces Arbovirus Dissemination in
 1072 *Aedes aegypti* Mosquitoes *In Vivo. J Virol* **93**, e00705-19, /jvi/93/18/JVI.007051073 19.atom (2019).
- 32. Blitvich, B. J. & Firth, A. E. Insect-specific flaviviruses: a systematic review of their
 discovery, host range, mode of transmission, superinfection exclusion potential
 and genomic organization. *Viruses* 7, 1927–1959 (2015).
- 1077 33. Colpitts, T. M., Barthel, S., Wang, P. & Fikrig, E. Dengue virus capsid protein binds
 1078 core histones and inhibits nucleosome formation in human liver cells. *PLoS One*1079 6, e24365 (2011).
- Mourão, D. et al. A histone-like motif in yellow fever virus contributes to viral
 replication. http://biorxiv.org/lookup/doi/10.1101/2020.05.05.078782 (2020)
 doi:10.1101/2020.05.05.078782.
- 35. Girardi, E. *et al.* Histone-derived piRNA biogenesis depends on the ping-pong
 partners Piwi5 and Ago3 in Aedes aegypti. *Nucleic Acids Res* 45, 4881–4892
 (2017).
- 36. Varjak, M. *et al.* Aedes aegypti Piwi4 Is a Noncanonical PIWI Protein Involved in
 Antiviral Responses. *mSphere* 2, e00144-17 (2017).
- 37. Parry, R. & Asgari, S. Aedes Anphevirus: an Insect-Specific Virus Distributed
 Worldwide in *Aedes aegypti* Mosquitoes That Has Complex Interplays with *Wolbachia* and Dengue Virus Infection in Cells. *J Virol* **92**, e00224-18,
 /jvi/92/17/e00224-18.atom (2018).
- 38. Zhang, G., Asad, S., Khromykh, A. A. & Asgari, S. Cell fusing agent virus and
 dengue virus mutually interact in Aedes aegypti cell lines. *Sci Rep* 7, 6935 (2017).
- 39. Nasar, F., Erasmus, J. H., Haddow, A. D., Tesh, R. B. & Weaver, S. C. Eilat virus
 induces both homologous and heterologous interference. *Virology* 484, 51–58
- 1096 (2015).
- 40. Kenney, J. L., Solberg, O. D., Langevin, S. A. & Brault, A. C. Characterization of a
 novel insect-specific flavivirus from Brazil: potential for inhibition of infection of

- arthropod cells with medically important flaviviruses. *J Gen Virol* 95, 2796–2808
 (2014).
- 41. Romo, H., Kenney, J. L., Blitvich, B. J. & Brault, A. C. Restriction of Zika virus
 infection and transmission in *Aedes aegypti* mediated by an insect-specific
 flavivirus. *Emerging Microbes & Infections* **7**, 1–13 (2018).
- 1104 42. Goenaga, S. et al. Potential for Co-Infection of a Mosquito-Specific Flavivirus,
- 1105 Nhumirim Virus, to Block West Nile Virus Transmission in Mosquitoes. *Viruses* 7,
 1106 5801–5812 (2015).
- 1107 43. Alefelder, S., Patel, B. K. & Eckstein, F. Incorporation of terminal
 1108 phosphorothioates into oligonucleotides. *Nucleic Acids Res* 26, 4983–4988
 1109 (1998).
- 44. Marques, J. T. *et al.* Loqs and R2D2 act sequentially in the siRNA pathway in
 Drosophila. *Nat Struct Mol Biol* **17**, 24–30 (2010).
- 45. Martin, M. Cutadapt removes adapter sequences from high-throughput
 sequencing reads. *EMBnet j.* **17**, 10 (2011).
- 46. Matthews, B. J. *et al.* Improved reference genome of Aedes aegypti informs
 arbovirus vector control. *Nature* 563, 501–507 (2018).
- 1116 47. Chen, X.-G. et al. Genome sequence of the Asian Tiger mosquito, Aedes
- albopictus, reveals insights into its biology, genetics, and evolution. *Proc Natl Acad*
- 1118 *Sci U S A* **112**, E5907-5915 (2015).
- 48. Langmead, B., Trapnell, C., Pop, M. & Salzberg, S. L. Ultrafast and memoryefficient alignment of short DNA sequences to the human genome. *Genome Biol*
- 1121 **10**, R25 (2009).
- 49. Zerbino, D. R. Using the Velvet de novo assembler for short-read sequencing
 technologies. *Curr Protoc Bioinformatics* Chapter 11, Unit 11.5 (2010).
- 1124 50. Bankevich, A. *et al.* SPAdes: a new genome assembly algorithm and its 1125 applications to single-cell sequencing. *J Comput Biol* **19**, 455–477 (2012).

