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1	Aedes anphevirus (AeAV): an insect-specific virus distributed worldwide in Aedes					
2	aegypti mosquitoes that has complex interplays with Wolbachia and dengue virus					
3	infection in cells					
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11	Running title: Anphevirus from Aedes aegypti					
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14						
15						
16						
17	Word count (abstract): 249					
18	Word count (importance): 161					
19	Word count (text): 5334					
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30 Abstract

Insect specific viruses (ISVs) of the yellow fever mosquito Aedes aegypti have been demonstrated 31 to modulate transmission of arboviruses such as dengue virus (DENV) and West Nile virus by the 32 mosquito. The diversity and composition of the virome of Ae. aegypti, however, remains poorly 33 understood. In this study, we characterised Aedes anphevirus (AeAV), a negative-sense RNA virus 34 from the order Mononegavirales. AeAV identified from Aedes cell lines were infectious to both Ae. 35 aegypti and Aedes albopictus cells, but not to three mammalian cell lines. To understand the 36 incidence and genetic diversity of AeAV, we assembled 17 coding-complete and two partial 37 genomes of AeAV from available RNA-Seq data. AeAV appears to transmit vertically and be 38 present in laboratory colonies, wild-caught mosquitoes and cell lines worldwide. Phylogenetic 39 analysis of AeAV strains indicates that as the Ae. aegypti mosquito has expanded into the 40 41 Americas and Asia-Pacific, AeAV has evolved into monophyletic African, American and Asia-Pacific lineages. The endosymbiotic bacterium Wolbachia pipientis restricts positive-sense RNA 42 viruses in Ae. aegypti. Re-analysis of a small RNA library of Ae. aegypti cells co-infected with 43 44 AeAV and Wolbachia produces an abundant RNAi response consistent with persistent virus replication. We found Wolbachia enhances replication of AeAV when compared to a tetracycline 45 cleared cell line, and AeAV modestly reduces DENV replication *in vitro*. The results from our study 46 improve understanding of the diversity and evolution of the virome of Ae. aegypti and adds to 47 previous evidence that shows Wolbachia does not restrict a range of negative strand RNA viruses. 48

49 **Importance**

The mosquito *Aedes aegypti* transmits a number of arthropod-borne viruses (arboviruses) such as dengue virus and Zika virus. Mosquitoes also harbour insect-specific viruses that may affect replication of pathogenic arboviruses in their body. Currently, however, there are only a handful of insect-specific viruses described from *Ae. aegypti* in the literature. Here, we characterise a novel negative strand virus, Aedes anphevirus (AeAV). Meta-analysis of *Ae. aegypti* samples showed that it is present in *Ae. aegypti* mosquitoes worldwide and

is vertically transmitted. *Wolbachia* transinfected mosquitoes are currently being used in biocontrol
as they effectively block transmission of several positive sense RNA viruses in mosquitoes. Our

results demonstrate that *Wolbachia* enhances the replication of AeAV and modestly reduces dengue virus replication in a cell line model. This study expands our understanding of the virome in *Ae. aegypti* as well as providing insight into the complexity of the *Wolbachia* virus restriction phenotype.

62 Introduction

The yellow fever mosquito *Aedes aegypti* is a vector of medically important viruses with worldwide distribution within the tropical and subtropical zones (1). *Ae. aegypti* is the principal vector of both dengue virus (DENV) and Zika virus (ZIKV) (Family: *Flaviviridae*) with estimates suggesting up to 390 million incidences of DENV infections a year (2), and approximately 400,000 cases of ZIKV during the 2015–2016 Latin American ZIKV outbreak (3).

The ability of mosquitoes to transmit viruses is determined by a complex suite of genetic and 68 69 extrinsic host factors (4-6). One developing area is the contribution of insect-specific viruses (ISVs), demonstrated not to replicate in mammalian cells, in the vector competence of individual 70 71 mosquitoes (7, 8). ISVs can suppress the anti-viral RNAi response as shown in Culex-Y virus (CYV) of the Birnaviridae family (9), or enhance the transcription of host factors; cell fusing agent 72 virus (CFAV) (Family: Flaviviridae) infection of Ae. aegypti Aa20 cells upregulates the V-ATPase-73 associated factor RNASEK allowing more favourable replication of DENV (10). ISVs have also 74 75 been shown to suppress or exclude replication of arboviruses; prior infection of Aedes albopictus C6/36 cells and Ae. aegypti mosquitoes with Palm Creek virus (PCV) (Family: Flaviviridae) has 76 been shown to supress replication of the zoonotic West Nile virus (WNV) and Murray Valley 77 encephalitis virus (Family: Flaviviridae) (11, 12). Also, it has recently been demonstrated in Aedes 78 79 cell lines that dual infection with Phasi Charoen-like virus (Family: Bunyaviridae) and CFAV restricts the cells permissivity to both DENV and ZIKV infection (13). 80

Metagenomic and bio-surveillance strategies have proved invaluable in describing the virome diversity of wild-caught *Culicinae* mosquitoes (14, 15). To date, six ISVs have been identified and characterised from wild-caught and laboratory *Ae. aegypti*; CFAV (16, 17), Phasi Charoen-like virus (Family: *Bunyaviridae*) (18), Dezidougou virus from the Negevirus taxon (19), Aedes densoviruses (Family: *Parvoviridae*) (20) and the unclassified Humaita-Tubiacanga virus (HTV)

86 (21). Recently, transcriptomic analysis of wild-caught *Ae. aegypti* mosquitoes from Bangkok,
87 Thailand and Cairns, Australia suggested possible infection of the mosquitoes with up to 27 insect88 specific viruses, the majority of which currently uncharacterized (22). This represents a narrow
89 understanding of the diversity of the circulating virome harboured by *Ae. aegypti* mosquitoes.

In this study, we identified and characterised a novel negative-sense RNA Anphevirus, putatively 90 91 named Aedes anphevirus (AeAV), from the order Mononegavirales in Ae. aegypti mosquitoes. 92 According to the most recent International Committee on Taxonomy of Viruses (ICTV) report (23), Xīnchéng mosquito virus (XcMV), assembled as part of a metagenomic analysis of Anopheles 93 94 sinensis mosquitoes in Xinchéng China, is the only member of the Anphevirus genus and closely related to members of Bornaviridae and Nyamiviridae (24). Originally thought to only encode for 95 four ORFs, presence of a number of closely related viruses to XcMV from West African Anopheles 96 97 gambiae mosquitoes (15) and West Australian Culex mosquitoes (25) suggests that members of 98 this taxon encode for six ORFs with a genome size of approximately 12kb.

The endosymbiotic bacterium *Wolbachia pipientis* was first shown to restrict RNA viruses in *Drosophila melanogaster* (26, 27). Transinfection of *Wolbachia* into *Ae. aegypti* restricted DENV and Chikungunya virus (Family: *Togaviridae*) replication in the host (28). In *Ae. aegypti* Aag2 cells, stably transinfected with a proliferative strain of *Wolbachia* (*w*MelPop-CLA), the endosymbiont restricts CFAV (29), but has no effect on the negative sense Phasi Charoen-like virus (Family: *Bunyaviridae*) (30). In addition to characterising AeAV, we also studied the effect of *Wolbachia* on AeAV replication and co-infection of AeAV and DENV in *Ae. aegypti* cells.

106 Results

Identification and assembly of the full Aedes anphevirus (AeAV) genome from *Wolbachia* infected Aedes cells

During replication of RNA viruses in *Ae. aegypti* mosquitoes, the RNA interference (RNAi) pathway cleaves viral dsRNA intermediates into 21nt short interfering RNAs (vsiRNAs) (31, 32). Using this fraction of reads from RNA-Seq data, it is possible to *de novo* assemble virus genomes (21, 33).

