Emergence of Zika virus: Role of founder effects, drift and mutational reversions in spread to the Americas

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Summary Paragraph

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Mosquito-borne viruses have recently spread globally, with major impacts on human health. Zika virus (ZIKV) emerged from obscurity in 2013 to spread from Asia to the South Pacific and the Americas, where millions of people were infected. For the first time, severe clinical manifestations, including Guillain Barré syndrome and congenital defects to the fetus of pregnant women, were detected. Phylogenetic studies have shown that ZIKV evolved in Africa and later spread to Asia, and that the Asian lineage is responsible for the recent epidemics. However, the reasons for the sudden emergence of ZIKV remain incompletely understood. Accumulating evidence on other arboviruses like chikungunya and West Nile suggest the likelihood that viral mutations could be determinants of change in ZIKV transmission efficiency responsible for efficient spread. Using evolutionary analyses, we determined that four mutations, which occurred just before ZIKV introduction to the Americas, represent direct reversions of previous mutations that accompanied spread many decades ago from ZIKV's native Africa to Asia and early circulation there. Experimental infections of mosquitoes, human cells, and mice with ZIKV strains with and without these mutations demonstrated that the original mutations reduced fitness for urban transmission, while the reversions restored fitness. increasing epidemic risk. Our findings help to explain the recent ZIKV emergence and underscore the major yet unpredictable impacts of founder effects and genetic drift on viral emergence. Because they involve random mutations that become fixed in a population when it undergoes bottlenecks during geographic spread and inefficient transmission, founder effects and other forms of drift can limit the ability of mosquitoborne viruses like ZIKV to emerge. Furthermore, their stochastic nature limits our ability to predict epidemics.

Zika virus (ZIKV), an arthropod-borne virus (arbovirus) in the genus *Flavivirus* discovered in 1947¹, remained obscure with little association with human disease until 2007. Then, small outbreaks occurred in Gabon² and Yap Island in Micronesia³, associated with mild dengue-like illness. In 2013, ZIKV spread to the South Pacific, where approximately half of the residents of French Polynesia were infected⁴, and an association with Guillain Barré syndrome (GBS) was discovered⁵. Spread though other areas of the South Pacific followed, and the first outbreak ever detected in the Americas was identified in 2015 northeastern Brazil⁶, accompanied by a dramatic increase in the incidence of microcephaly⁷ and other congenital malformations now termed congenital Zika syndrome (CZS)⁸. ZIKV continued to spread to nearly all countries in the Americas with the continued association with GBS and CZS, leading the World Health Organization to declare in 2016 a Public Health Emergency of International Concern.

Early phylogenetic studies combined with ZIKV detections and seroprevalence indicated that the virus originated in and remains widespread in Africa, and was introduced many decades earlier to Asia⁹. However, the reason(s) for its recent, dramatic spread to the Americas remain enigmatic. One hypothesis is that ZIKV increased its ability to be transmitted efficiently by mosquito vectors through adaptive mutations. This mechanism has been demonstrated for other arboviruses including West Nile¹⁰, Venezuelan equine encephalitis¹¹ and chikungunya (CHIKV) viruses¹². RNA arboviruses exhibit high mutational frequencies due to their lack of proof-reading during genome replication, providing the opportunity for rapid adaptation to changing infection and transmission opportunities¹³. The first evidence supporting this adaptive evolution hypothesis for ZIKV came from studies of an A188V amino acid substitution in the nonstructural protein 1 (NS1-A188V) that occurred just before ZIKV spread to the South Pacific and the Americas. This substitution slightly enhances ZIKV transmission by *Aedes* (*Stegomyia*) *aegypti* mosquitoes¹⁴, the principal epidemic vector¹⁵.

Directly reverting Zika virus mutations. To further test the hypothesis that adaptive evolution enhanced epidemic ZIKV transmission, leading to unprecedented spread, millions of infections, and the resulting detection of rare disease manifestations, we performed phylogenetic analyses to identify additional amino acid substitutions that preceded its spread from Asia to the Americas. Extended Data Fig. 1 shows a tree with four substitutions mapped to its branches that preceded epidemic spread, including NS1-A188V⁴. Tracings of these amino acids in ZIKV phylogenetic trees are shown in Extended Data Fig. 2. None of these mutations was noted in earlier, comprehensive phylogenetic studies¹⁶. Strikingly, when we traced the evolution of these 4 pre-epidemic amino acid substitutions, we found that they represented direct reversions of earlier mutations inferred to have accompanied spread from Africa to Asia. or early circulation in Asia. The simplest explanation for this mutation pattern is that that the preepidemic substitutions restored fitness (reproduction and survival ability) declines caused by founder effects and/or genetic drift during early, inefficient A. aegypti-borne interhuman transmission. This transmission inefficiency is consistent with the lack of recognized outbreaks in Asia before 2010, and bottlenecks that accompany the arbovirus transmission cycle can allow drift to dominate evolution if populations remain small.

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Founder effects result in a loss in genetic diversity that accompanies geographic introductions of small founder populations from a large, ancestral population. For humanamplified arboviruses like ZIKV, these typically involve a single infected traveler¹⁷. Furthermore, several virus population bottlenecks are known to punctuate the arbovirus transmission cycle during mosquito infection and dissemination of virus to the salivary glands. The resultant loss of genetic diversity can result in the fixation of random mutations, the majority of which are by chance deleterious for RNA viruses¹⁸ and other organisms. The hypothesis that African ZIKV strains lost fitness upon their introduction into Asia is also supported by their greater infectivity for A. aegypti mosquitoes compared to Asian and subsequent American ZIKV strains 19-21, as well as their greater virulence in mouse models of human infection^{21,22}. As an initial test of the hypothesis that ZIKV underwent a fitness decline upon its introduction into Africa, followed by fitness restoration, we assessed the fitness of African, Asian, and American ZIKV strains. We used competition fitness assays, where ZIKV strains differing by as little as one amino acid were generated from cDNA clones and mixed in a roughly 1:1 ratio based on Vero cell plaque-forming units (PFU). This inoculum mixture was then used to initiate infections. Following appropriate incubation to allow the two strains to compete for replication and, in mosquitoes, dissemination to the salivary glands, the virus mixture was again quantified by Sanger sequencing of RT-PCR amplicons to estimate the ratios of mixed nucleotides. The relative fitness value w was analyzed according to w=(f0/i0), where f0 is the final ratio of one competitor following infection, and i0 is the initial ratio in the inoculum or bloodmeal mixture; this ratio reflects relative fitness advantage of one competitor over the other (Fig. 1a). This approach has major advantages over performing individual strain infections with numerous host replicates; each competition is internally controlled, eliminating host-to-host variation that can reduce the power of experiments, and the virus strain ratios can be assayed with more precision than individual virus titers. Thus, competition assays have been used for many studies of microbial fitness²³, including arboviruses²⁴⁻²⁶. To confirm that our targeted 1:1 ratios were similar in mosquito experimental systems, where RNA:infectious virus ratios could differ based on different levels of mutant infectivity compared to those for Vero cells, we also determined infectious titers of the wt and mutants on C6/36 mosquito cells, and compared them with the Vero PFU and Sanger-derived ratios. These ratios were nearly indistinguishable with no significant differences (Extended data Fig. 3), indicating that frequency-based differences in fitness based on potential mutation effects on initial infectivity were not a concern.