- 1126 51. O'Leary, N. A. et al. Reference sequence (RefSeq) database at NCBI: current
- status, taxonomic expansion, and functional annotation. *Nucleic Acids Res* 44,
 D733-745 (2016).
- 1129 52. Altschul, S. F., Gish, W., Miller, W., Myers, E. W. & Lipman, D. J. Basic local
 1130 alignment search tool. *J Mol Biol* 215, 403–410 (1990).
- 1131 53. Fu, L., Niu, B., Zhu, Z., Wu, S. & Li, W. CD-HIT: accelerated for clustering the next-
- generation sequencing data. *Bioinformatics* **28**, 3150–3152 (2012).
- 54. Aguiar, E. R. G. R. *et al.* A single unidirectional piRNA cluster similar to the
 flamenco locus is the major source of EVE-derived transcription and small RNAs
 in Aedes aegypti mosquitoes. *RNA* 26, 581–594 (2020).
- 1136 55. Whitfield, Z. J. *et al.* The Diversity, Structure, and Function of Heritable Adaptive
 1137 Immunity Sequences in the Aedes aegypti Genome. *Curr Biol* 27, 3511-3519.e7
 1138 (2017).
- 1139 56. Palatini, U. *et al.* Comparative genomics shows that viral integrations are abundant

and express piRNAs in the arboviral vectors Aedes aegypti and Aedes albopictus.

1141 BMC Genomics **18**, 512 (2017).

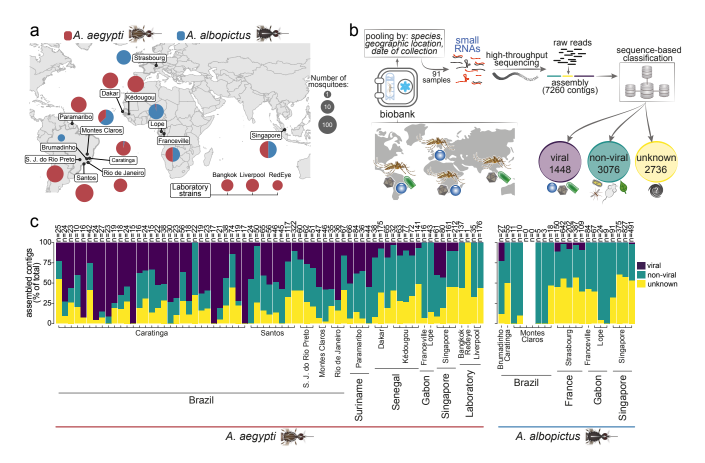
- 1142 57. Katoh, K., Misawa, K., Kuma, K. & Miyata, T. MAFFT: a novel method for rapid
 1143 multiple sequence alignment based on fast Fourier transform. *Nucleic Acids Res*
- **30**, 3059–3066 (2002).
- 1145 58. Miller, M. A., Pfeiffer, W. & Schwartz, T. The CIPRES science gateway: a 1146 community resource for phylogenetic analyses. in *Proceedings of the 2011*
- 1147 TeraGrid Conference on Extreme Digital Discovery TG '11 1 (ACM Press, 2011).
- 1148 doi:10.1145/2016741.2016785.
- 59. Darriba, D. *et al.* ModelTest-NG: A New and Scalable Tool for the Selection of DNA
 and Protein Evolutionary Models. *Mol Biol Evol* **37**, 291–294 (2020).
- 1151 60. Kumar, S., Stecher, G., Li, M., Knyaz, C. & Tamura, K. MEGA X: Molecular
- 1152 Evolutionary Genetics Analysis across Computing Platforms. *Mol Biol Evol* **35**,
- 1153 1547–1549 (2018).