The previously sequenced small RNA fraction of embryonic Ae. aegypti Aag2 cells and Aag2 cells 112 stably infected with Wolbachia (wMelPop-CLA strain) (34) was trimmed of adapters, filtered for 113 21nt reads and de novo assembled using CLC Genomics Workbench with a minimum contig 114 115 length of 100nt. The resulting contigs were then gueried using BLASTX against a local virus protein database downloaded from the National Centre for Biotechnology Information (NCBI). In 116 the Aag2.wMelPop-CLA assembly, four contigs between 396-1162nt were found to have amino 117 acid similarity (E value 9.46E-51) to proteins from two closely related Mononegavirales viruses: 118 Culex mononega-like virus 1 (CMLV-1) and Xīnchéng mosquito virus (XcMV); the type species for 119 120 the Anphevirus genus. No contigs from the Wolbachia negative Ae. aegypti Aag2 dataset showed any similarity to CMLV-1 or XnMV. 121

Subsequent RT-PCR analysis between RNA samples from Aag2 and Aag2. MelPop-CLA cell 122 lines indicated that this tentative virus was exclusive to the Aag2. MelPop-CLA cell line (Fig. 1A). 123 124 We hypothesised that the presence of any putative virus may have been the result of contamination during Wolbachia transinfection. The Wolbachia wMelPop-CLA strain was isolated 125 from the Ae. albopictus cell line RML-12 and transinfected into Aag2 (35) and the Ae. albopictus 126 C6/36 (C6/36. MMelPop-CLA) cell lines (36). RT-PCR analysis of RNA extracted from RML-12 and 127 128 C6/36 cells, as well as the Ae. aegypti cell line Aa20, showed that the putative virus was present only in RML-12 cells (Fig. 1A). 129

130 To recover the remainder of the virus genome, transcriptome RNA-Seg data from RML-12. MelPop-CLA and C6/36. MelPop-CLA cells were downloaded (37, 38) and de novo 131 132 assembled with automatic bubble and word sizes using CLC Genomics Workbench. BLASTN analysis of assembled contigs indicated that a complete 12,940 contig from the C6/36.wMelPop-133 CLA cells and two contigs (9624 and 3487nt) from RML-12. MelPop-CLA were almost identical 134 (99-100% pairwise nucleotide identity) to the virus like contigs assembled from Aag2. MelPop-135 CLA. We were then able to use this reference to recover the full genomes from Aag2. MelPop-136 137 CLA and RML-12. MelPop-CLA consensus mapping to this reference. To re-validate that AeAV 138 was only present in Aag2.wMelPop-CLA cells, reads from the Wolbachia-negative Aag2 cells were mapped to the representative genome, and only four reads were identified in the data. The result 139

from here and the RT-PCR analysis above (Fig. 1A) also confirm that the virus found in the *Wolbachia* transinfected cells originate from RML-12 cells in which *w*MelPop-CLA was originally transinfected and subsequently transferred to other cell lines.

143 Characterisation of Aedes anphevirus (AeAV)

AeAV genomes assembled in this study were between 12,455 to 13,011 nucleotides in length with a %GC content of 46.8% and encode for 7 non-overlapping ORFs (Fig. 1B). Phylogenetic analysis of the RNA-dependent RNA polymerase protein places AeAV within a well-supported clade of the unassigned *Anphevirus* genus, which are from the order *Mononegavirales* and closely related to members of *Bornaviridae* and *Nyamiviridae* (Fig. 1C).

149 All members of Mononegavirales have a negative-stranded RNA genome encapsidated within the capsid and the RNA polymerase complex (39). The RNA genome is used as the template by the 150 RNA polymerase complex to sequentially transcribe discrete mRNAs from subgenomic genes. 151 mRNA from each gene is capped and polyadenylated. To analyse the transcriptional activity of 152 153 AeAV, we used the poly-A enriched RNA-Seq libraries prepared from the Cali. Colombia laboratory 154 strain (40). Read mapping and coverage analysis of the AeAV genome showed that AeAV follows the trend of reduced transcriptional activity seen in other Mononegavirales species (41) with 155 approximately 50% reduction between ORF1 and ORF2 but an increased transcription between 156 ORF2 and ORF3 (Fig. 1B). The reduction in transcriptional activity of AeAV genes is conserved for 157 each sequential ORF with the least transcriptional activity for ORF7/L protein, that is conserved in 158 all AeAV strains in Poly(A) enriched RNA-Seg libraries (Fig. S1). 159

160 ORF1 of AeAV encodes a predicted 49kDa nucleoprotein with no transmembrane domains and closest pairwise amino acid identity (26%) to the nucleoprotein gene from Culex mononega-like 161 virus 1 (CMLV-1) from Culex mosquitoes in Western Australia (25). Protein homology analysis 162 using HHPred showed that ORF1 was a likely homolog of the p40 nucleoprotein of the Borna 163 disease virus (Probability 98.66%, E- value: 7.1e-10). ORF2 encodes an 11kDa protein with two 164 transmembrane domains in the N-terminus of the protein with no similarity to any proteins within 165 the non-redundant protein database or homologs as predicted by HHPred. ORF3 and ORF4 166 encode putative glycoproteins, 64kDa and 72kDa, respectively. ORF3 has no pairwise amino acid 167

similarity to any virus protein or homologs as per HHPred analysis. ORF4 was predicted to have a signal peptide in the N-terminus followed by a heavily O- and N-linked glycosylated outside region as well as two transmembrane domains in the C-terminus of the protein. ORF4 is most closely related to the glycoprotein from the Gambie virus identified from West African *An. gambiae* mosquitoes with 45% pairwise amino acid identity (15). Protein homology analysis predicted ORF4 to be a homolog of the Human Herpesvirus 1 Envelope Glycoprotein B (Probability 99.88%, Evalue 2.2e-22).

The presence of a Zinc-like finger (ZnF) domain in a small ORF proximal to the L protein previously reported in closely related viruses (15) (Fig. 2A and B), was identified in AeAV based on sequence alignment (Fig. 1B). Re-analysis of putative ORFs from CMLV-1 and CMLV-2 (25) showed the presence of this GATA-like ZnF domain in both of these viruses and the genus type species XcMV identified from *An. sinesis* (24) (Fig. 2C).

180 ORF6 encodes for a small 4kDa protein that has a single transmembrane domain in the Cterminus. This protein was almost missed in the prediction of ORFs due to having only 37 amino 181 acids, however, it has a strong transcriptional coverage in Poly-A datasets and exists in all 182 assembled strains (Fig. 1B and Fig. S1). It was predicted to share no structural homology or amino 183 acid identity with any previously reported peptide. In addition to this, we were able to identify small 184 transmembrane domain containing proteins proximal to or overlapping with the ZnF protein in 185 CMLV-1, CMLV-2 and XcMV (Fig. 2A), suggesting that this protein may be a conserved feature of 186 anpheviruses. 187

ORF7 encodes for the 226kDa L protein, has 41% pairwise amino acid identity with the RNA dependent RNA polymerase from CMLV-1. Protein domain analysis of the L protein showed the highly conserved *Mononegavirales* RNA dependent RNA polymerase, mRNA capping domain and a mRNA (guanine-7-) methyltransferase (G-7-MTase) domain conserved in all L proteins in *Mononegavirales* (42).

193 AeAV cis-regulatory elements

194 For identification of *cis*-regulatory elements in the AeAV genome, we used MEME (Multiple Em for Motif Elicitation) to search for overrepresented 5-50nt motifs (43). Using a 0-order Markov model, 195 196 one 32nt motif 3'-UUVCUHWUAAAAAACCCGCYAGUUASAAAUCA-5' was considered statistically 197 significant (E-value: 4.2e-010). Importantly the motif was proximal to each predicted virus gene ORF, suggesting it may be a potential promoter (Fig. 3A and C). No motif was found between ORF 198 5 and 6 in AeAV suggesting that these two genes may be under the control of a single cis-199 regulatory element. Interestingly, the complement of this motif appeared twice on the anti-genome 200 201 suggesting that it may be used in an anti-genome virus intermediate. We noticed that these motifs 202 localised to partial palindromic repeats and predicted that they may form stable secondary RNA structures. Using RNAfold, we were able to visualise and predict the MFE structure 20nt upstream 203 and downstream of the motif (44). All predicted *cis*-regulatory motifs formed partial or complete 204 stable secondary stem loops and hairpins with high base-pair probabilities (Fig. 3B). The exception 205 was predicted element 3, which is proximal to the second ORF; as this second gene is transcribed 206 less than ORF3 irrespective of its similarity to the motif, the lack of a stable stem loop structure 207 208 may be a novel transcriptional regulatory mechanism. The presence of two conserved 209 homopolymeric triplets in the overrepresented motif is very similar to "slippery" -1 ribosome frame shifting (RFS) sites XXX YYY Z (X=A, G, U; Y=A, U; Z=A, C, U) (45). It has been previously 210 demonstrated that similar 'slippery' sequence motifs followed by a predicted stem-loop structure is 211 212 a feature of rhabdovirus gene overlap regions (46). In AeAV, this feature appears in the intergenic 213 space and is unlikely to represent ribosomal frame shifting event and subsequent extension of a 214 protein. We also searched for additional slippery motifs in the AeAV genome. The genomic context 215 for each predicted "slippery motif" did not extend or produce additional ORFs.

AeAV infection is widespread in *Ae. aegypti* laboratory colonies, wild-caught mosquitoes and cell lines

Taking advantage of the currently published RNA-Seq data, we performed a meta-analysis of global incidence and genetic diversity of this virus. We were able to show that AeAV is ubiquitous in laboratory colonies, cell lines and wild-caught *Ae. aegypti* mosquitoes. During preparation of this manuscript a partial AeAV genome of 5313nt (Accession: MG012486.1) was deposited into NCBI nucleotide database from a study characterising the evolution of piRNA pathways across
arthropods (47). Using our AeAV genome reference, we were able to complete the CDS portion of
the genome and also 16 additional strains of AeAV with two additional incomplete genomes (Fig.
(Table S1).