Fitness of major Zika virus lineages, Initially, we performed competition assays with wildtype (wt) ZIKV strains representing the African, Asian pre-epidemic, and American epidemic genotypes (Fig. 1). The highly susceptible Rockefeller strain of A. aegypti mosquitoes was used along with interferon type I receptor-deficient A129 mice, models for human infection^{27,28}. Following infection with an approximately 1:1 mixture of the African Dakar 41525 and Asian Cambodian FSS13025 strains, or an equivalent mixture of FSS130025 and the Puerto Rican RRVABC59 strain, samples from mosquito bodies and mice were evaluated for changes in the initial ratio, indications of competitive fitness (Fig. 1a). Following oral mosquito infection and incubation, the African strain consistently and significantly won the competition with the Asian strain, and the Puerto Rican strain consistently outcompeted the Asian strain (Fig. 1b). These findings were reproducible when additional African, Asian and American ZIKV strains, as well as a Dominican Republic strain of A. aegypti with a low level of colonization (F6), were used (Extended Data Fig. 4). After three days of competition following subcutaneous infection of A129 mice to mimic human infection, the same relative fitness results were obtained when viremic blood (predicted to be the vertebrate host compartment where major selection for efficient mosquito infection occurs) was analyzed (Fig. 1c). Analysis of individual organ samples from mice on day 8 (after viremia had subsided to eliminate this confounding source of virus in organs) showed that these competition outcomes were consistent for replication throughout the mouse (Extended Data Fig. 5). Mosquitoes that fed upon these viremic mice were also evaluated after extrinsic incubation, with the same results of greater fitness of the African versus Asian ZIKV strain, and greater fitness of the American versus Asian strain (Fig. 1d).

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Next, primary human cells believed to be involved in seeding viremia were used for competition assays: fibroblasts and keratinocytes^{29,30}. The mixed ZIKVs were inoculated into cells and the culture supernatant were harvested, RT-PCR-amplified and Sanger sequenced 3 days post-infection (Extended Data Fig. 6a). The results showed that the African strain and post-epidemic American strains always out-competed the pre-epidemic Asian strain in both cell types (Extended Data Fig. 6b, c). In all these experimental models, the changes in competitor ratios appeared to be greater between the African versus Asian strains compared to the Asian versus American strains. Overall, these data supported the hypothesis that ZIKV underwent a major loss of fitness upon its introduction into Asia, followed by a partial restoration of fitness upon introduction into the Americas. We note that microcephaly and other central nervous system involvement is rare and not known to play a major role in generating viremia, so there should not be major selection for CNS tropism to enhance viremia and subsequent transmission by mosquitoes. Therefore, microcephaly may represent a chance pathogenic outcome that did not result from positive selection. Also, sexual ZIKV transmission is believed to play a minor role compared to vector-borne transmission.

Based on the evidence discussed above, we developed a revised adaptation hypothesis, depicted in Fig. 2a, that the 4 initial amino acid substitutions (and possibly others not undergoing reversion) represent founder effects that reduced the fitness of ZIKV for transmission in the epidemic cycle by reducing *A. aegypti* transmissibility and/or human viremia levels. The reversion of these 4 mutations, and possibly other adaptive mutations, then partially restored ZIKV fitness, allowing for efficient spread to the Americas and major epidemics.

Fitness effect of reversion mutations. To test this hypothesis that the 4 revertant mutations restored ZIKV fitness, we used *A. aegypti* mosquitoes and A129 mice in more detailed studies of the combinations of four amino acid substitutions that underwent reversion. For mosquito infections, we assayed bodies to evaluate infection and initial replication, legs to sample the hemolymph, which contains virus that has disseminated from the digestive tract (midgut) into the hemocoel for access to the salivary glands, and saliva collected in vitro to assess the virus population available for transmission (Fig. 2b). Competition between the African ZIKV strain and a mutant containing the four initial amino acid substitutions resulted in a consistent advantage for the African strain in all mosquito samples (Fig. 2c), as did infection of

A129 mice as sampled in major organs and blood (Fig. 2d, Extended Data Fig. 7a). In contrast, when the four reversion substitutions were placed into the pre-epidemic Asian strain to recapitulate ZIKV evolution just before introduction into the Americas, these substitutions resulted in a major fitness gain (Fig. 2e, f, Extended Data Fig. 7b). When mosquitoes were fed upon the viremic mice, the same outcome was observed (Fig. 2g, h). These results indicate that the four substitutions that directly reverted prior to the introduction of ZIKV into the Americas were major components of the fitness differences among African, Asian and American virus strains. At the same time, similar competition assays were performed in human fibroblasts and keratinocytes. The ZIKV strains containing these four initially substituted amino acids always won the competition in these human primary cells (Extended Data Fig. 8).

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To assess the roles of the individual amino acid substitutions in the overall fitness differences among ZIKV strains, we tested them individually using the same experimental systems. Placed individually into the African strain, all four initial mutations reduced overall fitness for infection, dissemination (two mutants showed no fitness effect in this compartment) and transmission by mosquitoes, although only the C-A106T and NS1-V188A results were significant (Fig. 3a-c). The same trends were observed in mosquito bodies, legs and saliva (Fig. 3a-c). When the reversions were placed into the Asian strain, only the NS1-A188V mutant produced a significant increase in ZIKV infection in the mosquito bodies, legs and saliva, indicating this mutation site may have a dominant role during the competition (Fig. 3d-f). To confirm this, the Dakar NS1-V188A and FSS NS1-A188V mutants were mixed with corresponding *wt* strains and inoculated into A129 mice. The ZIKVs containing valine in the 188th NS1 position exhibited a fitness advantage in both mice (Fig. 3g, h, Extended Data Fig. 9) and mosquitoes (Fig. 3i, j). the slight asymmetry in the fitness effects between some of the initial 4 mutations in the African strain and the reversions in the Cambodian pre-epidemic strain could reflect epistatic interactions that differ between the two virus strains.

Finally, the overall fitness of the sets of four mutations or reversions were tested over a complete experimental transmission cycle. To initiate the cycle, mosquitoes were inoculated intrathoracically (to ensure uniform infection) with mixtures of the African ZIKV strain and the four initial mutations, or the Asian strain with the four reversions (Fig. 4a). Following six days of extrinsic incubation (mosquitoes become infectious more rapidly after intrathoracic than oral infection), these mosquitoes were exposed to A129 mice for oral transmission. Three days later during peak viremia, naïve mosquitoes were allowed to feed on the mice, then these mosquitoes were assayed after 14 days of extrinsic incubation as in the above experiments. In salivary glands following incubation after mosquito inoculation, the initial four mutations decreased the fitness of the African ZIKV strain (Fig. 4b), and the four reversions increased the fitness of the Asian strain (Fig. 4c) as observed after oral infection (Fig. 2c, e). These fitness effects continued during infection of mouse blood and organs (Fig. 4d, e, Extended Data Fig. 10) and also in mosquitoes fed orally on these mice during viremia (Fig 4f, g). Overall, these results demonstrate that the four initial mutations reduced the fitness of the African ZIKV strain through a complete transmission cycle, while the four reversions at least partially restored fitness.

History of ZIKV fitness in Asia. Overall, our results strongly support the hypothesis that ZIKV underwent fitness declines upon its introduction into Asia many decades ago, and/or during its initial stages of circulation there when a lack of recognized outbreaks suggests limited urban circulation. An alternative hypothesis that would not rely on drift is that ZIKV host range changed upon its introduction into Asia, followed by another change (or reversion in host usage) just before introduction into the Americas; both of these changes could have resulted in positive selection that happened to involve the same four amino acids, resulting in reversion. The simplest form of this hypothesis would be that an ancestral sylvatic cycle involving different vectors and vertebrate hosts from those in Africa was the initial form of Asian transmission, and the urban cycle only developed there relatively recently accompanied by the reversions.

However, there is no evidence of enzootic ZIKV transmission in Asia, and the 1966 ZIKV isolation from *A. aegypti* mosquitoes in Malaysia suggests that human-amplified transmission has been ongoing in Asia for many decades. A simpler explanation is that this urban cycle was inefficient due to founder effects and drift, before reversion of these mutations increased transmission efficiency, facilitating spread to the South Pacific and the Americas. This same pattern of fitness loss accompanied CHIKV introduction into Asia about a century ago, when a major deletion in the 3' untranslated genome region occurred. This fitness of this Asian lineage remains incompletely restored to this day³¹. However, the inability of these four ZIKV reversion mutations to generate fitness comparable to that of African strains suggests that additional mutations, possibly also the result of founder effects or drift but without direct reversions, also limited transmission and spread in Asia. Additional mutations that occurred during circulation in Asia must be tested to expand our results to a comprehensive understanding of ZIKV evolution, and understanding the mechanisms of fitness effects of the four substitutions we studied require further study.

