Zika virus and temperature modulate *Elizabethkingia anophelis* in *Aedes albopictus*

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**Abstract**

**Background:** Vector-borne pathogens must survive and replicate in the hostile environment of an insect’s midgut before successful dissemination. Midgut microbiota interfere with pathogen infection by activating the basal immunity of the mosquito and by synthesizing pathogen-inhibitory metabolites.

**Methods:** The goal of this study was to assess the influence of Zika virus (ZIKV) infection and increased temperature on *Aedes albopictus* midgut microbiota. *Aedes albopictus* were reared at diurnal temperatures of day 28 °C/night 24 °C (L) or day 30 °C/night 26 °C (M). The mosquitoes were given infectious blood meals with 2.0 × 10⁸ PFU/ml ZIKV, and 16S rRNA sequencing was performed on midguts at 7 days post-infectious blood meal exposure.

**Results:** Our findings demonstrate that *Elizabethkingia anophelis albopictus* was associated with *Ae. albopictus* midguts exposed to ZIKV infectious blood meal. We observed a negative correlation between ZIKV and *E. anophelis albopictus* in the midguts of *Ae. albopictus*. Supplemental feeding of *Ae. albopictus* with *E. anophelis aegypti* and ZIKV resulted in reduced ZIKV infection rates. Reduced viral loads were detected in Vero cells that were sequentially infected with *E. anophelis aegypti* and ZIKV, dengue virus (DENV), or chikungunya virus (CHIKV).

**Conclusions:** Our findings demonstrate the influence of ZIKV infection and temperature on the *Ae. albopictus* microbiome along with a negative correlation between ZIKV and *E. anophelis albopictus*. Our results have important implications for controlling vector-borne pathogens.

**Keywords:** Zika virus, *Aedes albopictus*, Microbiome, *Elizabethkingia*

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Background

Zika virus (ZIKV), a Flavivirus of the family Flaviviridae with an ~11-kb positive-sense, single-stranded RNA genome [1], is classified as a member of the Spondweni group [2]. There are two main ZIKV lineages: one lineage with two genotypes from Africa and another from Asia [3, 4].

ZIKV is mainly transmitted by the bite of infected Aedes mosquitoes, although sexual, transplacental, and blood transfusion transmissions have also been documented [5]. There is no Food and Drug Administration (FDA)-approved medication or vaccine to treat or prevent ZIKV infection, despite the World Health Organization (WHO) declaring ZIKV a public health emergency of international concern [6]. Medications for symptomatic relief are the only source of relief for infected individuals [7]. Control measures based on preventing sexual transmission and controlling the
mosquito vector are the main prevention strategies against ZIKV [5]. An upsurge in insecticide resistance, as well as climate change, has expanded the geographical range of vector territories [8], highlighting the urgent need for alternative mosquito control approaches.

The interaction of gut microbial taxa and arboviruses of medical importance has been studied extensively [9–17] due to the potential for microbes to act as candidates for the development of novel therapeutic and transmission-blocking agents.

Flavobacteria have been shown to dominate the Anopheles mosquito midgut [18]. Elizabethkingia (EBK)—a gram-negative, rod-like, aerobic, non-fermenting, non-motile, and non-spore-forming flavobacterium—is widely distributed in Aedes [18–20] and Anopheles [21–23] mosquitoes. EBK has been isolated from both field-caught and laboratory-reared mosquitoes in Africa, Europe, and North America [20, 23, 24] and can be transmitted among mosquitoes through vertical, transstadial, or horizontal transmission [18, 22]. In addition, Elizabethkingia meningoseptica metabolites obtained from Anopheles stephensi midguts possess antiparasitic effects, as well as gametocyte toxicity [25]. Similarly, Elizabethkingia anophelis has a significant effect on Plasmodium parasite development, reducing the oocyst load when introduced at a low bacterial dose [21].

A previous metagenomics study aimed at understanding the effect of ZIKV on the dynamic bacterial community harbored by Aedes aegypti [26] demonstrated that midgut microbes belonging to the family Flavobacteriaceae were the most abundant bacterial taxa among ZIKV-infected mosquitoes. This was particularly the case among the ZIKV-exposed, blood-fed individuals. Flavobacteriaceae abundance was high during the first days of infection but decreased at 7 days post-infection (dpi) among ZIKV-exposed gravid individuals [26]. The aims of this study were to identify the bacterial species within the Aedes albopictus midgut that significantly correlate with ZIKV infectivity and replication, assess whether these microbes possessed antiviral properties, and characterize the influence of temperature on these interactions. Once identified, the symbionts associated with altered vector competence can potentially inform novel vector-borne pathogen control measures.