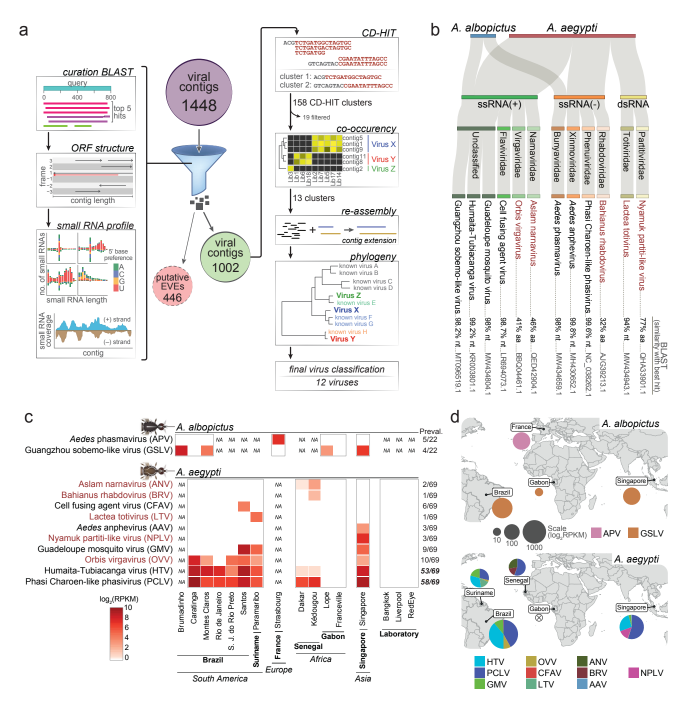
- 1154 61. Letunic, I. & Bork, P. Interactive Tree Of Life (iTOL) v4: recent updates and new
 1155 developments. *Nucleic Acids Res* 47, W256–W259 (2019).
- 1156 62. Barletta, A. B. F. et al. Microbiota activates IMD pathway and limits Sindbis

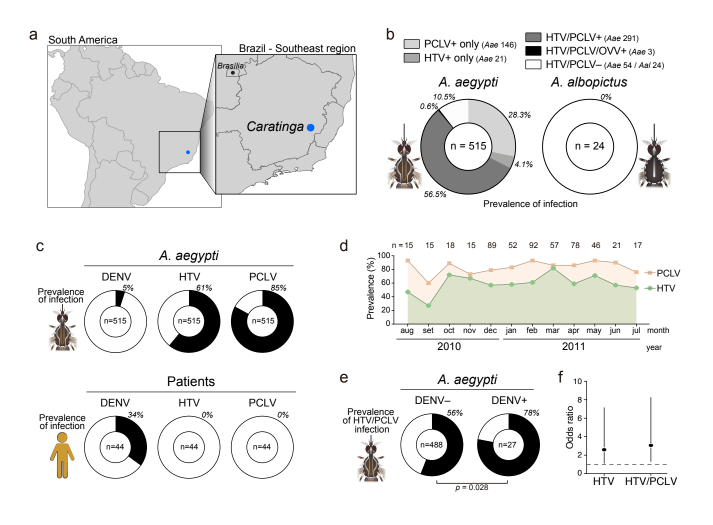
1157 infection in Aedes aegypti. *Parasit Vectors* **10**, 103 (2017).