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Parallels with CHIKV. Our results as well as the evolutionary history of ZIKV have remarkable parallels to those of CHIKV, which also evolved in Africa and spread many decades ago to Asia before its recent introduction into the Americas¹². Previous studies of CHIKV indicate that the virus arrived in Asia many decades ago with debilitating mutations in its 3' untranslated genome region (3'UTR), with fitness only partially restored over many decades through point mutations and a duplication³¹. Another mutation in the E1 glycoprotein, also a probable founder effect, prevented for many decades CHIKV adaptation for efficient transmission by A. albopictus mosquitoes in Asia, their native territory³². Then, just before or upon its introduction into the Americas in 2013, a duplication in the 3'UTR also improved CHIKV fitness³³. However, unlike ZIKV, another lineage of CHIKV has also undergone a series of adaptive mutations to enhance its ability to use A. albopictus for transmission with no evidence of founder effects, and the relative fitness advantages of these CHIKV mutations far exceed those of the four ZIKV mutations that we studied; for example, individual A. albopictus-adaptive CHIKV mutations showed relative fitness values of 5-40, while the combination of 4 ZIKV mutations we studies had a lower overall fitness effect than any of these individual CHIKV mutations (Extended Data Table 1).

The similarities in the presumed roles of founder effects in the evolution and emergence potential of ZIKV and CHIKV suggest that drift has been understudied as a factor in the emergence of arboviral and other RNA viral diseases, and that the stochastic nature of founder effects may limit our ability to ultimately predict the emergence of new viral diseases.

Finally, the high fitness of the African ZIKV strains in all experimental systems we utilized raises important questions about the risk of outbreaks and severe disease on that continent. The greater transmissibility of these African strains, even compared to American strains¹⁹, raises the question of why outbreaks have never been detected in Africa aside from one in Angola caused by a strain imported from the Americas³⁴. Limited African surveillance and the widespread presence of other infections that typically lead to Zika misdiagnosis in the absence of laboratory diagnostics could be responsible. Herd immunity in Africa, which could also play a role in limiting outbreaks as well as the incidence of congenital Zika syndrome, has been reported to range from 1-52%³⁵. However, some studies have used methods other than neutralization that could be highly cross-reactive with other endemic flaviviruses. We also cannot rule out the possibility that our models for human infection are not suitable to determine ZIKV fitness for human amplification. Additional surveillance and work with human cells and possibly nonhuman primate models are needed to further explore this possibility and to better understand the lack of Zika outbreaks in Africa caused by African lineage strains.

In addition to the fitness for transmission that we examined, the history of ZIKV's ability to cause GBS and CZS requires further study. It is possible that enhanced viremia just before spread to the Americas, suggested by our model systems involving human cells and A129 mice,

could at least partially explain the emergence of these severe disease outcomes if viremia magnitude is correlated with infection of the placenta. However, better surveillance in Africa is ultimately needed to determine if American or recent Asian strains are unique in this pathogenic potential.

Methods.

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Mouse and mosquito studies were performed in accordance with the guidance for the Care and Use of Laboratory Animals of the University of Texas Medical Branch (UTMB). The protocol (protocol number 1708051 for mouse) were approved by the Institutional Animal Care and Use Committee (IACUC) at UTMB. All the mouse manipulations were performed under anesthesia by isoflurane.

Mice, mosquitoes, cells and viruses

A129 mice, which are deficient in type I interferon receptors, were bred and maintained in animal biosafety level 2 (ABSL2) facilities at UTMB. Sex-matched mice, randomly selected and 6-to-8-week-old, were used. The Aedes aegypti Rockefeller strain and A. aegypti Dominican Republic strain (F6) were maintained in an incubator at 28°C and 80% humidity. Vero cells (CCL81) and C6/36 cells for virus rescue, proliferation and titration were purchased from the American Type Culture Collection (Bethesda, MD, USA) and maintained in Dulbecco's modified Eagle's medium (Gibco, Waltham, MA, USA) with 10% heat-inactivated fetal bovine serum (Atlanta Biologicals, Flowery Branch, GA, USA), and supplemented with 1% Tryptose phosphate broth (Gibco, Waltham, MA, USA) for C6/36 cells. The human primary dermal fibroblast cells and human epidermal keratinocyte cells were purchased from Lonza (Walkersville, MD, USA) and were maintained in FGM-2 BulletKit (Lonza) and KGM-Gold BulletKit (Lonza) media, respectively. All cells were verified and tested negative for mycoplasma. The parental African lineage ZIKV Dakar 41662 strain (KU955592), and 41671 stain (KU955595), and Asian lineage ZIKV Dominican Republic R114916 (KX766028), and Honduras HN-ME 59 stains were obtained from the World Reference Center for Emerging Viruses and Arboviruses at UTMB. The ZIKV DK-WT (Dakar wild-type 41525 strain, KU955591), DK-A106T (capsid 106th amino-acid), DK-A1V (pre-membrane 1st amino-acid), DK-V188A (nonstructural protein 1 188th amino-acid), DK-V872M (nonstructural protein 5 872nd amino-acid) were constructed using standard site-directed mutagenesis of cDNA clones and rescued on Vero cells using electroporation of transcribed RNA. The FSS-WT (FSS13025 wild-type strain, KU955593), FSS-T106A, FSS-V1A, FSS-A188V and FSS-M872V, as well as reversion mutants placed in the ZIKV Dakar strain, were constructed in the same manner, along with the DK-4M and FSS-4M that combined all four mutations on the matched backbone.

315 Competition assay

To better understand the relative fitness of different ZIKV strains and mutants, we conducted competition assays between two virus strains or a wild-type and mutant strain in various models. Initial 1:1 virus:mutant mixtures were made based on PFU titers determined in Vero cells, and virus ratios used to calculate fitness were determined by Sanger sequencing of RT-PCR amplicons. The PFU:genome ratio was consistent among all of our ZIKV strains and constructs, as estimated from the inocula and bloodmeals by comparing Vero cell PFU to genome copies as estimated from real-time RT-qPCR standard curves. To confirm that ratios were also similar in mosquito experimental systems, where RNA:infectious virus ratios could differ based on different levels of infectivity compared to Vero cells, we also determined infectious titers of the wt and mutants on C6/36 mosquito cells, and compared then with the Vero-infectious and Sanger-derived ratios.

Mosquitoes. The *Aedes aegypti* Rockefeller strain and *A. aegypti* Dominican Republic strain (F6) were used for the following studies. Triplicate mixtures of viruses were diluted to 6 log₁₀ PFU/ml mixed at a 1:1 ratio based on Vero PFU titers. Virus mixtures were subsequently mixed at a ratio of 1:1 with PBS-washed sheep blood cells (Hemostat laboratories, Galveston, TX,

USA). Aliquots of blood meals were collected to verify the initial virus proportions in the inoculum. Following a 14-day extrinsic incubation, saliva expectorated into capillary tubes, legs representing virus disseminated into the hemocoel, or remaining mosquito bodies were collected separately in a 2ml Eppendorf Safe Lock tube with 250µl DMEM (2% FBS, 1% Sodium Pyruvate) with 25 mg/ml Amphotericin B and a stainless steel bead (Qiagen, Hilden, Germany). Samples were stored at -80°C.

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- A129 mice. Sex-matched A129 mice (6-8 weeks old) were used for ZIKV competition assays.