**Methods**

**Virus**

The ZIKV strain used in this study was HND 2016–19563 (GenBank accession no. KX906952), which was isolated in 2016 in New York state from an infected patient who had traveled to Honduras [27]. To create stock virus for this study, the virus was isolated and passaged three times on Vero cell culture and once on C6/36 cell culture. Stocks were frozen at −70 °C prior to use. Chikungunya virus (CHIKV) IDR 140025961 was isolated from a patient in the Dominican Republic in 2014, while the dengue virus serotype 2 (DENV-2) NI2B strain 306 was isolated from a patient in Nicaragua in 2007. All virus samples were treated similarly.

**Mosquito sampling and screening for ZIKV infection**

The Intergovernmental Panel on Climate Change (IPCC) predicts a 2–4 °C mean global temperature rise over the next century due to global warming [28]. We modeled a 2 °C increase (M temperature regimen) over our baseline temperatures (L temperature regimen), representing the mean temperatures during peak transmission in Suffolk County (NY), where the Ae. albopictus were collected in 2015 (kindly provided by I. Rochlin). Ae. albopictus were colonized at the New York State Department of Health (NYSDOH) Arbovirus Laboratory.

The 15th generation following laboratory colonization of Ae. albopictus were vacuum-hatched and maintained at 28 °C under standard rearing conditions.

To initiate the experiment aimed at exposing them to ZIKV, the mosquitoes were hatched and reared under two thermal conditions: 14 h light at 30 °C/10 h dark at 26 °C (M temperature regimen), or 14 h light at 28 °C/10 h dark at 24 °C (L temperature regimen).

At 8 days post-eclosion, female adults were orally exposed to blood meals consisting of either 1:1 dilution of defibrinated sheep blood plus 2.5% sucrose, sodium bicarbonate (to adjust pH to 8.0) and the virus, or a non-infectious blood meal (NIBM) containing a final concentration of 2.5% sucrose solution. Infectious blood meals contained 2.00E+08 PFU/ml ZIKV HND [27]. The female mosquitoes were exposed to blood meals in a 37 °C preheated Hemotek membrane feeding system (Discovery Workshops, Accrington, UK) with a porcine sausage casing membrane. After 1 h, the mosquitoes were anesthetized with CO₂ and immobilized on a pre-chilled tray connected to CO₂. Engorged females were separated and placed in three separate 0.6 L cardboard cartons (30 individuals/carton). Blood-fed females were maintained on 10% sucrose solution provided ad libitum.

At 7 dpi, the female mosquitoes were immobilized using triethylamine (Sigma-Aldrich, St. Louis, MO, USA). To examine for ZIKV dissemination in the mosquito, the legs were removed before dissections. To assess transmission, saliva was collected under sterile conditions by inserting the proboscis of each female mosquito into a capillary tube containing ~20 µl fetal bovine serum (FBS) plus 50% sucrose 1:1 for 30 min. The mixture was then ejected into 125 µl of mosquito diluent (MD) (20% heat-inactivated FBS in Dulbecco phosphate-buffered
saline plus 50 µg/ml penicillin/streptomycin, 50 µg/ml gentamicin, and 2 µg/ml Fungizone [Sigma-Aldrich, St. Louis, MO, USA]). The midgut was dissected from surface-sterilized individual mosquitoes and rinsed twice in sterile phosphate-buffered saline (PBS) before transfer to sterile microcentrifuge tubes and storage at −80 °C until testing. Mosquito carcass and legs were then placed in individual tubes containing 500 µl MD and a bead. All samples were held at −80 °C until assaying. An absolute quantity of ZIKV in the legs, carcass, and saliva was obtained by isolating RNA using TRIzol (Thermo Fisher Scientific, Waltham, MA, USA). A ZIKV-specific quantitative polymerase chain reaction (qPCR) assay targeting the NS1 region was used for detection (Table 1). Serially diluted ZIKV of known quantities was used to generate the standard curve and extrapolate viral load (Additional file 1: Figure S1). Midguts obtained from Ae. albopictus adults maintained on 10% sucrose solution were used as a control.

Sequencing and analysis of microbiome of Aedes mosquito midguts
A total of 29 samples were included in this study: 10 midguts exposed to infected blood meal (IBM), 10 midguts exposed to NIBM, and nine midguts exposed to 10% sucrose only (non-blood-fed [NBF]), along with one negative control (NC), one non-template control (NTC), and one positive spike (PC). Ae. albopictus midgut, legs, carcass, and saliva samples were screened for virus at 7 dpi. The infection and dissemination status of each individual was determined prior to downstream processing. The complementary DNA (cDNA) from individual midguts was prepared from the RNA samples using the iScript cDNA kit (Bio-Rad, Hercules, CA, USA). Thereafter, 16S rRNA was amplified using the 16S primer set and conditions from Table 1.