226 AeAV was present in colonies of Ae. aegypti established from eggs collected in Bakoumba, Gabon (48) and also from Rabai, Kenya (designated K2, K14) as well as four mated hybrid strains (49). In 227 228 colonies wild caught from locations in the Americas (47), full genomes of AeAV were assembled from Miami, USA, Cali, Colombia (40), and Chetumal, Mexico (50, 51) laboratory strains. Partial 229 230 genomes of AeAV were assembled from Cayenne and St-Georges, French Guiana (52). AeAV was identified in colonies established from eggs collected in Chaiyaphum and Rayong, Thailand 231 (49, 53) as well as Jinjang, Malaysia (54). AeAV was also identified from the widely used Bora-232 Bora reference strain from French Polynesia (55). AeAV was also present in eight pools of wild-233 234 caught Ae. aegypti mosquitoes used for ZIKA bio-surveillance in Miami, Florida (56) as well as Nakhon Nayok, and Bangkok, Thailand (18, 22). 235

In *Aedes* cell lines, AeAV was assembled from RNA-Seq data from the larval *Ae. aegypti* line CCL-125 originally produced in Pune, India (57) and sequenced by the Arthropod Cell Line RNA-Seq initiative, Broad Institute (broadinstitute.org). With the exception of RNA-Seq data from the three *Aedes* cell lines stably infected with *Wolbachia* (RML-12.*w*MelPop-CLA, C6/36.*w*MelPop-CLA and Aag2.*w*MelPopCLA), AeAV was not identified in any other available C6/36 or Aag2 RNA-Seq libraries.

242 Genetic variation and evolution of AeAV strains

To assess relatedness and evolution between AeAV strains, a Maximum likelihood phylogeny (PhyML) was undertaken of the CDS region of all strains with complete genomes (Fig. 5A). The unrooted radial phylogenetic tree indicated three strongly supported monophyletic lineages associated with the geographic origin of the sample. We have designated these lineages of AeAV as African, American and Asia-Pacific (Fig. 5B).

248 In the American lineage of AeAV, all strains that are associated with Wolbachia-infected Aedes cell lines (RML-12. MMelPop-CLA, C6/36. MMelPop-CLA and Aag2. MMelPopCLA) are almost identical 249 (99.55-99.86% identity), supporting the hypothesis that contamination of C6/36 and Aag2 cell lines 250 251 infected with Wolbachia is likely from the original RML-12 cell line. AeAV from the eight wild-caught pools of Ae. aegypti mosquitoes from Florida, USA (56) and the laboratory colony established from 252 wild collected samples Florida (47) were almost identical (99.86% pairwise identity) with only 17nt 253 differences over the CDS region. The three African lineage strains of AeAV were slightly closer in 254 pairwise nucleotide identity to the American strains (92.65-93.15%) than the Asia/Pacific strains 255 256 (91.63%-91.74%). All samples that originated from Thailand form a monophyletic group and are closely related to other Thai strains (99.23-99.62%). 257

We hypothesised that AeAV may have been harboured as part of the virome of Ae. aegypti 258 mosquitoes as Ae. aegypti expanded from its sub-Saharan African location into the Americas and 259 260 Asia-Pacific (58). Phylogenetic studies of the Ae. aegypti genome support the origin of Ae. aegypti from Africa into the New World (Americas) and a subsequent secondary invasion of Ae. aegypti 261 aegypti from the New World to the Asia-Pacific region (59, 60). Comparing the evolution of the Ae. 262 aegypti nuclear genome with the evolution of AeAV indicates that the Asia-Pacific strains of AeAV 263 264 have not evolved from the currently circulating American strain lineage. This may indicate that the virus was established independently in both the New-world Americas and also in the Asia-Pacific 265 266 (Fig. 5B).

267 Anphevirus-like insertions into the Ae. aegypti genome

The Ae. aegypti genome has a large repertoire of virus genes and partial viral genomes, termed 268 269 Endogenous Viral Elements (EVEs) (61, 62). To explore the possibility of anphevirus-like insertions 270 within the Ae. aegypti genome, we queried the most recent Liverpool genome (Aaegl5) with the Aag2.wMelPop-CLA reference the VectorBase 271 AeAV strain using BLASTN suite (https://www.vectorbase.org/blast). There were numerous hits of nucleotide similarity (67-70%) of 272 273 500-1704nt regions dispersed throughout the Ae. aegypti genome. EVEs are acquired through recombination with long terminal repeat (LTR) retrotransposons (62). We present one ~20kb 274 portion of Chromosome 2 of the Ae. aegypti genome (Fig. 6) with four anphevirus-like insertions 275

276 and close proximity to LTR retrotransposable fragments in unidirectional orientation. This suggests 277 insertion of viral elements through LTR retrotransposases and a long evolutionary history of 278 challenge with anphevirus-like species in *Ae. aegypti*.

Aedes Anphevirus (AeAV) replicates in *Aedes* cell lines but does not replicate in three mammalian cell lines

Supernatant of Aag2. *w*MelPop-CLA cells was infectious to both *Ae. aegypti* cells (Aa20), and *Ae. albopictus* C6/36 cells over a five-day time course through RT-qPCR analysis (Fig. 7A). Generally, there was significantly more relative AeAV genome copies detected in C6/36 cells at 1 and 5 dpi compared to Aa20 cells. There were also significantly more anti-genome copies of AeAV in C6/36 cells as compared to Aa20 cells is not unexpected since C6/36 cells are RNAi deficient and generally RNA viruses replicate more efficiently in the cells (63-65).

We assumed that AeAV is an insect-specific virus based on its phylogenetic position, however, to 288 289 test if AeAV can replicate in mammalian cells, we inoculated human hepatocellular carcinoma cells (Huh-7), African green monkey cells (Vero), and baby Hamster Kidney (BSR) cells with medium 290 from AeAV-infected cells and performed RT-PCR for AeAV RNA genome abundance over a 7-day 291 time course. While AeAV RNA (most likely from the inoculum) could be detected by RT-PCR at 292 293 day 1 and 3 after inoculation, it did not increase over time and was visibly reduced in the mammalian cells by 5/7 dpi (Fig. 7B). AeAV was also not detected in the An. gambiae cell line 294 MOS-55 transinfected with wMelPop-CLA from RML-12-wMelPop-CLA (36) sequenced by the 295 Arthropod Cell Line RNA-Seq initiative, Broad Institute (broadinstitute.org). Taken together, the 296 results suggest that AeAV infection may be restricted within the subfamily Culicinae or even the 297 Aedes genus and is insect specific. 298

299 Wolbachia pipientis infection in Ae. aegypti cells enhances AeAV replication

As *Wolbachia* is being deployed in the field to reduce dengue transmission, we were interested to find out if it has any effect on replication of AeAV. We extracted RNA from Aag2.*w*MelPop-CLA cells and a previously tetracycline cured Aag2.*w*MelPop-CLA cell line (66), and tested the effect of *Wolbachia* infection on AeAV genome and anti-genome copies. AeAV genomic RNA copies were significantly greater in *Wolbachia*-infected (Aag2.*w*MelPop-CLA) cells than those in tetracycline cleared (Aag2.*w*MelPop-CLA.Tet) *Ae. aegypti* cells, however, there was no statistically significant difference between the relative anti-genome copies between the two cell lines (Fig. 8A).

To explore the host small RNA response to AeAV, clean reads from previously prepared sRNA 307 libraries from Aag2. MMelPop-CLA and Aag2 (34) were mapped to the AeAV genome. In the 308 309 cytoplasmic fraction of the Aag2. MelPop-CLA sample 870,012 of 4,686,954 reads (18.56%) mapped to AeAV. In the nuclear fraction 420,215 of 11,406,324 reads (3.68%) mapped to the 310 311 genome. In the combined Aag2 sRNA library, of 8,600,821 clean reads only four reads mapped to the AeAV genome. The mapping profile of 18-31nt reads mapped from the Aag2.wMelPop-CLA 312 library to AeAV indicates a higher proportion 27-31nt viral derived PIWI RNAs (vpiRNAs) than the 313 21nt vsiRNAs (Fig. 8B). 314

Analysis of the profile of mapped AeAV vsiRNAs fairly ubiquitously targeted the AeAV genome and anti-genome (Fig. 8C). Hotspots in the vpiRNA mapping profile appeared to target the 3'UTR and ORF1 and the 5'UTR of the AeAV anti-genome (Fig. 8D). Biogenesis of vpiRNAs are independent of Dicer-2 with a bias for adenosine at position 10 in the sense position and a uracil in the first nucleotide of antisense polarity (67). This "ping-pong" characteristic signature was apparent in the vpiRNA reads from the cell line (Fig. 8E and F).