 The A129 mouse is defective for the interferon type I receptor and susceptible to ZIKV infection in vivo with high titered viremia. Mixtures of ZIKV strains (total 1×10⁴ PFU/mouse) were inoculated intradermally to simulate a mosquito bite. Titers of viruses before mixing were plaque assayed to verify proper concentrations. Also, an aliquot of the inoculum was reserved for estimating the initial ratio of viruses. Mice were bled retro-orbitally on day 3 post infection representing the peak. Serum was collected from blood via centrifugation at 2,000 g for 5 minutes and stored at -80°C until analysis. Eight days post infection, all infected mice were euthanized and necropsied and major organs collected in a 2ml Eppendorf Safe Lock tube with 500µl DMEM (2% FBS, 1% Sodium Pyruvate) and a stainless-steel bead. Samples were stored at -80°C.
 - **Human primary cells.** The human fibroblasts and keratinocytes were purchased and qualified by ATCC. Cells were seeded into 12-well plates 1 day before infection to allow them reach 90% confluence. Mixtures of ZIKVs (1 PFU/cell MOI) were added into the cells and incubated at 37°C for 2 hours. Titers of viruses before mixing were plaque assayed to verify proper concentrations. Also, an aliquot of the inoculum was reserved for estimating the initial ratio of the viruses. After incubation, the cells were washed 3 times with 1ml of PBS. The infected cells were cultured, and the supernatant were collected from days 1-5. Samples were stored at -80°C for further analysis.
- Sample preparation and Nucleic acid extraction. Aliquots of mosquito lysates, mouse serum, mouse organ lysates and human primary cell supernatant samples were prepared in RNeasy Mini Kit (Qiagen) lysis buffer RLT (400µl), and nucleic acids were subsequently extracted according to the manufacturer's protocol. After extraction, a portion of the RNA was immediately applied to a one step RT-PCR assay and the remaining material was archived at -80°C.
 - **Quantitative real-time RT-PCR assays**. Prior to Sanger sequencing analysis, virus-positive samples were identified by quantitative real-time RT-PCR, which was performed using a QuantiTect Probe RT-PCR Kit (Qiagen) on the LightCycler 480 system (Roche, Rotkreuz, Switzerland) following the manufacturer's protocol. The primers and probes are listed in Extended Data Table 2. The absolute quantification of ZIKV RNA was determined using a standard curve with in vitro-transcribed, full-length ZIKV RNAs.
 - Reverse transcriptase PCR. 400-500 bp of RT-PCR product were synthesized and amplified from extracted RNA using a SuperScript™ III One-Step RT-PCR kit (Invitrogen, Carlsbad, CA, USA). 20µl reactions were assembled in PCR 8-tube strips through the addition of 10µl 2X reaction mix, 0.4µl SuperScript™ III RT/Platinum™ Taq Mix, 0.8µl 10 µM specific forward primer, 0.8µl 10 µM specific reverse primer (see Extended Data Table 2), 4µl of extracted RNA and 6µl Rnase-free water. RT was completed using the following protocol: 1) 55°C, 30 min; 94°C, 2 min; 2) 94°C, 15s; 60°C, 30s; 68°C, 1min; 40 cycles; 3) 68°C, 5 min; 4) indefinite hold at 4°C. 2µl of generated PCR products were loaded to 1% DNA agarose gel to verify the size. The left samples were purified by a QIAquick PCR Purification kit (Qiagen) according to the manufacturer's protocol.

Sanger sequencing and electropherogram peak height analysis. Sequences of the purified RT-PCR products were generated using a BigDye Terminator v3.1 cycle sequencing kit (Applied Biosystems, Austin, TX, USA). The products of the sequencing reactions were then purified using a 96-well plate format (EdgeBio, San Jose, CA, USA), and then analyzed on a 3500 Genetic Analyzer (Applied Biosystems). The peak electropherogram height representing each mutation site and proportion of each competitor was analyzed using the QSVanalyser program³⁷.

intrathoracic inoculation of mosquitoes.

The intrathoracic microinjection of ZIKV into mosquitoes has been described previously³⁸. Female mosquitoes were anaesthetized on a cold tray (0°C). Then, a defined titer (1 pfu in 100 nl total inoculum volume) of mixed ZIKV strains were inoculation into the mosquito thorax with a Nanoject III (Drummond, Pennsylvania, USA). Six days later, the infected mosquitoes were allowed to feed on A129 mice to transmit ZIKV.

Membrane blood feeding.

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- Seven-day-old female mosquitoes were placed into mesh-covered cartons and provided cotton balls containing 10% sucrose. Complement-inactivated sheep blood was mixed with different ZIKV combinations for mosquito feeding via a Hemotek system membrane feeder (5W1, Hemotek limited, Lancashire, UK). The total viral titers of the two competing ZIKVs in the blood meals were 1x10⁶ PFU/ml. Fully engorged mosquitoes were incubated for 14 days until harvest.
 At that time, live mosquitoes were killed by freezing and homogenized individually (TissueLyser II, Qiagen) for RNA isolation and real-time qPCR detection. The infected mosquito RNAs were RT-PCR amplified, followed by Sanger sequencing.
 - Mosquito feeding on infected mice.
- Ten female mosquitoes were placed into mesh-covered cartons for blood feeding following sugar-starvation for 24 hrs. ZIKV-infected A129 mice were anaesthetized with ketamine and placed on the top of the cartons for 20 min of feeding in the dark. Then, fully engorged mosquitoes were transferred to new containers and incubated for 14 days prior to RT-PCR, followed by Sanger sequencing of amplicons.

Plaque assay.

The plaque assay was performed on Vero cells following procedures described previously³⁹. In brief, Vero cells were seeded into 12-well plates 12 to 16 hours prior to the plaque assay. 25 μ l of each sample were used for 10-fold serial dilutions. For each dilution, 200 μ l were added to 12-well plates with 90% confluent Vero cells. The cells were incubated at 37 °C with 5% CO₂ for 1h with gentle rocking every 15 min. After that, 1 ml of overlay (DMEM, 2% FBS, containing 0.8% methyl cellulose with 1% antibiotics) was added onto each well. The plates were cultured at 37 °C with 5% CO₂ for 4 days until clear plaques formed. The plates were fixed in 4% formaldehyde solution for 2 hours and stained with 1% crystal violet.

Focus-forming assay.

The FFA was performed on C6/36 mosquito cells as described previously¹⁹. In brief, C6/36 cells were seeded into 12-well plates 12-16 h prior to the assay. A 25 μl volume of each sample was used for 10-fold serial dilutions. For each dilution, 200 μl were added to 12-well plates with 90% confluent cells. The cells were incubated at 30°C with 5% CO₂ for 1h with gentle rocking every

15 min. After a 5-day incubation, plates were fixed with methanol / acetone (1:1), washed by PBS, and blocked with 2% FBS/PBS before overnight incubation with mouse anti–ZIKV antibody (4G2). Plates were washed and incubated with goat anti-mouse secondary antibody conjugated to horseradish peroxidase (KPL, Gaithersburg, MD, USA), then washed and developed with aminoethyl carbazole solution (Enzo Diagnostics, Farmingdale, NY, USA) prepared according to the manufacturer's protocol for detection of infection foci.

Construction of ZIKV mutant infectious clones.

The ZIKV FSS13025 full-length cDNA infectious clone pFLZIKV-FSS³⁹ (referred as FSS-WT in text) was used as the backbone for engineering FSS-T106A, FSS-V1A, FSS-A188V, FSS-M872V and FSS-4M mutants. The ZIKV Dakar 41525 full-length cDNA infectious clone pFLZIKV-DK (referred as DK-WT in text) was used as the backbone for engineering DK-A106T, DK-A1V, DK-V188A, DK-V872M and DK-4M mutants. Standard overlapping PCR was performed to amplify the cDNA fragment between unique restriction enzyme sites that contained the corresponding mutations. Afterward, the fragments were cloned into pFLZIKV-FSS or pFLZIKV-DK plasmids and propagated in E. coli strain Top 10 (ThermoFisher Scientific, Framingham, MA, USA). All restriction enzymes were purchased from New England BioLabs (Massachusetts, USA). The plasmids were validated by restriction enzyme digestion and Sanger DNA sequencing. All primers were synthesized from Integrated DNA Technologies and are available upon request.