The barcoded high-throughput amplicon sequencing of the bacterial 16S V3-V4 hypervariable rRNA was completed at Wadsworth Center’s Advanced Genomic Technologies Core (WCAGT). PCR reactions were carried out in a total volume of 50 µl: 5 µM of 16S rRNA V3-V4 hypervariable region primers consisting of an Illumina barcode (Table 1), 2 µl cDNA, 8 µl deionized filter sterilized water, and 36 µl AccuStart II PCR SuperMix (Quanta Biosciences, Beverly, MA, USA). To ensure identification of any potential contamination at either the PCR stage or the sequencing step, two negative controls (water and non-template control) were included at the PCR step and sequenced. A positive spike was included during the sequencing run. A fragment size of ~460 base pairs (bp) of each sample, as well as the negative controls, was submitted to the WCAGT for sequencing. Automated cluster generation and paired-end sequencing (250-bp reads) were performed using the Illumina MiSeq 500-cycle reagent kit.

| Table 1 | Primer sequences and reaction conditions |
|---------|------------------------------------------|
| Primer name | Primer sequence | Annealing temperature (°C) |
| Ribosomal protein S7_F | GAG ATC GAG TTC AAC AGC AAG A | 55 |
| Ribosomal protein S7_R | GAG AAG TTC TTC TCA AGC TCA C | 55 |
| Elizabethkingia_upoB_F | CTC CGG AAG GAC CAA ACA TTG | 55 |
| Elizabethkingia_upoB_R | CAA CCG TCC AGT CAG ATC C | 55 |
| Elizabethkingia_upoB_F | GCT AAG ACA CAT GGT TGA TGA | 55 |
| Elizabethkingia_upoB_R | ACC ACC GAA CTG AGC TTT AC | 55 |
| 16S Microbiome_F | TCG TCG GCA GCG TCA GAT GTG TAT TAT AAG AGA CAG CCT ACG GGN GGC WGC AG | 55 |
| 16S Microbiome_R | GTC TCG TGG GCT CGG AGA TCT GTA TAA GAG ACA GGA CTA CHV GGG TAT CTA ATC C | 60 |
| ChikF1 | AAGCTCGGCTCTTTCCAAAG | |
| ChikR1 | CCAATTTGCTCCTGTCTCTCCT | 60 |
| Chik Probe [60, 61] | CCAATTTGCTCCTGTCTCTCCT | 60 |
| DenF1 | AAGGACTAGAGGTATTAGAGGAGACC | 60 |
| DenR1 | GGGCTGTCGCTGCTGCATTGATG | 60 |
| Den Probe 1–3 [62] | AACAGCATATGACGCTGGGAGAGACC | 60 |
| ZIKV 1086 | CGGCTGCCCACAAGCAAG | 60 |
| ZIKV 1162c | CCACAATGCCTCTTTCCAGACAT | 60 |
| ZIKV 1107-FAM [4] | AGSCTACCTTGACAAGCAGTCAGACACTCAA | 60 |
Data analysis was carried out on QIITA (https://qiita.ucsd.edu/) and MicrobiomeAnalyst [29]. In summary, to convert the de-multiplexed FASTQ files to the format used by QIITA, the command split libraries FASTQ was used. The clustering of the sequences into operational taxonomic units (OTUs) was done via closed-reference OTU picking based on the Greengenes 16S reference database, which matches the sequences based on a 97% sequence identity threshold before assigning taxonomies. A BIOM-formatted OTU table was subsequently generated, and in order to reduce the potential for alpha and beta diversity biases, the data were rarefied to a depth of 30,000 reads per sample. Finally, to visualize the taxonomic profiles of each sample, taxa bar plots were generated using the rarefied data artifact. Following the generation of the relative quantity of each taxon, we generated taxa which were deposited in the NCBI GenBank Short Read Archives with accession numbers PRJNA608507, PRJNA608505, PRJNA608503, PRJNA608501, PRJNA608500, PRJNA608499, PRJNA608472, PRJNA608496, PRJNA608495, PRJNA608493, PRJNA608492, PRJNA608491, PRJNA608489, PRJNA608483, PRJNA608471, PRJNA608482, PRJNA608470, PRJNA608456, PRJNA608450, PRJNA608481, PRJNA608449, PRJNA608448, PRJNA608480, PRJNA608479, PRJNA608477, PRJNA608437, PRJNA608434, PRJNA608427, and PRJNA608428.

The alpha diversity profiling (number of species of taxa in a single sample) was calculated using the Chao1 diversity measure, and significance testing was done using analysis of variance (ANOVA) in MicrobiomeAnalyst. The Chao1 index estimates the richness of taxa in a sample by extrapolating the number of rare organisms that may have been omitted due to under-sampling. The index assumes that the number of observations for a taxon has a Poisson distribution and thus corrects for variance [30]. Beta diversity, a comparison of microbial community composition between two samples or two conditions, was analyzed by utilizing the Bray–Curtis dissimilarity statistic [31]. The distance matrix generated after comparing each sample to the other was visualized for dissimilarity between samples using the principal coordinates analysis method.