Persistent infection of AeAV in Aa20 cells modestly reduces replication of DENV-2 genomic RNA

Recently, it has been demonstrated that in Aedes cell lines experimentally infected with two ISVs 323 replication of DENV and ZIKV was reduced (13). To test if there was any interaction between 324 AeAV and the subsequent challenge of cells with DENV, we generated an Aa20 cell line inoculated 325 with medium from RML-12 and maintained it for three passages. Aa20 cells persistently infected 326 with AeAV were challenged with 0.1 and 1 multiplicities of infection (MOI) of DENV-2 ET-100 327 328 strain. RT-qPCR analysis of DENV-2 genomic RNA showed that accumulation was on average less in AeAV infected Aa20 cells as compared to the control (Fig. 9A and B). This reduction in 329 DENV-2 genome copies was statistically significant at MOI of 0.1 at both three and five days post 330

infection. No AeAV-related RT-qPCR product was detected in mock-infected Aa20 cells (data not
shown). We also examined the effect of DENV infection on AeAV RNA levels in the AeAV
persistently infected cells. RT-qPCR analysis showed no significant effect on AeAV levels between
1 and 3 days post-DENV infection, however, AeAV genomic RNA levels significantly declined at 5
days-post DENV infection (Fig. 9C). There was no significant difference in the results between 0.1
and 1 MOI of DENV.

337 Evidence for vertical transmission of AeAV

338 We were fortunate to explore the potential vertical transmission of AeAV by using RNA-Seq data of 339 uninfected and infected mated individuals from a study characterising the genetic basis of olfactory preference in Ae. aegypti (49). Briefly, McBride and collegues used eggs from a number of Ae. 340 aegypti species in Rabai, Kenya to establish laboratory colonies for RNA-Seq analysis. We 341 identified AeAV in the domestic K2 and K14 colonies, which was seemingly absent from the other 342 343 Rabai (K18, K19, K27) colonies. The K27 colony was interbred with the strain K14, which we found 344 to be AeAV positive. In all the four resultant hybrid colonies, which were subjected to RNA-Seq analysis, we were able to *de novo* assemble identical K14 AeAV strain genomes (Fig. 10). 345

The possibility of vertical transmission also is supported by the presence of AeAV in both RNA-Seq data from the sperm of adult male mosquitoes (54) and the female reproductive tract (53).

348 Discussion

349 The ability of AeAV to propagate in Ae. aegypti and Ae. albopictus cell lines but not in the three mammalian cell lines suggests that AeAV is most likely an ISV, although this needs to be further 350 confirmed using cell lines from other species. To the best of our knowledge, this is the first 351 comprehensive characterisation of any Anphevirus species within Mononegavirales and the first 352 Mononegavirales virus species within Ae. aegypti. Although as this manuscript was under revision, 353 the complete genome sequence of AeAV and its phylogenetic relationship with other ISVs was 354 published in a short communication (68). While we have demonstrated that AeAV is spread 355 worldwide in Ae. aegypti mosquitoes, we have limited understanding as to how prevalent AeAV is 356 in individual mosquitoes in wild populations, its tissue tropism, or potential impacts on the host. 357

Although it is likely that AeAV is maintained in wild populations of *Ae. aegypti* mosquitoes through vertical transmission, it is possible that AeAV could be maintained though venereal transmission as we were able to identify the whole AeAV genome from a dataset prepared from the sperm of adult male *Ae. aegypti* mosquitoes (54).

362 To our knowledge, the oldest continually maintained colony of laboratory mosquitoes with AeAV present comes from Jinjang, Malaysia, which was established from wild-collected samples from as 363 364 early as 1975 (54). This suggests vertical transmission rates of AeAV are high or a high incidence of AeAV within the colony. In comparison, in both French Guiana colonies, we were unable to 365 366 recover the complete genomes from these strains. It is unlikely that this is due to insertion of AeAV into the nuclear Ae. aegypti genome as numerous reads mapped to the ORF7/L region of the 367 reference strain; however, there was not enough coverage to reach consensus of the full genome. 368 As these libraries were prepared from homogenates of mosquitoes, it seems likely that the 369 370 incidence of infection within these colonies may be lower; however, further analysis would have to be conducted. 371

In our analysis, AeAV was not detected in any of the widely used Liverpool (LVP) and Rockefeller/UGAL, as well as derived "white eye mutant" strains of *Ae. aegypti.* Analysis of published ncRNA-Seq data from Australian Townsville and Cairns colonies of wild-caught Australian mosquitoes (69, 70) suggests that there is no RNAi response or presence of AeAV in these mosquitoes. Evidence from this study and others suggests that widely used laboratory strains of *Ae. aegypti* harbour a diverse and heterogeneous virome composition and may contribute to the variable vector competence between these colonies.

In our analysis, the geographic origin of the RNA-Seq samples matched the resulting phylogenetic relationship of each strain. The presence of AeAV in the *Ae. albopictus* cell line RML-12, presumably the origin of the AeAV contamination in other *Aedes* cell lines transinfected with adapted *w*MelPop strain, was the only *Ae. albopictus* sample in our analysis. During our analysis, we queried all of the 266 currently available *Ae. albopictus* RNA-Seq datasets (TaxonID: 7160) uploaded to the SRA, none of which indicated presence of AeAV. We hypothesise that AeAV from RML-12 is likely due to contamination as the cell line is often mischaracterised as originating from

Ae. aegypti (71, 72). In many laboratories that study arbovirus interactions, more than one Aedes 386 cell line is maintained. As the RML-12 AeAV strain is genetically placed within the American 387 lineage, and the namesake of the cell line, Rocky Mountain Laboratories, (71) located in Montana, 388 389 USA suggests possible contamination from domestic Ae. aegypti mosquito samples. While no other Ae. albopictus RNA-Seq data were positive for AeAV in this study, we cannot rule out the 390 possibility that AeAV could exist in American populations of Ae. albopictus. Ae. aegypti and Ae. 391 albopictus co-exist in North America and compete for larval habitats of discarded tires and other 392 393 artificial containers (73).

394 RNAi response is commonly observed in mosquitoes against RNA viruses. This response includes miRNA, siRNA and piRNA pathways (74, 75). Similarly, we found a large number of 21nt vsiRNAs 395 produced against AeAV in infected cells that were evenly mapped to both sense and antisense 396 strands, indicating that dsRNA intermediates produced during replication must be the target of the 397 398 host cell RNAi response. In addition, a large number of vpiRNAs were found mapped to the 5'UTR, ORF1 and the 3'UTR of the AeAV genome. These vpiRNAs had the typical ping-pong signature 399 (U₁-A₁₀) of secondary piRNAs. This signature has also been found in vpiRNAs produced during 400 alphavirus (76) and bunyavirus (63, 77) infections, but not in vpiRNA-like small RNAs in most 401 flaviviruses, such as DENV (78), ZIKV (79, 80) and an insect-specific flavivirus (81). We found a 402 403 higher proportion of small RNAs from Aag2-wMelPop-CLA cells that mapped to AeAV are vpiRNAs (about 50%) as compared to less than 10% vsiRNAs. Literature suggests that when the siRNA 404 pathway is compromised more vpiRNAs are produced. This has been shown in RNAi deficient 405 406 C6/36 cells when infected with Sindbis virus, Rift Valley fever virus or La Crosse virus (63-65). The RNA-Seq data from C6/36. MMelPop-CLA cells were for long transcripts rather than small RNAs, 407 408 therefore we were not able to confirm if in those cells there are higher proportion of vpiRNAs than vsiRNAs. The over-representation of vpiRNAs in respect to vsiRNAs has also been demonstrated 409 410 in negative-sense Bunyavirales members PCLV and RVFV (30, 65) in Aag2 cells for all segments of the genome. It remains to be seen however if the higher vpiRNA to vsiRNA ratio in 411 Aag2. *w*MelPop-CLA could be due to suppression of the siRNA pathway by AeAV, or alternatively, 412 Wolbachia may have an effect on the siRNA pathway. It seems however more likely that as 413

414 negative sense RNA viruses produce less dsRNA replicative intermediates, these could be simply 415 less targeted by the siRNA pathway and are unable to be resolved by sRNA profiling. These 416 possibilities require further investigations, and it remains to determine the role of the vpiRNAs in 417 AeAV replication or host anti-viral response.