RNA transcription, electroporation and virus recovery.

The full-length cDNA clone plasmids of all ZIKVs were linearized with restriction enzyme Clal prior to RNA transcription. The linearized DNAs were purified by phenol-chloroform extraction and ethanol precipitation. RNA transcripts of ZIKVs were synthesized using the T7 mMessage mMachine kit (Ambion, California, USA) *in vitro*. The quantity and quality of RNAs were verified by Spectrophotometry (DS-11, DeNovix, Delaware, USA) and Agarose gel electrophoresis. For RNA transfection, 10 µg of transcribed RNA were electroporated into 8×10⁶ Vero cells using the Gene Pulser Xcell™ Electroporation Systems (Bio-rad, California, USA) under as described previously³⁹. The viral culture supernatant was harvested when obvious cytopathic effects (CPE) of the transfected Vero cells were observed, and the titers of all ZIKVs were measured on Vero cells using plaque assays³⁹.

Statistics.

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For virus competition experiments in mosquitoes, A129 mice and human primary cells, relative replicative fitness values for viruses in each sample were analyzed according to w=(f0/i0), where i0 is the initial ratio of one competitor and f0 is the final ratio after competition. Sanger sequencing (initial timepoint T0) counts for each virus strain being compared (for example, wild-type versus mutant strains) were based upon average counts over several replicate samples of inocula or bloodmeals per experiment (effectively averaging the replicates), and post-infection (timepoint T1) counts were taken from samples of individual subjects, such that for each strain the ratio between counts at T0 and T1 (T1/T0) reflects a measure of a sample from a single mosquito, mouse or cell culture well in a specific experiment for that strain. Typically, multiple experiments were performed, so that f0/i0 was clustered by experiment. To model f0/i0, the ratio T0/T1 was found separately for each subject in each strain group, log (base-10) transformed to an improved approximation of normality, and modeled by analysis of variance with relation to group, adjusting by experiment to control for clustering within experiment. Specifically, the model was of the form Log10_CountT1overCountT0 ~ Experiment + Group. Fitness ratios

between the two groups [the model's estimate of w=(f0/i0)] were assessed per the coefficient of the model's Group term, which was transformed to the original scale as 10^coefficient. This modeling approach compensates for any correlation due to clustering within experiment similarly to that of corresponding mixed effect models, and is effective since the number of experiments was small.

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Statistical analyses were performed using R statistical software (R Core Team, 2019, version 3.6.1). In all statistical tests, two-sided alpha=.05. Catseye plots⁴⁰, which illustrate the normal distribution of the model-adjusted means, were produced using the "catseyes" package⁴¹.

490 Data availability.

Extended Data and source data for generating main figures are available in the online version of the paper. Any other information is available upon request.

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Author Contributions

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S.C.W., J.L., Y.L., C.S., and P.-Y.S. designed the experiments and/or wrote the manuscript; J.L. and Y.L. performed the majority of the experiments and analyzed the data; S.C., B.T.D.N., and P.-Y.S. developed the reverse genetic system for ZIKV strains; R.M. and N.V. provided the mosquito colonies; R.M. assisted in mosquito collection; S.L.H. assisted in designing the competition assay; G.H.R. and S.R.A provide the A129 mice colonies; C.R.A. assisted with statistics analyses; K.P. provided ZIKV parental strains and contributed experimental suggestions. All authors reviewed, critiqued and provided comments on the text.

510 Competing Interests statement

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The authors declare that they have no competing financial interests. Correspondence and requests for materials should be addressed to S.C.W. (sweaver@utmb.edu) or P.-Y.S. (peshi@utmb.edu).

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Figure legends and Table.

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Figure 1. African lineage ZIKV and post-epidemic Asian lineage ZIKV had the fitness advantage versus pre-epidemic Asian lineage ZIKVs in both A129 mice and mosquitoes

- a, Experimental design of competition fitness assays. African lineage ZIKV and post-epidemic 630 Asian lineage ZIKV were mixed with pre-epidemic Asian lineage ZIKV at an approximate ratio of 1:1 respectively. The initial 1:1 ratios of mixed virus were confirmed by RT-PCR, followed by Sanger sequencing of replicons and polymorphic nucleotide peaks analysis. The ZIKV mixture was engorged by mosquitos through membrane blood feeding or biting an infected A129 mouse at the viremia peak 3 days post-infection. The mosquitoes were sacrificed after 14 days of 635 incubation, the ZIKV population amplified by RT-PCR, and the amplicon Sanger-sequenced. The blood and organs of infected mice were collected 3- or 8-days post-infection, respectively. All the mosquitoes and mouse specimens were subjected to RT-PCR amplification and Sanger sequencing to compare the ZIKV strain ratio after competition, and each point represents a single mosquito or mouse sample. Relative fitness values were compared among multiple 640 competition replicates to determine if fitness of the competitors differed. b, Fitness comparison between African (Dakar 41525), Asian pre-epidemic (FSS13025) and American (PRVABC59) ZIKV strains in A. aegypti. The relative replicative fitness value w was analyzed according to w=(f0/i0), where i0 is the initial ratio of the Dakar 41525 or PRVABC59 competitor to the FSS13025 strain, determined from Sanger sequence electropherogram peaks, and f0 is the 645 final ratio. c, Fitness comparison of African (Dakar 41525), Asian pre-epidemic (FSS13025) and American (PRVABC59) ZIKV strains in mouse blood, using the same method as for panel b. d, Fitness comparison of African (Dakar 41525), Asian pre-epidemic (FSS13025) and American (PRVABC59) ZIKV strains in mosquitoes after feeding on the viremic mice shown in panel c, and extrinsic incubation, using the same method as for panel b. **b-d**. The distribution of the model-adjusted means is illustrated by catseve plots with shaded +/- standard errors overlaid by 650 scatterplots of individual relative fitness values; scatterplots have been randomly jittered horizontally for clarity, and are shown on the log (base-10) scale such that comparisons are against a null value of 1.
- Figure 2. Four amino acids determine much of the fitness difference between different ZIKV strains in both mosquitoes and A129 mice.
 - a. The hypothetical diagram of fitness changes following the spread of ZIKV from Africa to Asia and the Americas, including strains used for fitness assays. b, Schematic representation of competition fitness assay in mosquito bodies, legs and saliva. c, e, Fitness comparison between African and Asian lineage, and the four amino-acid substitution mutants (DK-4M and FSS-4M) in mosquito bodies, legs and saliva. The mosquitoes acquired the ZIKV mixture through membrane blood feeding and were assayed at 14 days post-infection. The body (carcass), legs and saliva were collected from each individual mosquito and subjected to RT-PCR amplification and Sanger sequencing to determine relative fitness values; each point represents a single mosquito or mouse sample. d, f, Competition between DK-WT and DK-4M(d), FSS-WT and FSS-4M (f) in mouse blood collected 3 days after infection. g, h, Fitness of DK-4M (g) and FSS-4M (h) in mosquitoes after oral infection from viremic mice. The mosquitoes acquired the virus through biting a viremic mouse 3 days after infection and were tested after 14 days of extrinsic incubation. c-h, The distribution of the model-adjusted means is illustrated by catseye plots with shaded +/- standard error bars overlaid by scatterplots of subject measures; scatterplots have been randomly jittered horizontally for clarity, and are shown on the log (base-10) scale such that comparisons are against a null value of 1.