Phylogenetic analysis of the Elizabethkingia (EBK) isolated from Ae. albopictus midguts

In order to validate the identity of EBK taxa identified in the Ae. albopictus midguts, the rpoB gene (accession number MT096047) was amplified from the midgut cDNA using the rpoB primer pairs and PCR conditions as described in Table 1. The 558-bp PCR amplicon was excised from an agarose gel and cleaned using the QIAquick gel extraction kit (Qiagen, UK). The amplicons were Sanger-sequenced at the WCAGT. To construct the EBK rpoB phylogenetic tree, Molecular Evolutionary Genetic Analysis (MEGA) software [32] was utilized.

A total of nine gene sequences published with GenBank accession numbers CP023010.2, KY587659.1, CP016377.1, KY587657.1, CP035811.1, CP035809.1, CP011059.1, CP039929.1, and MN327643.1 were added to the phylogenetic analysis.

Clustering and biomarker analysis

Patterns of relative abundance of taxa species in response to different experimental factors were analyzed in MicrobiomeAnalyst. To provide an estimate of the most important taxa for classification of data resulting from different experimental factors, biomarker analysis was carried out using the random forest algorithm within MicrobiomeAnalyst. The default setting of the number of trees to grow and number of predictors to try (500 and 7, respectively) was applied with the randomness setting left on. The random forest algorithm is a supervised classification algorithm of trees created by using bootstrap samples with training data and random feature selection in tree induction. It is an ensemble of unpruned classifications or regression trees trained with the bagging method [33].

Analysis of the impact of supplemental feeding of Ae. albopictus with E. anophelis aegypti on their survivability and vector competence for ZIKV

The Elizabethkingia strain associated with the Ae. albo - pictus in this study was identified as E. anophelis aegypti (Fig. 3). Elizabethkingia anophelis aegypti cultured from Ae. aegypti midguts was used for supplemental feeding of Ae. albopictus, which were reared to adult stage as described previously. Septic adult Ae. albopictus were fed a supplemental diet of E. anophelis aegypti grown to optical density at 600 nm (OD$_{600}$) = 1.0 [(3.16E$^{+08}$ colony-forming units per milliliter (CFU/ml)] (N = 30) and OD$_{600}$ = 0.5 (1.0E$^{+08}$ CFU/ml) (N = 30), after which the survivability was assessed. Sterile PBS was used as a negative control (N = 30), and the differences in survivability were calculated using the Log-rank test.

To assess the impact of E. anophelis aegypti supplemental feeding on Ae. albopictus vector competence for ZIKV, Ae. albopictus mosquitoes were reared to the adult stage, and upon emergence, Ae. albopictus adults were antibiotic-treated with 20% penicillin, 20% streptomycin, and 75ug/ml gentamicin in a 10% sucrose solution for 8 days (Fig. 4b). The cotton strips that were used to deliver the antibiotic treatment were changed daily. Both groups were maintained on filter-sterilized 10%
sucrose solution. The effectiveness of the sterility rearing was confirmed by CFU assays on randomly selected adult female midguts prior to blood-feeding or bacterial challenge (Fig. 4c). *Elizabethkingia anophelis aegypti* was introduced to a total of *n* = 75 antibiotic-treated *Ae. albopictus* through a sugar meal (cotton strips moistened with 1.5% sucrose solution containing *E. anophelis aegypti* to a final concentration of 1.0E+08 CFU/ml) [9]. The control consisted of septic *Ae. albopictus* (*n* = 75) without exposure to *E. anophelis aegypti*. The cotton strips were withdrawn after 24 h and replaced with sterile cotton strips moistened with sterile 10% sucrose solution. Both the experimental (antibiotic-treated *Ae. albopictus* infected with *E. anophelis aegypti*) and control groups (septic *Ae. albopictus*) were deprived of sucrose solution for 18–24 h and offered 2.00E+08 PFU/ml ZIKV blood meal as described previously. The midgut, carcass, saliva, and legs were obtained from each individual of each population at 7 dpi. RNA was isolated from the carcass, saliva, and legs, and screened for ZIKV. The infection, dissemination, and transmission percentage rates were calculated and compared between the two groups using Chi-square tests. The assay was carried out in two independent experiments.

**Assessment of the broad antiviral effect of E. anophelis in vitro**

The antiviral effect of *E. anophelis aegypti* was assessed in the Vero and C636 cell lines. A total of 3 ml/well of cells (2.0 x 10^5 cells/ml) was used to seed six-well plates that were incubated at 37 °C (Vero) and 28 °C (C636) and 5% CO₂ until they attained 90% confluence. Upon attaining 90% confluence, the experimental wells were inoculated with 75 μl of 1.0E+08 CFU/ml of *E. anophelis aegypti* or *Escherichia coli*, while the control wells were inoculated with either *E. anophelis aegypti* or *E. coli* that were heat-inactivated for 6 h at 90 °C. The experiment was performed on two separate days, and each assay was performed in triplicate.