418 The effects of Wolbachia on virus restriction are variable and depend on Wolbachia strain, virus family and transinfected host (82). Wolbachia was shown to enhance AeAV replication in Ae. 419 420 aegypti cells in this study. Recent studies have demonstrated that Wolbachia has no effect on multi-segmented negative-sense RNA viruses; for example, Phasi Charoen-like virus (PCLV) 421 422 (Family: Bunyaviridae) in wMelPop-CLA strain infected Aag2 cells (30), La Crosse virus (Family: Bunyaviridae) and vesicular stomatitis virus (Family: Rhabdoviridae), which is in the same order as 423 AeAV, in wStri strain in Ae. albopictus C710 cell line (83), and the Rift Valley fever virus (RVFV) 424 (Family: Phenuiviridae), in Culex tarsalis mosquitoes transinfected with a somatic Wolbachia 425 426 (strain wAlbB) had no effect on RVFV infection or dissemination (84), respectively. RVFV and PCLV belong to the Bunyavirales order and AeAV belongs to Mononegavirales, distinct orders of 427 negative strand RNA viruses. All three viruses have conserved features that may provide insight 428 into how they might be protected from restriction by Wolbachia. The genome of all negative-sense 429 430 ssRNA viruses is both encapsidated within the nucleoprotein (85, 86) and is attached to RNA dependent RNA polymerase within the virion (87). The RNA dependent polymerase complex 431 carries out both transcription of virus genes and replication of the genome. 432

Wolbachia has been shown to restrict a number of positive-sense RNA virus species from the 433 434 Togaviridae and Flaviviridae families (82). After fusion and entry into the host cell, the genomes of Togaviridae and Flaviviridae species are released into the cytoplasm and translated directly into 435 polypeptide protein(s). These polypeptide proteins are processed by viral and cellular proteases to 436 generate the mature structural and non-structural proteins which are then used to replicate the 437 genome (88). While the exact mechanism for RNA virus restriction in Wolbachia-infected insects 438 439 has remained elusive, it has been shown that restriction of RNA viruses by Wolbachia happens early in infection (89, 90). In the Ae. albopictus cell line C710 stably infected with wStri, the 440 polypeptide of ZIKV is not produced as determined by immunoblot at one day post-infection (90). 441

Additionally, *Wolbachia* exploits host innate immunity to establish a symbiotic relationship with *Ae. aegypti* (91). Perhaps the combination of protection of the RNA nucleocapsid genome or genome segments when released in the cytoplasm or activity of the RNA-dependent RNA polymerase may aid in evasion of host immune response enhanced by *Wolbachia* or *Wolbachia* effector molecules (92). However, a recent study suggested increases in infection of *Ae. aegypti* mosquitoes by insect-specific flaviviruses when they harbour *Wolbachia w*Mel strain (93).

The ability of AeAV to modestly reduce DENV-2 genomic RNA in a persistently infected cell line was unexpected. While it has previously been shown that members of the same virus family can provide super exclusion of additional viruses (12, 94), little work has been undertaken to look at cross viral family exclusion effects. Our results showed that less DENV replication occured in the presence of AeAV, with the difference particularly significant at lower MOI. If this suppressive effect also occurs in mosquitoes, enhancement of AeAV in *Ae. aegypti* mosquitoes infected with *Wolbachia* may be beneficial in terms of DENV suppression.

As *Ae. aegypti* is perhaps the most important vector of arboviruses worldwide, further work should be undertaken in understanding and characterising the virome of this mosquito and effects on mosquito life-history traits. Our findings provide new insights into the evolution and genetic diversity of AeAV across a wide geographic range as well as providing valuable insights into the virus features and families restricted by *Wolbachia* in mosquito hosts and its effects on arboviruses they transmit.

461 Materials and Methods

462 Cell lines maintenance and experimental infection with AeAV

Ae. aegypti cell line (Aag2) stably infected with *Wolbachia* (denoted Aag2.*w*MelPop-CLA) as previously described for the C6/36.*w*MelPop-CLA (35), with its previously generated tetracyclinetreated line (66), and both *Ae. albopictus* C6/36 cell line (57) and RML-12 cell lines were maintained in 1:1 Mitsuhashi-Maramorosch and Schneider's insect medium (Invitrogen) supplemented with 10% Fetal Bovine Serum (FBS, Bovogen Biologicals). Aa20 cells established from *Ae. aegypti* larvae (95) were maintained in Leibovitz's L15 medium supplemented with 10% FBS (France – Biowest) and 10% Tryptose phosphate broth at 27°C. African green monkey cells (Vero) were maintained in Opti-MEM I Reduced-Serum Medium supplemented with 2% FBS and 10 mL/L Penicillin-Streptomycin (Sigma-Aldrich). Human hepatocellular carcinoma cells (Huh-7) were maintained in Opti-MEM I Reduced-Serum Medium supplemented with 5% FBS and 10 mL/L Penicillin-Streptomycin (Sigma-Aldrich). BSR cells (a clone of Baby Hamster Kidney-21) were maintained in Dulbecco's Modified Eagle Medium (Gibco), 2% FBS and 10 mL/L Penicillin-Streptomycin (Sigma-Aldrich). All mammalian cells were kept at 37°C with 5%CO₂.

For experimental infection of cells, 10⁶ cells of *Aedes* or mammalian cells were seeded in a 12-well plate. Subsequently, supernatant from Aag2.*w*MelPop-CLA cells was collected, centrifuged at 2150×g for 5 min to remove cells and debris, and used as an AeAV inoculation source. One Aa20 cell line was experimentally inoculated with RML-12 cell supernatant and kept as a persistently infected AeAV cell line. Cells were collected at 1, 3 and 5 days post-inoculation for *Aedes* cell lines for RT-qPCR analysis and 1, 3, 5 and 7 days for mammalian cell lines for RT-PCR analysis.

482 Aedes anphevirus (AeAV) and dengue virus (DENV-2) interaction assay

The third passage of Aa20 cells persistently infected with AeAV (denoted Aa20_{AeAV}) and Aa20 mock were seeded at the density of 3×10⁵ cells in 12-well plates overnight. Cells were then infected with the East Timor (ET-100) DENV-2 strain at a multiplicity of infection (MOI) of 0.1 and 1, cells were rocked for an hour at room temperature and supernatant was discarded and replaced with fresh medium. Cells were collected at 0, 1, 3 and 5dpi after infection with mock collected at 5 dpi. Cells were subjected to RNA extraction to quantify the DENV-2 genomic RNA levels by RTqPCR as described below.

490 Assembly and identification of AeAV strains from RNA-Seq data

For detection of AeAV in previously published RNA-Seq data, we used the assembled RML-12 491 AeAV genome as a BLASTN query for all available Ae. aegypti (taxonID: 7159) RNA-Seq data 492 493 within the Sequence Read Archive (SRA) on NCBI. SRA run files with positive hits of 90-100% identity and an E value <2E-30 were downloaded and converted to fastq using the NCBI SRA 494 495 toolkit https://www.ncbi.nlm.nih.gov/sra/docs/toolkitsoft/ for further analysis. FastQC (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/) was used for quality checking of fastq 496

files and adapter identification. Fastq files were then imported into CLC Genomics Workbench
(10.1.1) and were adapter and quality trimmed (<0.02; equivalent Phred quality score of 17,
ambiguous nucleotides: 2).

Two strategies were used to assemble strains of AeAV; fastq files from the same source of *Ae. aegypti* were pooled and *de novo* assembled using the CLC Genomics Workbench assembly program with automatic bubble and word sizes. This was sufficient to assemble the full coding sequences (CDS) of most strains of AeAV. Table S1 contains *de novo* assembly statistics from each dataset used.

505 If *de novo* assembly did not produce the complete AeAV genome, to complete further sections of the AeAV genome clean reads were mapped to the C6/36. MelPop-CLA strain of AeAV with 506 stringent alignment criteria (Match score:1 Mismatch cost:2 length fraction 0.89 and similarity 507 fraction 0.89) to exclude false-positive mapping that derives from Endogenous Viral Elements 508 509 (EVEs). To confirm accuracy of assembly, the largest contigs of consensus mapping were 510 extracted and then used as a reference for re-mapping and manually checked. Final sequences of the virus genomes were obtained through the majority consensus of the mapping assembly and 511 were given Coding Complete (CC) or Standard Draft (SD) genome guality ratings (96). 512

513 RNA isolation, strand specific cDNA synthesis and RT-qPCR

514 Total RNA was extracted from mosquito cells using QIAzol Lysis Reagent (QIAGEN) and treated with Turbo DNase (Thermo Fisher Scientific) as per manufacturer's instructions. RNA quality and 515 quantity were evaluated using a BioTek Epoch Microspot Plate Spectrophotometer. For the 516 production of AeAV genome and anti-genome cDNA, two cDNA reactions were generated using 517 600ng of RNA and SuperScript III Reverse Transcriptase (Thermo Fisher Scientific). The genome 518 cDNA strand was synthesised using a forward primer to AeAV (AeAVGenome-RT 5'-519 AGACTTCTAAGCCTGCCCACA -3'), and the AeAV anti-genome cDNA strand was synthesised 520 using a reverse orientation primer (AeAVAntiGenome-RT 5'- ACACTTGCCATGTGCTCAG -3'). 521 522 Aedes Ribosomal protein subunit 17 (RPS17) primers (Ae. aegypti: AeRPS17-gR 5'-GGACACTTCGGGCACGTAGT-3' AalRPS17q-R 5'-523 and Ae. albopictus ACGTAGTTGTCTCTCTGCGCTC -3') were used for reference gene cDNA synthesis. Following 524

525 RT, qPCR with AeAV primers (AeAV-qF 5'-GACAATCGCATTGGCTGCAT-3' and AeAV-qR 5'-526 CCCGAGACAATCGGCTTCTT - 3') as well as primer pairs for the RPS17 genes (*Ae. aegypti:* 527 AeRPS17-qF 5'- CACTCCGAGGTCCGTGGTAT -3' and *Ae. albopictus:* AalRPS17-qF 5'-528 CGCTGGTTTCGTGACACATC -3') were undertaken. RPS17 was used for normalizing data as 529 described previously in *Ae. aegypti* cells (97).