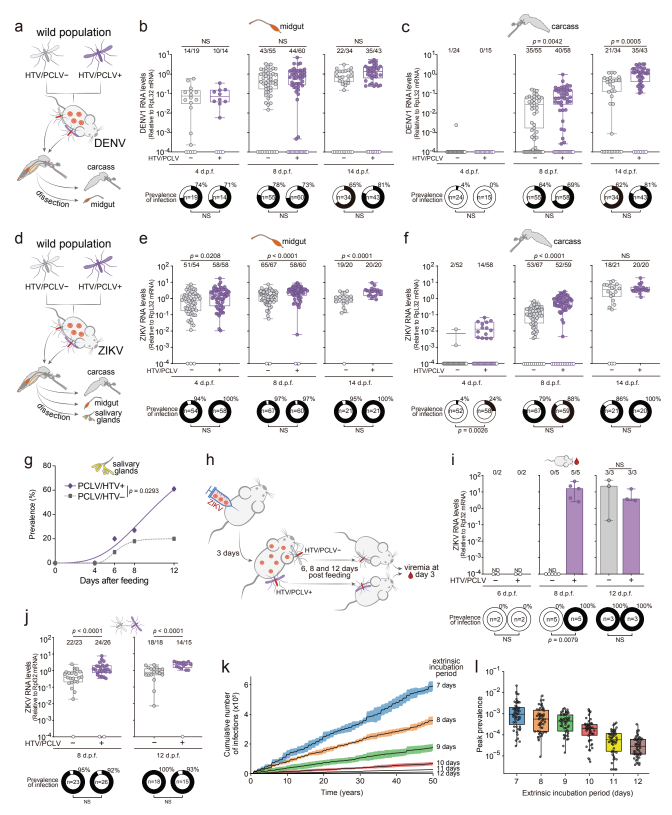
- 1158 63. Donald, C. L. *et al.* Full Genome Sequence and sfRNA Interferon Antagonist
 1159 Activity of Zika Virus from Recife, Brazil. *PLoS Negl Trop Dis* **10**, e0005048 (2016).
- 64. Bolger, A. M., Lohse, M. & Usadel, B. Trimmomatic: a flexible trimmer for Illumina
 sequence data. *Bioinformatics* **30**, 2114–2120 (2014).
- 1162 65. Patro, R., Duggal, G., Love, M. I., Irizarry, R. A. & Kingsford, C. Salmon provides
- fast and bias-aware quantification of transcript expression. *Nat Methods* 14, 417–
 419 (2017).
- 66. Robinson, M. D. & Oshlack, A. A scaling normalization method for differential
 expression analysis of RNA-seq data. *Genome Biol* **11**, R25 (2010).
- 67. Robinson, M. D., McCarthy, D. J. & Smyth, G. K. edgeR: a Bioconductor package
 for differential expression analysis of digital gene expression data. *Bioinformatics*
- **26**, 139–140 (2010).
- 1170 68. Korotkevich, G. et al. Fast gene set enrichment analysis.
 1171 http://biorxiv.org/lookup/doi/10.1101/060012 (2016) doi:10.1101/060012.
- 1172 69. Di Tommaso, P. *et al.* T-Coffee: a web server for the multiple sequence alignment
 1173 of protein and RNA sequences using structural information and homology
 1174 extension. *Nucleic Acids Res* **39**, W13-17 (2011).
- 1175 70. Guindon, S. *et al.* New algorithms and methods to estimate maximum-likelihood
 1176 phylogenies: assessing the performance of PhyML 3.0. *Syst Biol* **59**, 307–321
 1177 (2010).
- 1178 71. Lefort, V., Longueville, J.-E. & Gascuel, O. SMS: Smart Model Selection in PhyML.
 1179 *Mol Biol Evol* 34, 2422–2424 (2017).
- 1180 72. Lambert, D. Zero-Inflated Poisson Regression, with an Application to Defects in
 1181 Manufacturing. *Technometrics* 34, 1 (1992).

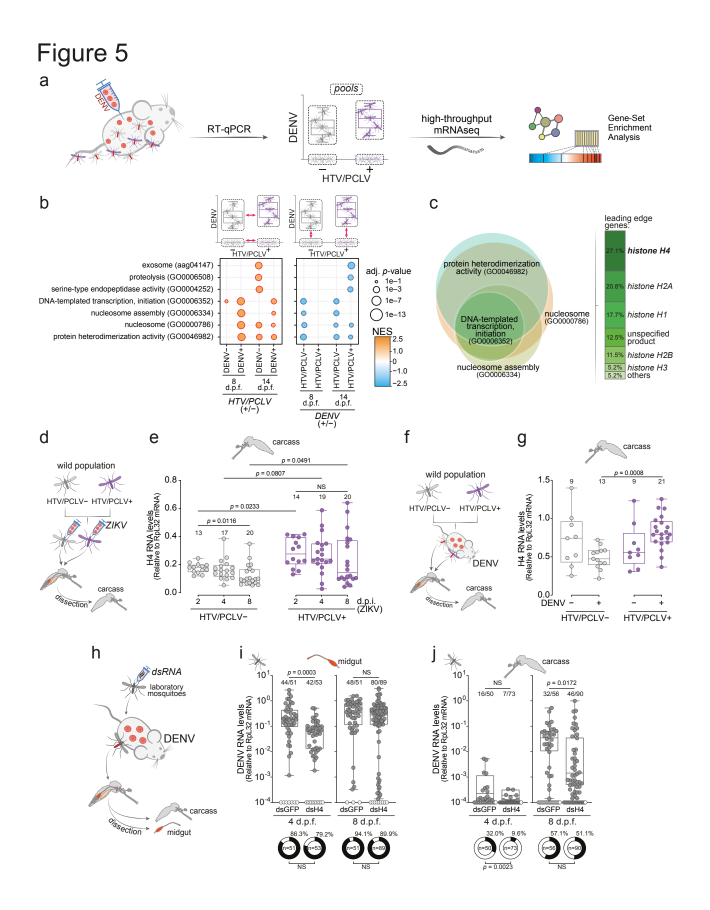
- 1182 73. Hilbe, J. M. Negative Binomial Regression. (2007).
- 1183 74. Cameron, A. C., Cameron, A. C., & Cambridge University Press. Regression
- 1184 *analysis of count data.* (Cambridge University Press, 1998).
- 1185 75. Zeileis, A., Kleiber, C. & Jackman, S. Regression Models for Count Data in *R. J.*
- 1186 Stat. Soft. **27**, (2008).
- 1187