At 24 h post-infection (hpi) with bacteria (*E. anophelis aegypti* or *E. coli*), Vero cell monolayers were infected at a multiplicity of infection (MOI) of 0.1 infectious particles/cell with ZIKV, CHIKV, or DENV. C636 cells were inoculated with ZIKV at 24 hpi with *E. anophelis aegypti* or *E. coli*. The supernatant was harvested at 24 hpi with the virus. The viral load was compared between the experimental and control supernatants. RNA was extracted from the supernatant using the TRIzol procedure (Thermo Fisher Scientific, Waltham, MA, USA) and the virus quantified by qPCR as described previously, using the primers in Table 1. To assess the effect of timing on virus replication, 75 μl of 8.0 log₁₀ CFU/ml of *E. anophelis aegypti* or *E. coli* was co-inoculated with 0.1 MOI of ZIKV. Differences in virus quantities were measured using ANOVA in GraphPad Prism.

In order to assess the cell-independent effect of *E. anophelis aegypti* antiviral activity, U4.4 cells (*Ae. albopictus*) were co-infected with *E. anophelis aegypti* and ZIKV.

The *E. anophelis aegypti* bacterial strain was streaked on Luria-Bertani (LB) plates and incubated in a 10% CO₂ incubator at 28 °C overnight. A single colony was picked using a sterile plastic loop and used to inoculate 500 ml of sterile LB broth. The inoculated broth medium was incubated overnight at 28 °C in an orbital shaker shaking at 225 rpm. The *E. anophelis aegypti* overnight culture was subsequently diluted to OD_{600} = 1.0 (8.5 log₁₀ CFU/ml), OD_{600} = 0.5 (8.0 log₁₀ CFU/ml), and OD_{600} = 0.2 (7.7 log₁₀ CFU/ml).

Individual wells were seeded with 3 ml of 2 x 10^5 U4.4 cells/ml suspended in minimum essential medium without FBS and incubated at 28 °C for 1 h before adding FBS to each well at a final concentration of 20%. The U4.4 cells were then incubated for 5 days in a 10% CO₂ incubator at 28 °C. After 5 days, the supernatant was carefully removed, and the plates were infected with *E. anophelis aegypti* as described previously at OD_{600} = 1.0 (8.5 log₁₀ CFU/ml), OD_{600} = 0.5 (8.0 log₁₀ CFU/ml), and OD_{600} = 0.2 (7.7 log₁₀ CFU/ml) for three biological replicates. After 1 h of incubation at 28 °C, 3 ml of M&M media containing 20% FBS was added to each well and incubated at 28 °C.

To determine whether the U4.4 cells were successfully infected with *E. anophelis aegypti*, a single well from each replicate of the different dilutions of *E. anophelis aegypti* infection was harvested at 2 dpi.

The cells were spun at 1000 rpm and washed three times with sterile PBS. After the third wash, 100 μl of the washed cells was used to inoculate 3 ml of LB media that was incubated overnight at 28 °C in an orbital shaker at 225 rpm. The experimental wells were infected with 0.1 MOI ZIKV as described previously, as were the ZIKV quantities. Supernatants were harvested from the two treatments at both 2 dpi and 4 dpi. RNA was extracted from the cells using TRIzol (Thermo Fisher Scientific, Waltham, MA, USA). The ZIKV quantities were measured as described previously.

**Results**

**Influence of temperature, blood feeding, and ZIKV infection on microbial profile of Ae. albopictus**

A total of 12 out of 30 *Ae. albopictus* reared in the L temperature regimen disseminated the virus, with a virus titer that ranged from 1 to 2 log₁₀ PFU/ml, while 26 out of 30 individuals reared in the M temperature regimen disseminated the virus, with a virus titer that ranged from 1
to $5 \log_{10}$ PFU/ml (Additional file 5). None of the individuals reared in the L temperature regimen transmitted the virus in the saliva, while three individuals reared in the M temperature regime had positive saliva. The viral titer range in saliva was $1–2 \log_{10}$ PFU/ml (Additional file 6).

Illumina MiSeq 16S rRNA sequencing resulted in 6,089,629 reads. The number of reads for individual samples ranged from 531,220 to 34,833, with a mean of 152,448. Both the positive spike-in (PhiX) and negative controls (water and non-template control) had a limited number of sequence reads, ranging from 1620 to 2823, and hence were discarded in downstream analysis. In order to control for alpha and beta diversity biases, each sample was rarefied to a total frequency of 34,000 reads.

A total of 1878 OTUs were identified in this study. The core phyla identified were Bacteroidetes, Proteobacteria, Actinobacteria, Cyanobacteria, and Firmicutes.