530 For quantitation of DENV-2 genome copies in Aa20 cells, two SuperScript III reverse transcription 531 (Thermo Fisher Scientific) reactions with 1000ng of RNA were prepared. For DENV-2 genome 532 copies reaction, the reverse primer (DENV2-qR 5'- CAAGGCTAACGCATCAGTCA -3') and in a 533 separate cDNA synthesis reaction the *Ae. aegypti* RPS17 primers as described above. 534 Subsequently, qPCR for DENV-2 was carried out using the DENV-2 primer pair (DENV2-qF 5'-535 GGTATGGTGGGCGCTACTA -3' and DENV2-qR), and RPS17 was used as a normalising control 536 as also described above.

Each time-point for experimental infection was run with three biological replicates and two technical replicates. Platinum SYBR Green Mix (Invitrogen) was used for qPCR with 20ng of RT products in a Rotor-Gene thermal cycler (QIAGEN) as described above. The relative abundance of AeAV RNA and DENV-2 to the host reference gene was determined by qGENE software using the $\Delta\Delta C_t$ method and analyzed using GraphPad Prism.

To test for the replication of AeAV in mammalian cells, 1000ng of total RNA extracted from the cells was extracted and used for first stand synthesis with SuperScript III reverse transcriptase with the AeAVGenome-RT primer. qPCR was subsequently carried out using AeAVGenome-RT primer and the qPCR primer AeAVqR2 5'-ATGAAAGTATGGATACACACCGCG-3'. Products were then visualised on a 1% agarose gel.

547 Virus genome annotation

Potential ORFs of AeAV were analysed using NCBI Open Reading Frame Finder 548 (https://www.ncbi.nlm.nih.gov/orffinder/) with a minimal ORF length of 150. ORFs were cross 549 550 referenced with mapping from Poly(A) enriched transcriptomes (Fig. S1) to reduce false positive identification of ORFs. For determination of putative domains in AeAV, ORFs were translated and 551 Conserved Domain Search (CD 552 searched against the Service Search)

(https://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi). For protein homology detection, we used
the HHPred webserver on translated AeAV ORFs (98).

To best discriminate N-terminus transmembrane domains from signal peptides, we used the consensus TOPCONS webserver (99). Glycosylation sites were predicted by the NetNGlyc 1.0/ NetOGlyc 4.0 Server (http://www.cbs.dtu.dk/services/).

558 Phylogenetic analysis

559 For placement of AeAV within the order *Mononegavirales*, ClustalW was used on CLC Genomics 560 Workbench to align amino acid sequences of 30 L proteins of the most closely related 561 *Mononegavirales* species as determined by BLASTp of the NCBI non-redundant database. A 562 Maximum likelihood phylogeny (PhyML) was constructed using the Whelan and Goldman (WAG) 563 amino acid substitution model with 1000 bootstraps. Accession numbers and associated hosts 564 used to produce the phylogenetic tree are available in Table S1.

To determine relatedness between different strains of AeAV, genomes that were coding complete and greater than 30x coverage were trimmed of 3'UTR and 5'UTR regions and aligned using the ClustalW algorithm on CLC Genomics Workbench. A Maximum likelihood phylogeny (PhyML) was constructed. A Hierarchical likelihood ratio test (hLRT) with a confidence level of 0.01 suggested that the General Time Reversible (GTR) +G (Rate variation 4 categories) and +T (topology variation) nucleotide substitution model was the most appropriate. 1000 bootstrap replicates were performed with 95% bootstrap branching support cut-off.

572 Statistical analysis

573 Statistical analysis Unpaired t-test was used to compare differences between two individual 574 groups, while One-way ANOVA with Tukey's post-hoc test was carried out to compare differences 575 between more than two groups. Data that did not pass the normality test were re-analysed by the 576 non-parametric Wilcoxon test indicated in their relevant figure legends.

577 Accession numbers

578 All complete virus genome sequences generated in this study have been deposited in Genbank 579 under the accession numbers (MH037149-MH037164). The incomplete Cayenne and St Georges 580 French Guinana AeAV contigs have been placed in supplementary File S1 alongside

581 Aag2. *w*MelPop-CLA and C6/36. *w*Mel-CLA strains.

582 Acknowledgements

- 583 This project was funded by the Australian Research Council grant (ARC, DP150101782) to SA and
- 584 University of Queensland scholarship to RP. For the supply of the DENV-2 (ET-100) strain, we
- thank Dr Daniel Watterson and Professor Paul Young from the University of Queensland. We
- 586 thank Dr Daniel Watterson and Dr Jody Hobson-Peters for providing Huh-7 and BSR cells,
- respectively. The authors would like to thank the technical assistance of Dr Sultan Asad, Dr
- 588 Kayvan Etebari and Hugo Perdomo Contreras, as well as current and former members of the
- 589 Asgari lab for their fruitful conversations.

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 912 server for consensus prediction of membrane protein topology and signal peptides.
 913 Nucleic Acids Res 43:W401-407.
- 914 **Figure legends:**
- Figure 1. Presence of Aedes anphevirus (AeAV) in insect cell lines, and genome
- organisation and phylogeny of the virus. A) RT-PCR analysis of Aedes cell lines Aag2,
- 917 Aag2. *w*MelPop-CLA (Pop), *w*MelPop-CLA.Tet (Pop-T), Aa20, RML-12, and C6/36 for the
- presence of AeAV. RPS17 was used as a loading control. **B)** Genome organisation of the
- 919 Cali, Colombia AeAV genome strain and subgenomic gene transcription profile.
- 920 Transmembrane domains (TMD) are depicted as boxes with dashed lines and signal

peptide is depicted as a blue box. NP, nucleoprotein; G, glycoprotein; ZnF, zinc-like finger; 921 RdRP, RNA dependent RNA polymerase. C) AeAV is a member of the Anphevirus genus 922 (red), related to members of the Nyamiviridae (pink) and Bornaviridae (purple) in an 923 924 unassigned family within the order *Mononegavirales*. A multiple sequence alignment of the RNA dependent RNA polymerase and the mRNA capping domain was used to create a 925 Maximum likelihood phylogeny. The phylogeny is arbitrarily rooted. 1000 bootstraps were 926 performed and branches with bootstrap values greater than 85% are highlighted. Branch 927 length represent expected numbers of substitutions per amino acid site. Genbank protein 928 accession numbers are Bolahun virus variant 1 (AOR51366.1), Culex mononega-like virus 929 1 (CMLV1) (ASA47369.1), Culex mononega-like virus 2 (CMLV2) (ASA47322.1), Gambie 930 virus (AOR51379.1), Xincheng Mosquito Virus (XcMV) (YP 009302387.1), Borna disease 931 virus (YP 009269418.1), Canary bornavirus 1 (YP 009268910.1), Loveridges garter 932 snake virus 1 (YP_009055063.1), Parrot bornavirus 1 (AEG78314.1), Variegated squirrel 933 bornavirus 1 (SBT82903.1), Midway nyavirus (YP 002905331.1), Nyamanini nyavirus 934 (YP 002905337.1), Sierra Nevada virus (YP 009044201.1), Soybean cyst nematode 935 socyvirus (YP 009052467.1), Farmington virus (YP 009091823.1), Beihai rhabdo-like 936 virus 3 (APG78650.1), Beihai rhabdo-like virus 5 (YP_009333422.1), Beihai rhabdo-like 937 virus 6 (YP 009333413.1), Drosophila unispina virus 1 (AMK09260.1), Hubei diptera virus 938 11 (YP_009337182.1), Hubei orthoptera virus 5 (YP_009336728.1), Hubei rhabdo-like 939 virus 7 (YP 009337121.1), Orinoco virus (ANQ45640.1), Sanxia Water Strider Virus 4 940 (YP_009288955.1), Shuangao Fly Virus 2 (AJG39135.1), Wenling crustacean virus 12 941 (YP 009336618.1), Wenzhou Crab Virus 1 (YP 009304558.1), Wenzhou tapeworm virus 942 943 1 (YP_009342311.1), Wuchan romanomermis nematode virus 2 (YP_009342285.1).