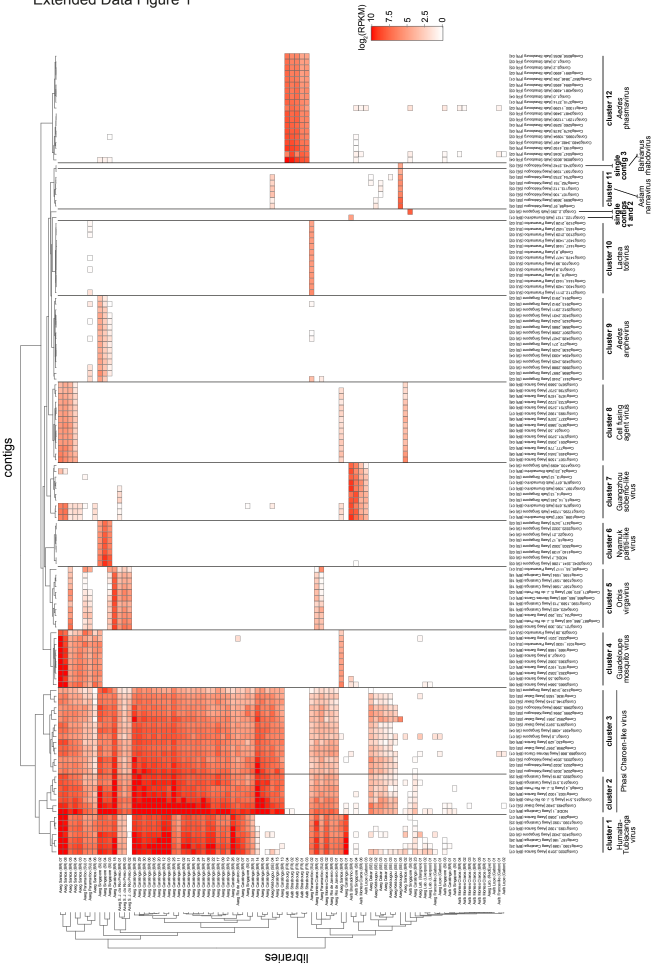


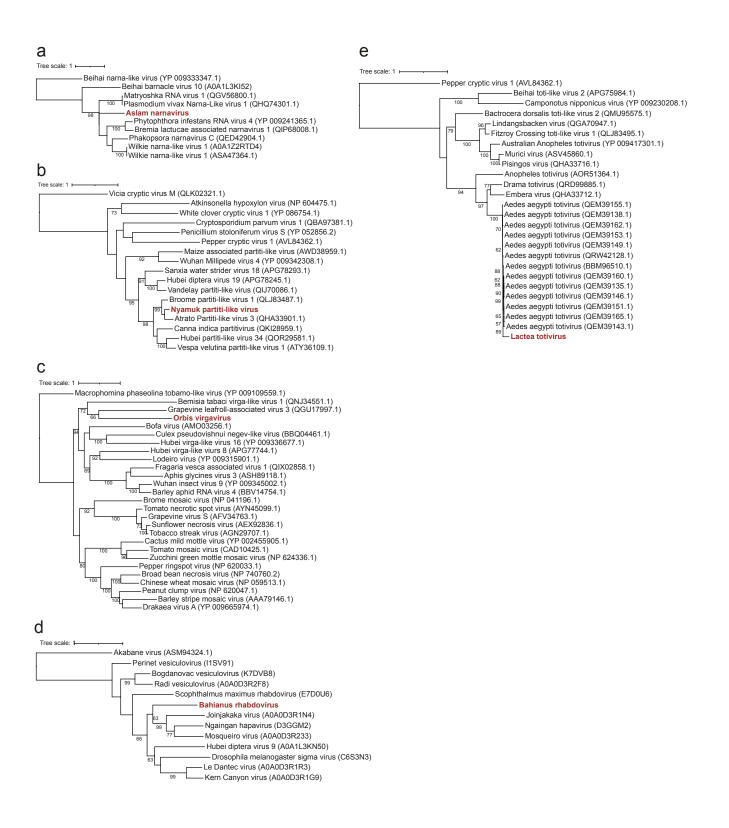


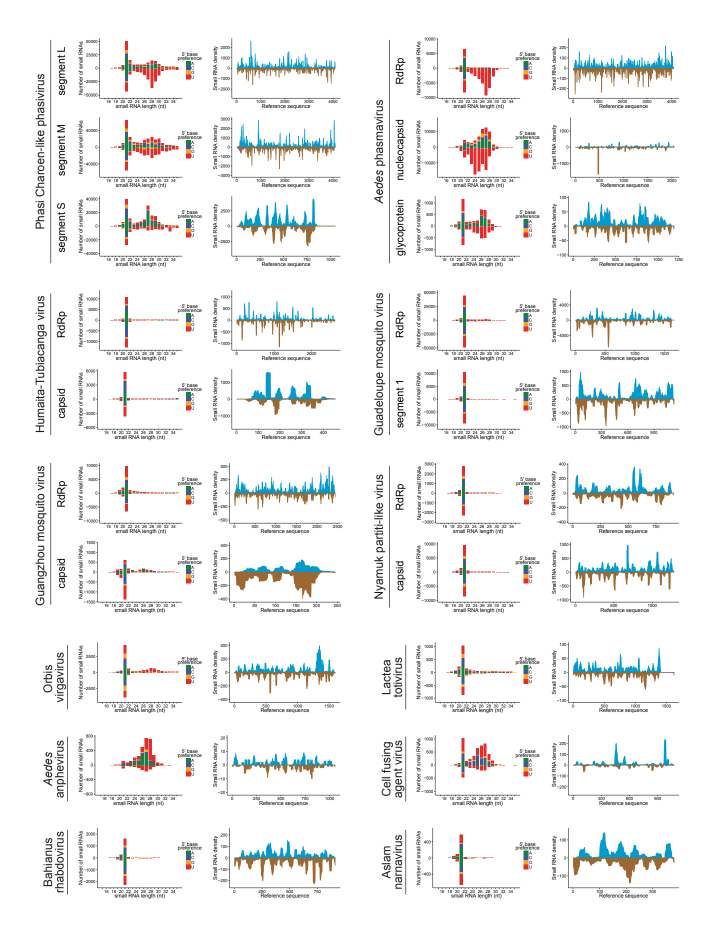