Overall, midgut taxa diversity was impacted by feeding [permutational multivariate analysis of variance (Beta-diversity analysis; PERMANOVA test $F$-value = 13.22; $R$-squared = 0.5042; $P$-value < 0.001; Fig. 1a)]. Increasing the rearing temperature did not result in significant changes in taxa diversity (Beta-diversity analysis; PERMANOVA test; $F$-value: 2.1707; $R$-squared: 0.074415; $P$-value < 0.1; Alpha-diversity index: Chao1: $p$-value: 0.85872; $t$-test = 0.17997; $P = 0.9$; Fig. 1b and Additional file 2: Figure S2). Despite this, at the individual taxa level, Elizabethkingia was unique in that it was identified at higher proportions at the lower rearing temperature regime (Fig. 2b, Additional file 3: Figure S3). Blood meal intake, irrespective of the infection status, decreased the taxon diversity (PERMANOVA; $F$-value: 20.007; $R$-squared: 0.42562; $P$-value < 0.001; Fig. 1c). Additionally, ZIKV infectious blood meal uptake significantly reduced the individual midgut bacterial diversity (Alpha-diversity

**Fig. 1**  Microbial taxa diversity identified in *Ae. albopictus* midguts.  

**a**  The meal component significantly clustered the midgut taxa [PERMANOVA] $F$-value: 13.22; $R$-squared: 0.5042; $P$-value < 0.001.  

**b**  Increase in temperature did not have an impact on *Ae. albopictus* midgut microbial richness [PERMANOVA] $F$-value: 2.1707; $R$-squared: 0.074415; $P$-value < 0.1.  

**c**  Blood meal intake, irrespective of the infection status, altered the midgut microbial taxa [PERMANOVA] $F$-value: 20.007; $R$-squared: 0.42562; $P$-value < 0.001.
Elizabethkingia was significantly associated with the midgut exposure to an infectious blood meal (Fig. 2c).

**Phylogenetic analysis of Elizabethkingia**

To taxonomically classify the *Elizabethkingia* strain that we identified in the midgut of the *Ae. albopictus*, we sequenced the *Elizabethkingia rpoB* gene amplified from the midgut cDNA of *Ae. albopictus*. The phylogenetic analysis demonstrated that the *Elizabethkingia* strain identified in our study has 100% sequence homology to *E. anophelis*, which was originally isolated from the midgut of *Anopheles gambiae* [34]. Our strain has 99% homology to *E. endophytica*, originally isolated from sweet corn [35], and was closely related to the *E. meningoseptica* genomospecies 4 [36], *E. miricola* (isolated from condensation water in the space station) [37], and *E. bruuniana* [38]. *Elizabethkingia anophelis* has 92% homology to *E. meningoseptica*, originally isolated from human cerebrospinal fluid [39] (Fig. 3).

**Elizabethkingia anophelis colonization of *Ae. albopictus* midguts is associated with reduced ZIKV infection rates and viral loads**

We assessed the impact of *E. anophelis aegypti* colonization of *Ae. albopictus* midguts on survivability and vector competence for ZIKV. Colonization of *Ae. albopictus* midguts with 3.16E+08 CFU/ml and 1.0E+08 CFU/ml did not impact their survivability [log-rank (Mantel-Cox)
test \([\chi^2 = 2.345; \text{df} = 2; \ P = 0.31]\) (Fig. 4a). To study the impact of \textit{E. anophelis aegypti} on the vector competence of \textit{Ae. albopictus} for ZIKV, conventionally reared \textit{Ae. albopictus} was utilized as a positive control; antibiotic-treated \textit{Ae. albopictus} exposed to 1.0E+08 CFU/ml \textit{EBK} constituted the experimental group. The \textit{E. anophelis aegypti}-treated group demonstrated increased blood-feeding rates (40/75) relative to the conventionally reared individuals (19/75) (Chi-square test \([\chi^2 = 12.3207; \ P = 0.0004]\) (Additional file 7). In spite of the differences in blood meal intake, \textit{E. anophelis aegypti}-treated individuals demonstrated a 14% lower average ZIKV infection rate than the conventionally reared \textit{Ae. albopictus}, which was 37% (Fisher’s exact test, \(P = 0.0003; \text{Fig. 4d}\)). In spite of the differences in blood meal intake, \textit{E. anophelis aegypti}-treated individuals demonstrated a 14% lower average ZIKV infection rate than the conventionally reared \textit{Ae. albopictus}, which was 37% (Fisher’s exact test, \(P = 0.0003; \text{Fig. 4d}\)). There was no difference observed in the dissemination rates [conventionally reared (5%); \textit{E. anophelis aegypti}-treated (7%) (Fisher’s exact test, \(P = 0.7673\)], and no mosquito transmitted the virus at the measured time point (Additional file 8). Despite the fact that feeding on a ZIKV infectious blood meal was associated with a higher percentage of \textit{E. anophelis aegypti} relative to other taxa (Fig. 2a), the fold change in expression of the \textit{Elizabethkingia rpoB} gene from ZIKV-infected \textit{Ae. albopictus} \((N = 22)\) correlated negatively with ZIKV absolute quantities (Spearman \(r = −0.3985; \ P < 0.05\); (Fig. 4e). ZIKV viral loads ranged from 1.00E+03 to 1.00E+11 ZIKV RNA copies/ml, while the log fold change of \textit{E. anophelis aegypti} ranged from 0.016 to 158.49 CFU/ml (Additional file 9).