Figure 2: Conservation of GATA-like Zinc finger (ZnF) domain and small transmembrane domain containing protein between tentative members of the *Anphevirus* taxon. A) Genome orientation of previously discovered viruses within the *Anphevirus* taxon, and B) two viruses within a closely related clade. Predicted ORF encoding the ZnF domain is indicated by a black square. Predicted ORFs containing transmembrane domains are indicated by dashed lines. Genbank accession numbers are shown below virus name. NP, nucleoprotein; G, glycoprotein; ZnF, zinc-like finger; RdRP, RNA dependent RNA polymerase. C) Alignment of predicted GATAlike ZnF protein sequence (C-X(2)-C-X(17-20)-C-X(2)-C) between three representative strains of AeAV (Miami, USA ; Pune, India; Rabai, Kenya) and predicted ZnF domain proteins from Figure 2A and B.

Figure 3. Aedes anphevirus (AeAV) cis-regulatory elements. A) Location and orientation of 954 955 predicted *cis*-regulatory element in AeAV indicated by numbered red arrows: downwards indicating 956 genome and upwards arrow indicating anti-genome. B) Predicted minimum free energy (MFE) 957 RNA structure of the region surrounding the motif for each element using the RNAfold web server. Colour indicates probability of base-pairing and motif is indicated by the black line. C) Sequence of 958 959 the conserved motif as predicted by MEME as well as location and the statistical confidence of the motif. Sequences are written 3' to 5' and anti-genome motif sequences 1 and 7 are depicted as 960 961 reverse complement for visual clarity.

Figure 4. Aedes anphevirus (AeAV) has worldwide distribution in *Ae. aegypti* laboratory
colonies, cell lines and wild-caught mosquitoes. Locations of mosquito collection from RNASeq data that were positive for AeAV (Table S1). Points refer to collection sites from American
(orange), Asia-Pacific (blue) and African (green) locations.

966 Figure 5. Aedes anphevirus (AeAV) strains have evolved into African, Asia-Pacific and American lineages. A) Maximum likelihood phylogeny (PhyML) between AeAV strains using a 967 General Time Reversible (GTR) + G +T model with 1000 bootstraps. Branch lengths represent 968 expected numbers of substitutions per nucleotide site. For visual clarity, the RML-12 clade and 969 970 Miami clades were collapsed and single examples were shown. B) Evolutionary history of worldwide sampling of Ae. aegypti adapted from (59, 60) from 1504 SNPs species. Bootstrapped 971 neighbour-joining network based on population pairwise chord-distances from with node support 972 973 over 90% is shown on relevant branches. New World (American) populations in yellow, and Asia-974 Pacific populations are shown in light blue. We have truncated the tree and rooted to the Ae. 975 aegypti formosus (Aef) shown as a red branch.

Figure 6. Genomic context for anphevirus-like insertions into the *Ae. aegypti* genome. A 21,
242nt portion of chromosome 2 depicting anphevirus insertions (red) with predicted ORFs that
encode for LTR retrotransposase elements (yellow).

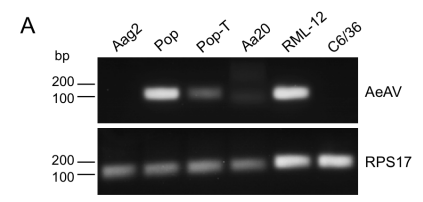
Figure 7. Aedes anphevirus (AeAV) is infectious to Aedes cell lines but does not replicate in
Huh-7, Vero and BSR vertebrate cell lines. (A) RT-qPCR of AeAV genome and anti-genome in a
five-day time course in *Ae. aegypti* Aa20 cells, and *Ae. albopictus* C6/36 cells. Error bars represent
the SEM of three biological replicates. (B) RT-PCR of AeAV genome in a seven-day time course in
Human hepatocellular carcinoma cells (Huh-7), African green monkey cells (Vero), Baby Hamster
Kidney (BSR). M, Mock infected cells.

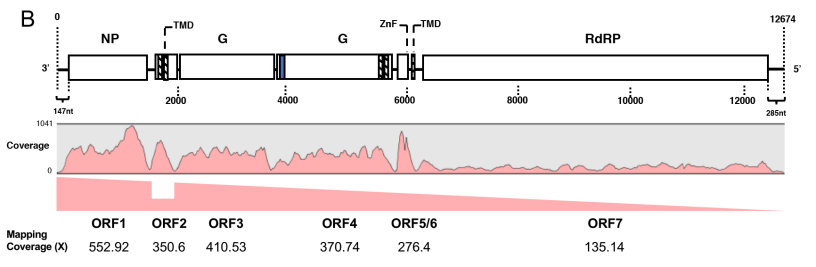
Figure 8. Aedes anphevirus (AeAV) genome replication is enhanced by Wolbachia infection 985 in Ae. aegypti cells and produces abundant vsiRNAs and vpiRNAs. A) RT-qPCR of the AeAV 986 genomic (gRNA) and antigenomic RNA in tetracycline cured Aag2.wMelPop-CLA cells (Pop-tet) 987 988 and Aag2.wMelPop-CLA cells (Pop) relative to RPS17. Error bars represent the SEM of six (genome) and three (antigenome) biological replicates. n.s, not significant; **, p < 0.01. B) Mapping 989 990 profile of pooled small RNA fraction in Aag2. MelPop-CLA cells. C) Alignment of the 21-nt sRNA reads (representing siRNAs), and D) the 26-31nt reads (representing piRNAs) mapped to the 991 AeAV antigenome (blue) and genome (red) in Aag2. MelPop-CLA cells. Relative nucleotide 992 frequency and conservation of the 28nt small RNA reads that mapped to the E) genome, and the 993 F) anti-genome of AeAV in Aag2. *w*MelPop-CLA cells. 994

Figure 9. Aedes anphevirus (AeAV) reduces dengue virus replication in Aa20 cells. Aa20 cells persistently infected with AeAV were infected with (A) 0.1 and (B) 1 MOI of dengue virus serotype 2 (DENV-2). Total RNA was extracted at 0, 1, 3 and 5 days following DENV-2 inoculation and analysed by RT-qPCR. (C) RT-qPCR analysis of AeAV persistently infected Aa20 cells infected with 0.1 and 1 MOI of DENV-2 using specific primers to the AeAV genome. Error bars represent the SEM of three biological replicates. n.s, not significant; *, p < 0.05; **, p < 0.01.

Figure 10. Aedes anphevirus (AeAV) is potentially vertically transmitted. A) Diagram showing
 the parental (K14, K27) and hybrid strains (GP1, GP2, HP1, HP2) from (49). B) Table showing
 assembly statistics and BLASTN similarity of AeAV assembled from K14 and K27 hybrid strains.