Figure 3. The fitness comparison of 4 individual mutant viruses against wild-type ZIKV strains in mosquitoes and A129 mice.

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a-c, The relative fitness of individual mutants in the Dakar strain, tested in mosquito bodies (**a**), legs (**b**) and saliva (**c**). **d-f**, The fitness comparison of individual reversion mutants in the FSS13025 strain, tested in mosquito bodies (**d**), legs (**e**) and saliva (**f**). **g,h**. Each point represents a single mosquito sample The fitness comparison of DK-V188A (**g**) and FSS-A188V (**h**) versus wild-type strains in mouse blood. **i**, **j**, The fitness comparison of DK-V188A (**i**) and FSS-A188V (**j**) in mosquitoes after oral infection from viremic mice. **a-j**, The distribution of the model-adjusted means is illustrated by catseye plots with shaded +/- standard error overlaid by scatterplots of subject measures; scatterplots have been randomly jittered horizontally for clarity, and are shown on the log (base-10) scale such that comparisons are against a null value of 1.

Figure 4. Fitness comparison of DK-4M and FSS-4M against wild-type ZIKV strains during a mosquito-host-mosquito transmission cycle.

a, Schematic representation of the study design. 10 pfu of ZIKV WT and mutant viruses were 690 intrathoracically injected into mosquitoes. After 6 days of extrinsic incubation, the infected mosquitoes fed on a A129 mice to transmit virus. Naïve mosquitoes were then employed to bite the infected A129 mice 3 days post-mouse infection. After an additional 14 days, the mosquitoes were harvested. All the mosquitoes and mice samples collected during the transmission cycle were analyzed by RT-PCR and Sanger sequencing. Each point represents a 695 single mosquito or mouse sample. b, c, Fitness comparison of DK-4M (b) and FSS-4M (c) against wild-type viruses in mosquito salivary glands 6 days post-intrathoracic injection. d, e, Fitness of DK-4M (d) and FSS-4M (e) versus wild-type viruses in mouse blood. f, g, The fitness comparison of DK-4M (f) and FSS-4M(g) versus wild-type viruses in mosquitoes after oral infection from viremic mice. b-a. The distribution of the model-adjusted means is illustrated by 700 catseye plots with shaded +/- standard error overlaid by scatterplots of subject measures; scatterplots have been randomly jittered horizontally for clarity, and are shown on the log (base-10) scale such that comparisons are against a null value of 1.

Extended Data Figures and Tables

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Extended Data Fig. 1. Phylogenetic analysis of representative ZIKV strains.

a, Phylogenetic tree of representative Zika virus (ZIKV) strains based on 73 complete open reading frame sequences. The evolutionary distances were computed using the Maximum Composite Likelihood method. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap tests were shown next to the branches (1000 replicates). Evolutionary analyses were performed in MEGA X, by using the Maximum Likelihood method. **b**, Four reversion directly reverting mutations compared among representative African, Asian and American lineage strains.

Extended Data Fig. 2. The trace history of four directly reverting amino acids in the evolutionary tree of Zika virus.

Trace history of four ZIKV amino acids performed in Mesquite (Version 3.61, http://www.mesquiteproject.org). These include: a. Amino acid 106 substitution of the capsid protein; b. Amino acid 1 of the pre-membrane (prM) protein; c. amino acid 188 of the nonstructural protein 1 (NS1), and; d. amino acid 872 of the nonstructural protein 5 (NS5) based on 73 complete zika virus open reading frame sequences.

Extended Data Fig. 3. ZIKV WT and 4M mutant strain had a similar initial ratio when titrated on mammalian or insect cells, or estimated by the Sanger sequencing method.

The initial ratio of DK-WT/DK-4M (a) and FSS-WT/FSS-4M (b) were calculated by the virus titer determined on Vero and C6/C36 cells or by the electropherogram peak height based on Sanger sequencing result. The titers of the WT and mutant viruses were determined by FFA (Focusforming assay) on the cells before they were mixed. The virus mixture of WT and mutant viruses were determined by Sanger sequencing of RT-PCR amplicons.

Extended Data Fig. 4. Fitness comparisons with additional Zika virus strains Dominican Republic *Aedes aegypti* mosquitoes.

a, The fitness of 2 African ZIKV strains (DK 41662) and (DK 41671) in competition with the Asian FSS13025 strain in mosquitoes. b, The fitness of 2 American ZIKV strains (R114916) and (HN-MF59) in competition with the FSS13025 Asian strain in mosquitoes. c. The fitness of ZIKV African strain Dakar 41525, Asian pre-epidemic strain FSS13025 strain and American strain PRVABC59 in a low generation (F6) colonized *A. aegypti* strain from the Dominican Republic. Each point represents a single mosquito or mouse sample a-c, The distribution of the model-adjusted means is illustrated by catseye plots with shaded +/- standard error overlaid by scatterplots of subject measures; scatterplots have been randomly jittered horizontally for clarity, and are shown on the log (base-10) scale such that comparisons are against a null value of 1.

Extended Data Fig. 5. Fitness comparisons of African and post-epidemic against a preepidemic ZIKV strains in different organs of A129 mice.

a, Fitness comparison between African (Dakar 41525) and Asian pre-epidemic (FSS13025)
 ZIKV strains in different organs of A129 mice after 8 days of infection, when viremia had ended.
 b, The fitness comparison between Asian pre-epidemic (FSS13025) and American

(PRVABC59) ZIKV strains in different organs of A129 mice after 8 days of infection, when viremia had ended. Each point represents a single mouse sample $\bf a$, $\bf b$, The distribution of the model-adjusted means is illustrated by catseye plots with shaded +/- standard error overlaid by scatterplots of subject measures; scatterplots have been randomly jittered horizontally for clarity, and are shown on the log (base-10) scale such that comparisons are against a null value of 1. * P < 0.05, ** P < 0.01, *** P < 0.001.

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Extended Data Fig. 6. Fitness comparison of African and post-epidemic against a preepidemic ZIKV stain in human primary cells.

a, Schematic representation of the study design. The mixed ZIKVs were inoculated into human fibroblast and keratinocyte cells. The RNAs in the culture supernatant were isolated, amplified by RT-PCR, Sanger-sequenced 3 days post infection. **b**, **c**, The fitness of ZIKV African strain (Dakar 41525), Asian pre-epidemic (FSS13025) strain and American strain (PRVABC59) in human primary fibroblast (**b**) and keratinocyte (**c**) cells. Each point represents a single culture sample The distribution of the model-adjusted means is illustrated by catseye plots with shaded +/- standard error overlaid by scatterplots of subject measures; scatterplots have been randomly jittered horizontally for clarity, and are shown on the log (base-10) scale such that comparisons are against a null value of 1.

Extended Data Fig. 7. Fitness comparisons of Dakar-4 amino-acid mutant and FSS13025-4 amino-acid mutant against wild-type viruses in different organs of A129 mice.

a, Fitness comparison between Dakar 4-amino acid mutant (DK-4M) and the wild-type strain in different organs of A129 mice. b, The fitness comparison between FSS 4 amino-acid mutant (FSS-4M) and wild-type strain in different organs of A129 mice. Each point represents a single mouse sample. a, b, The distribution of the model-adjusted means is illustrated by catseye plots with shaded +/- standard error overlaid by scatterplots of subject measures; scatterplots have been randomly jittered horizontally for clarity, and are shown on the log (base-10) scale such that comparisons are against a null value of 1. * P < 0.05, ** P < 0.01, *** P < 0.001, n.s. not significant.