**Infection of Vero cells with \textit{E. anophelis} results in a broad-spectrum reduction of viral load**

To clarify the potential for a broad-spectrum antiviral effect of \textit{E. anophelis aegypti}, we performed in vitro assays on both \textit{Ae. albopictus} and mammalian cells. A broad-spectrum \textit{E. anophelis aegypti} antiviral effect
was measured by significantly lower viral loads associated with Vero cells sequentially inoculated with *E. anophelis aegypti/ZIKV* (ANOVA test, \( P < 0.0001 \)).
and *E. anophelis aegypti*/DENV (ANOVA test, *P* < 0.0001; Fig. 5a; Additional file 10).

This antiviral property was specific to *E. anophelis aegypti*. We demonstrated no difference in viral loads when *E. anophelis aegypti* was replaced by *E. coli*, *E. coli*/ZIKV (ANOVA test, *P* = 0.9867), *E. coli*/CHIKV (ANOVA test, *P* > 0.9999), or *E. coli*/DENV (ANOVA test, *P* = 0.9987; Fig. 5b; Additional file 11). It is noteworthy that the antiviral effect was only measured when *E. anophelis* and the virus were inoculated sequentially, but not when co-inoculated with *E. anophelis aegypti*/ZIKV (ANOVA test, *P* = 0.6433; *E. anophelis aegypti*/ZIKV (ANOVA test, *P* = 0.9987; Fig. 5c; Additional file 12). When both *E. anophelis aegypti* and *E. coli* were used to inoculate C636 cells, we observed excessive bacteria growth (*E. anophelis aegypti* and *E. coli*) after 24 hpi, and ZIKV replication was attenuated in C636 cells co-inoculated with *E. anophelis aegypti*/ZIKV (ANOVA test, *P* = 0.9987; Additional file 12). When both *E. anophelis aegypti* and *E. coli* were used to inoculate C636 cells, we observed excessive bacteria growth (*E. anophelis aegypti* and *E. coli*) after 24 hpi, and ZIKV replication was attenuated in C636 cells co-inoculated with *E. anophelis aegypti*/ZIKV (ANOVA test, *P* = 0.9987; Additional file 12). Viability assays demonstrated 90% viability of *E. anophelis aegypti*-infected cells at 2 dpi, suggesting direct cell killing was not responsible for decreased viral replication (Additional file 14). We performed an in vitro assay of *E. anophelis aegypti*/ZIKV on the *Ae. albopictus*-derived U4.4 cell line and measured a significant reduction in ZIKV titers in all *E. anophelis aegypti*/ZIKV-infected U4.4 cells relative to the control (ANOVA; Kruskal–Wallis test, *P* < 0.01; Fig. 6b; Additional file 15).

**Discussion**

Recently, ZIKV has become a focus of intense research due to its unprecedented global spread, [4, 40–44] leading to outbreaks in different parts of the world. An upsurge in insecticide resistance, as well as climate change, has resulted in expansion of the geographical range of vector territories [8], which suggests an urgent need for alternative approaches to control mosquitoes and the pathogens they transmit.

Due to their potential as candidates for development of therapeutics and transmission-blocking agents, investigators have studied the interaction of gut microbial taxa and arboviruses of medical importance [9–17]. *Aedes* midgut bacterial species associated with ZIKV are poorly understood. The aim of this study was to assess the impact of increased temperature on the *Ae. albopictus* midgut microbiome and identify bacterial species that are associated with altered ZIKV infectivity and *Ae. albopictus* competence.

**Reduction in abundance of midgut *E. anophelis* due to temperature increase**

Given that the global increase in temperature is projected to accelerate [45, 46], a comprehensive
assessment of the potential effects of climatic change on vectorial capacity requires an assessment of the influence of temperature on the mosquito microbiome. Studies have shown that temperature is an important component in shaping the microbial communities of organisms [47–51]. In an effort to understand how climate change might ultimately alter microbial taxa that influence vectorial capacity for ZIKV, we measured the effect of increases in temperature on the midgut microbial community. Specifically, we observed a reduction in levels of E. anophelis albopictus, which our data suggest is associated with decreased levels of ZIKV and competence. These data suggest that future increases in temperatures could alter microbial profiles and therefore further exacerbate documented increases in transmission resulting directly from increases in viral replication at higher temperatures.