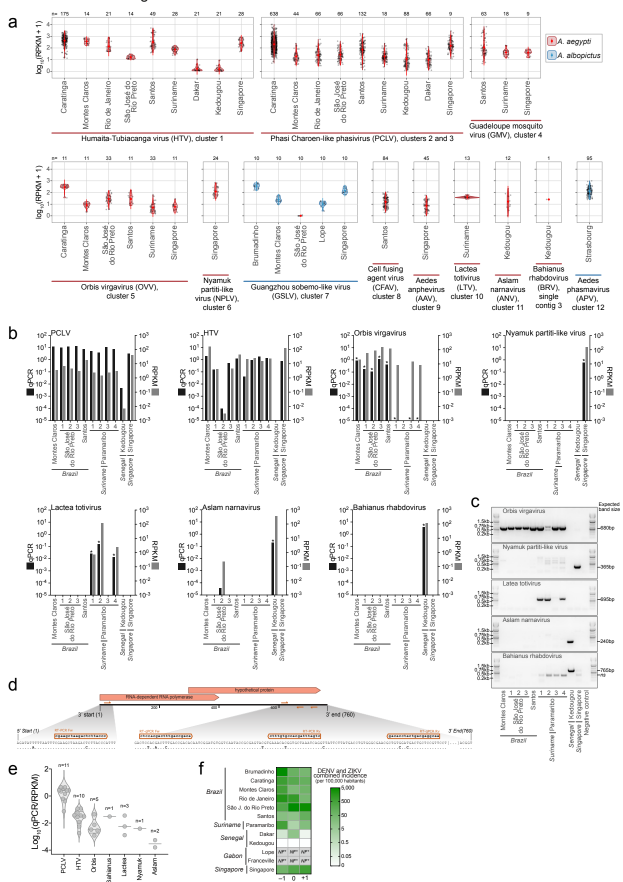












-1 0 +1 Year of mosquito capture

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