Extended Data Fig. 8. Fitness comparisons of Dakar-4 amino-acid mutant and FSS13025-4 amino-acid mutant versus wild-type ZIKV strains in human primary cells.

a, **b**, Fitness comparison between Dakar 4-amino acid mutant (DK-4M) and wild-type strain in human primary fibroblast (**a**) and keratinocyte (**b**) cells. **c**, **d**, The fitness comparison between FSS13025 4-amino acid mutant (FSS-4M) and wild-type strain in human primary fibroblast (**c**) and keratinocyte (**d**) cells. Each point represents a single culture sample. **a-d**, The distribution of the model-adjusted means is illustrated by catseye plots with shaded +/- standard error overlaid by scatterplots of subject measures; scatterplots have been randomly jittered horizontally for clarity, and are shown on the log (base-10) scale such that comparisons are against a null value of 1.

790 Extended Data Fig. 9. Fitness comparison between NS1-188 mutants against wild-type ZIKV strains in different organs of A129 mice.

a, Fitness comparison between Dakar NS1-V188A mutant and wild-type strain in different organs of A129 mice. **b**, Fitness comparison between FSS13025 NS1-A188V mutant and wild-

- type strain in different organs of A129 mice. Each point represents a single mouse sample **a**, **b**, The distribution of the model-adjusted means is illustrated by catseye plots with shaded +/- standard error overlaid by scatterplots of subject measures; scatterplots have been randomly jittered horizontally for clarity, and are shown on the log (base-10) scale such that comparisons are against a null value of 1. * P < 0.05, ** P < 0.01, *** P < 0.001, n.s. not significant.
- 800 Extended Data Fig. 10. Fitness comparison between African-4 amino-acid mutant and FSS13025-4 amino-acid mutant against wild-type viruses during mosquito-mouse-mosquito transmission cycle.

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- **a**, Fitness comparison between Dakar 4-amino acid mutant (DK-4M) and wild-type strain in different organs of A129 mice after being bitten by infected mosquitoes. **b**, Fitness comparison between FSS 4-amino acid mutant (FSS-4M) and wild-type strain in different organs of A129 mice biting by infected mosquitoes. Each point represents a single mosquito or mouse sample **a**, **b**, The distribution of the model-adjusted means is illustrated by catseye plots with shaded +/-standard error overlaid by scatterplots of subject measures; scatterplots have been randomly jittered horizontally for clarity, and are shown on the log (base-10) scale such that comparisons are against a null value of 1. * P < 0.05, ** P < 0.01, *** P < 0.001.
 - Extended Data Table 1. The relative replicative fitness of ZIKV mutant strains in mosquitoes, mice and human primary cells.
- The relative replicative fitness of ZIKV mutant strains in mosquitoes, mice and human primary cells, derived from the data in the figures, are presented as the final ratio divided by the initial ratio of the two competing viruses, as used previously to compare adaptive chikungunya virus mutations³⁶. P values are based on differences from equal fitness or a relative fitness value of 1.
- Extended Data Table 2. Primers and probes for gene cloning, qPCR, RT-PCR and Sanger sequencing.

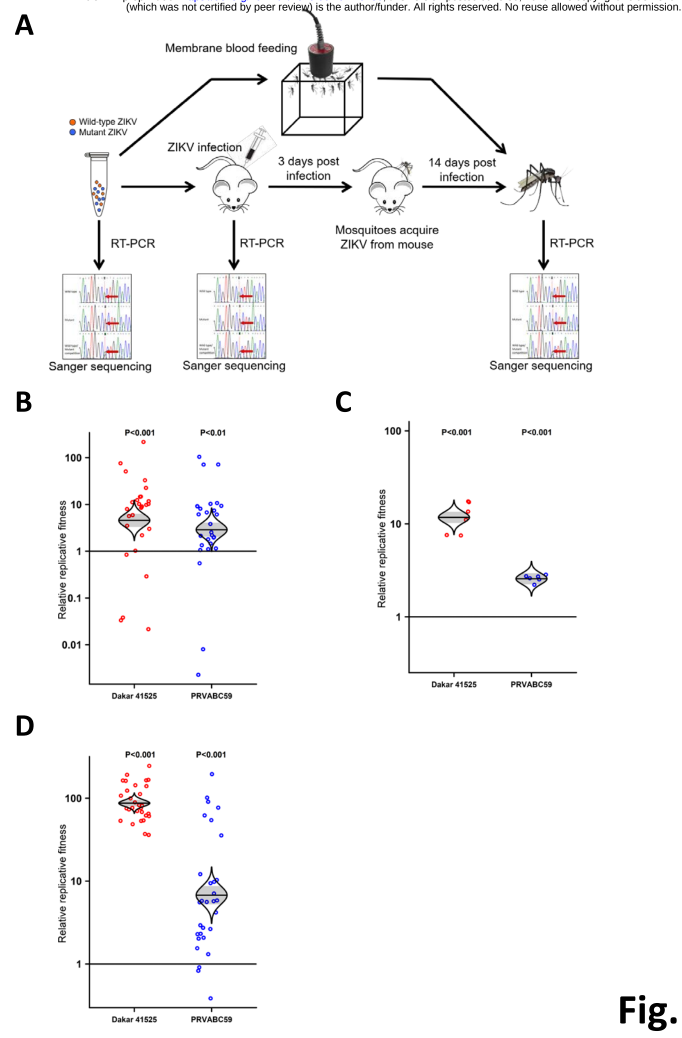
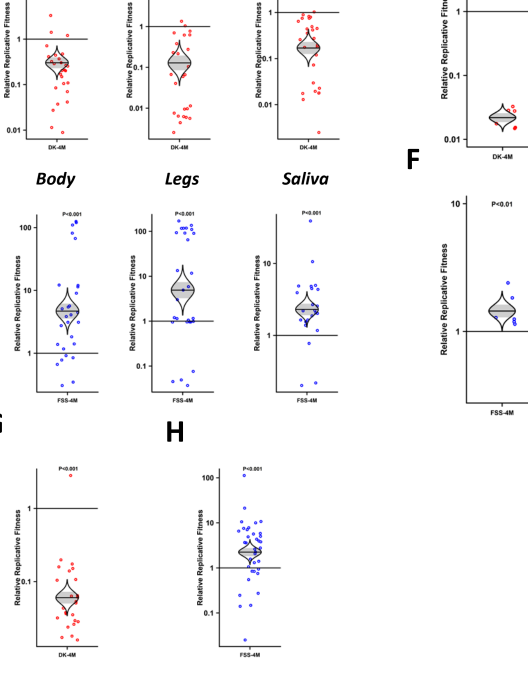