Figure 1





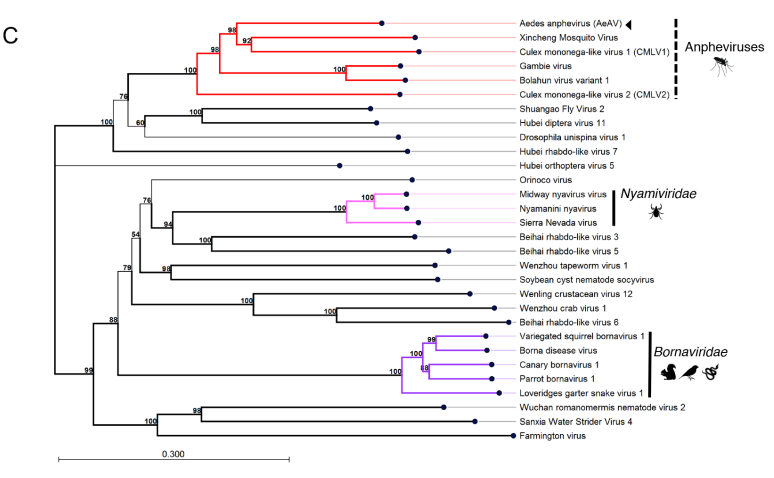
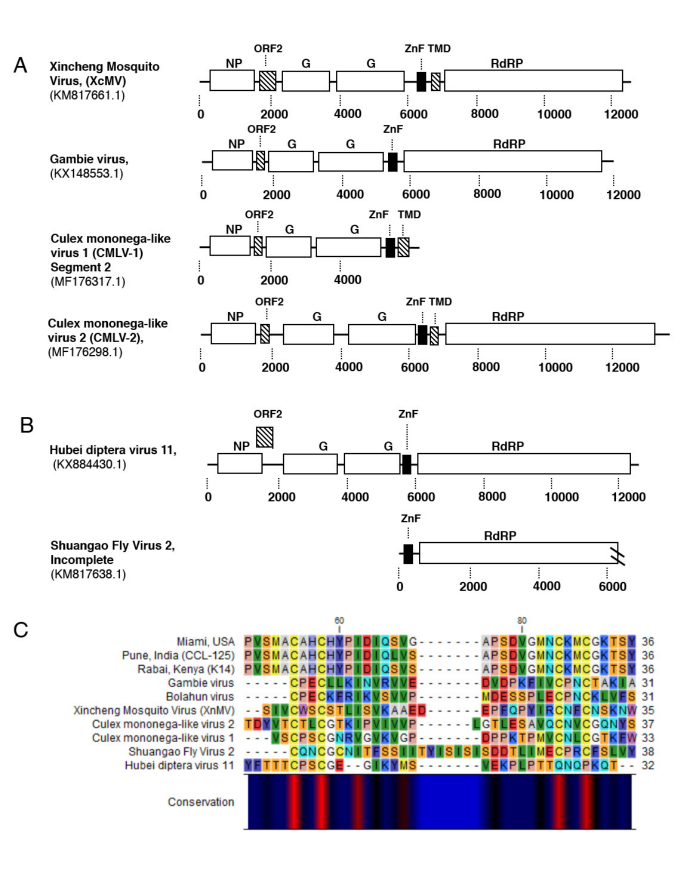
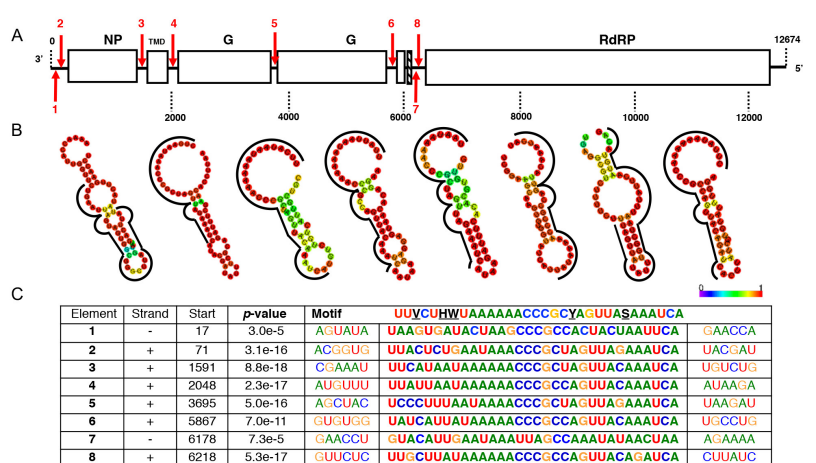
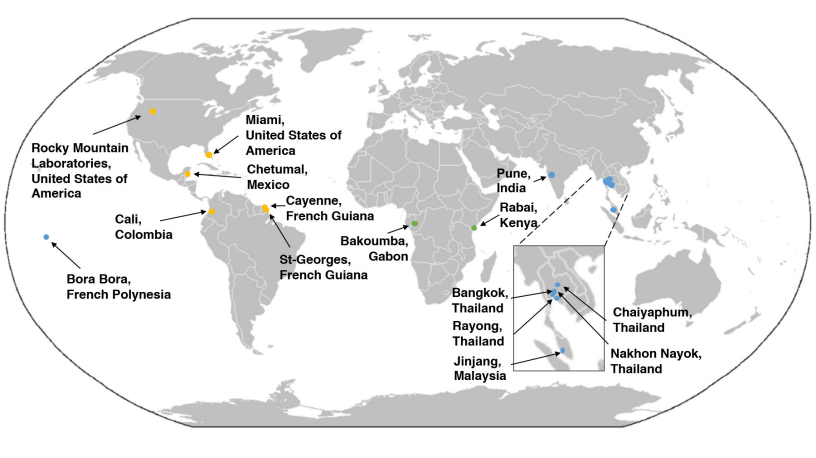
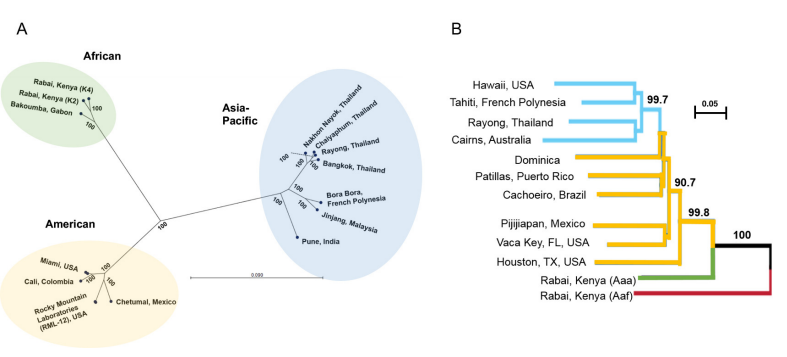


Figure 2

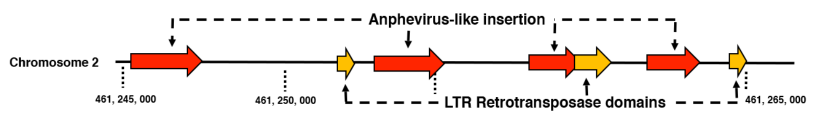


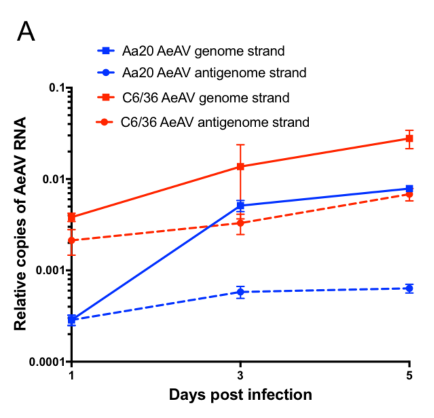




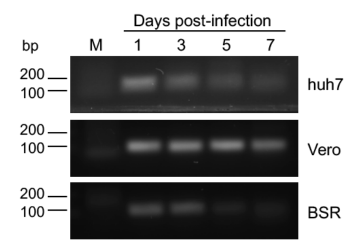


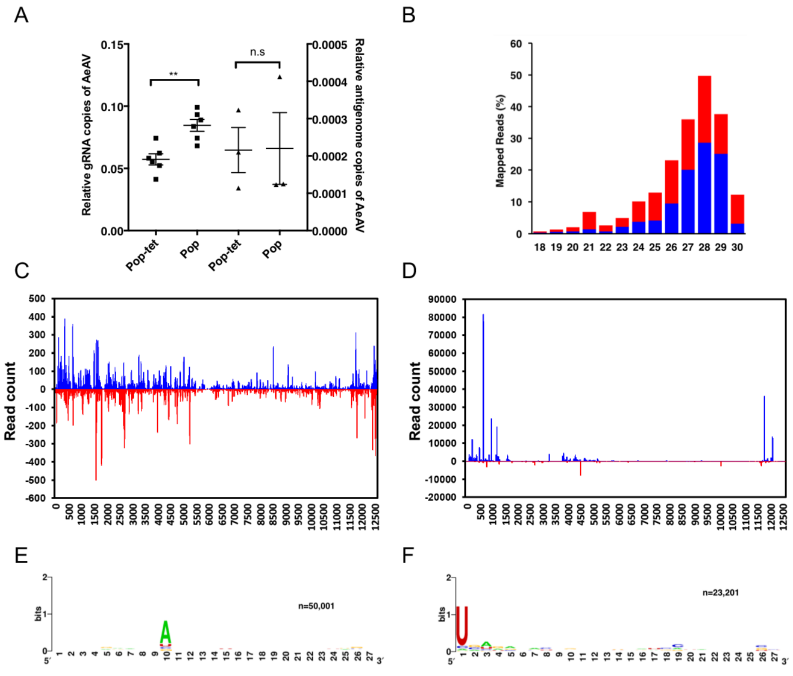




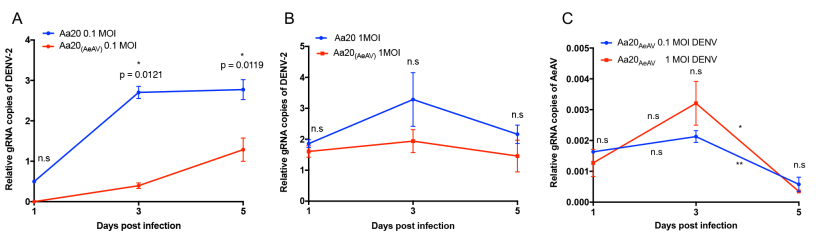


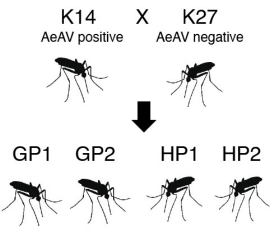












(4/4) AeAV positive

Parental Strains		K14	K27	
Assembled AeAV genome		13,012	-	
Hybrid Strains	GP1	GP2	HP1	HP2
Assembled AeAV genome	13,059	13,034	13,010	7,191; 5,962
BLASTN identity to K14	100%	100%	100%	100%