Impact of ZIKV infection and blood meal on Ae. albopictus gut microbiota diversity
A significant reduction in Ae. albopictus midgut microbial diversity resulting from ZIKV infection and/or blood meal digestion was evident, yet the effect on individual genera was highly variable. Our results are consistent with the findings of a previous study assessing the total gut microbiota diversity of Aedes japonicus and Aedes triseriatus following infection with La Crosse orthobunyavirus, which revealed higher midgut bacterial richness among the midguts obtained from newly emerged individuals versus lower richness among midguts exposed to viral infection or a non-infectious blood meal [52]. In addition, our study corroborates the finding of a study aimed at assessing the impact of West Nile virus infection on the bacterial diversity of Culex pipiens, which revealed higher bacterial diversity at the genus level among the unexposed individuals relative to both exposed infected and the exposed uninfected individuals [53].

Mosquitoes exposed to non-infectious blood meals demonstrated substantial increases in the relative abundance of E. anophelis albopictus as compared to unfed individuals, clearly indicating that the gut environment is more amenable to this species’ growth following blood meal digestion. This alteration in microbial composition could be beneficial to Ae. albopictus. Specifically, E. anophelis has been shown to produce OxyR regulon and antioxidants that, in turn, could protect the mosquitoes from the oxidative stress associated with blood digestion. Elizabethkingia anophelis has also been shown to possess hemolytic activity and may therefore help facilitate erythrocyte digestion in the mosquito midgut [54].
Correlation of native with viral load suggest block viral infection, both the in vitro data and negative these data suggest that may primarily act to corroborates the results of other studies [55–59]. While not translate to increased virus dissemination, which [21]. Taken together, these results suggest that E. anopheles aegypti maintains its antiviral activity with both intact and deficient RNA interference responses. Past studies show that E. anophelis attenuates Plasmodium parasite development [21]. Taken together, these results suggest that E. anophelis has the capacity to directly or indirectly elicit a broad antipathogen phenotype. Further studies delineating the potential antiviral mechanism of E. anophelis are needed, as this could lead to the development of novel control measures.

We observed higher infection rates in the conventionally reared Ae. albopictus relative to the E. anophelis aegypti-infected Ae. albopictus group. However, this did not translate to increased virus dissemination, which corroborates the results of other studies [55–59]. While these data suggest that E. anophelis may primarily act to block viral infection, both the in vitro data and negative correlation of native E. anophelis with viral load suggest that increasing doses could further inhibit viral replication and dissemination. An antiviral effect was only measured when E. anophelis aegypti and the virus were inoculated sequentially. While the mechanistic basis is unknown, this suggests that establishment and/or replication of E. anophelis is required to effectively perturb viral infection and replication.

Conclusions
To the best of our knowledge, our study provides the first description of broad-spectrum antiviral association of E. anophelis. We demonstrate both a negative correlation of E. anophelis and ZIKV in Ae. albopictus and an association with reduced viral loads of ZIKV, DENV, or CHIKV in vitro. Further, E. anophelis was found to be temperature-sensitive, with decreased levels identified at higher temperatures. Together, these findings have significant implications for our understanding of the interactions of bacterial and viral agents in mosquitoes, not to mention how climate change could influence these interactions. Identifying the mechanistic basis for these associations could improve our understanding of disease epidemiology and ultimately inform the development of novel vector control measures.

Abbreviations
ZIKV : Zika virus; EBK : Elizbethkingia; DENV: Dengue virus; CHIKV : Chikungunya virus; NY: New York state; NYS: New York State Department of Health; MD : Mosquito diluent; PBS: Phosphate-buffered saline; IBM: Infectious blood meal; NIBM : Non-infectious blood meal; NBF: Non-blood-fed; NC: Negative control; NTC: Non-template control; PC: Positive control; OTUs: Operational taxonomic units; MOI: Multiplicity of infection; Hpi: Hours post-infection; CFU: Colony-forming units; MEGA: Molecular Evolutionary Genetic Analysis.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s13071-021-05069-7.
the current study. AV, DN, KL and CA: Involved in the study design and participated in the writing of the paper. All authors read and approved the final manuscript.

Funding
This publication was supported by the Cooperative Agreement Number U01CK000509 funded by the Centers for Disease Control and Prevention. Its content is solely the responsibility of the authors and do not necessarily represent the official views of the Centers for Disease Control and Prevention or the Department of Health and Human Services. These studies were also partially funded by the National Institutes of Health awards R21AI131683 and R21 AI138344.

Availability of data and materials
Datasets generated during the current study are available in GenBank, as well as included in this manuscript.

Declarations

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
The authors declare they have no competing interests.

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Received: 5 August 2021 Accepted: 15 October 2021

Published online: 12 November 2021

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