Fig. 1



Α

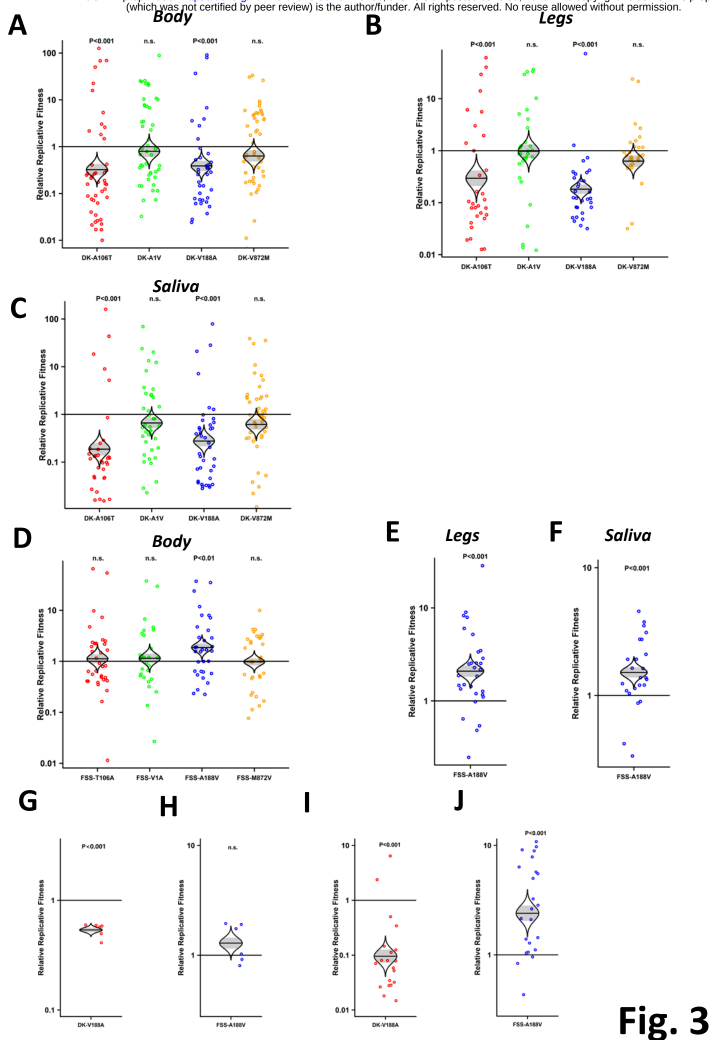
Fitness for epidemic transmission

10

E

G

Fig. 2



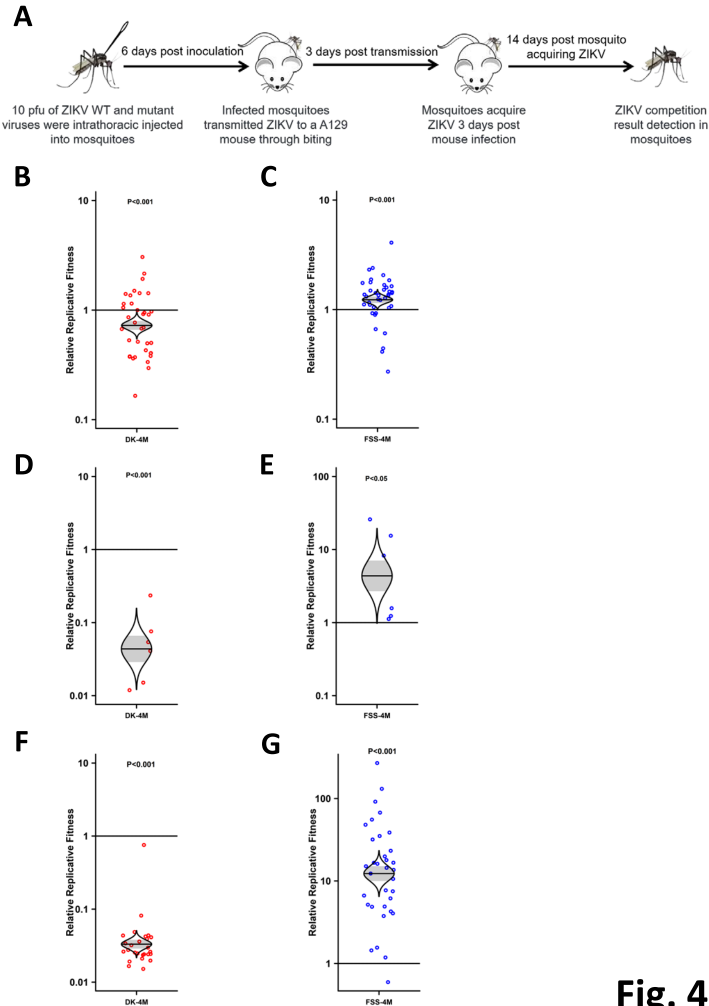
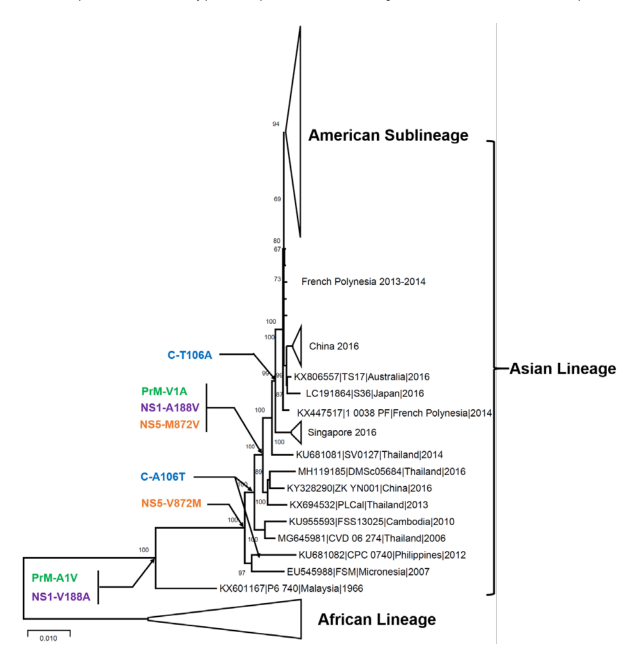


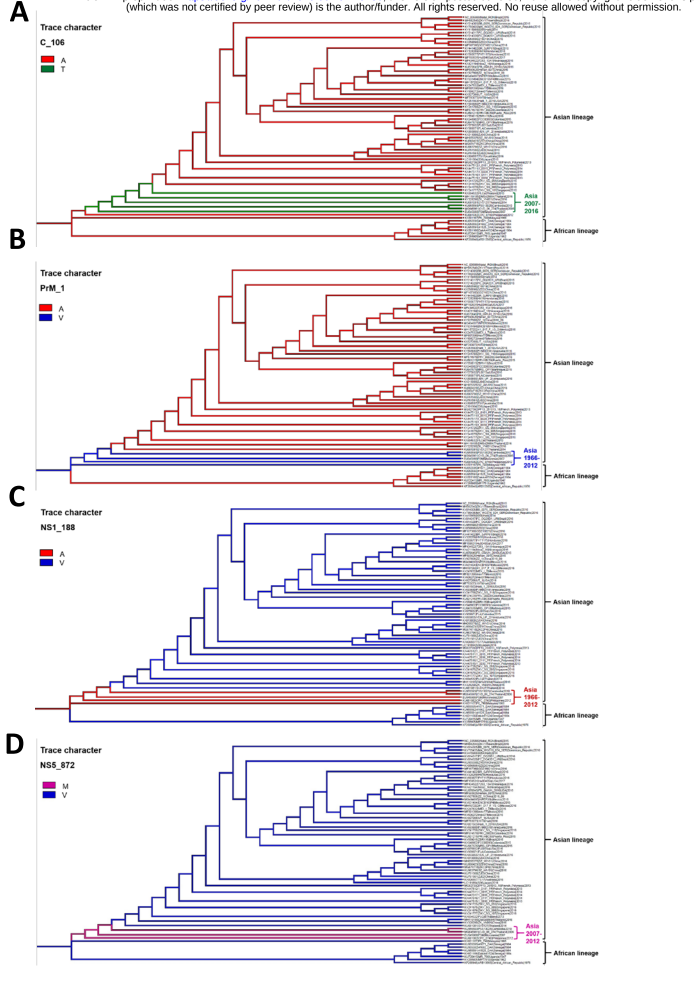
Fig. 4





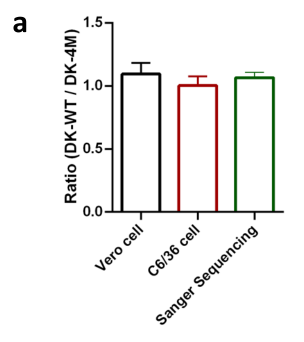
B

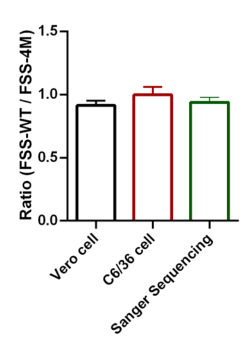
Position		African Lineage		Asian Linea	ge (Pre-epidemic)	Asian Lineage (American)		
Gene	Amino acid	Codon	Amino acid	Codon	Amino acid	Codon	Amino acid	
Capsid	106	GCT	Α	ACA	Т	GCA	A	
PrM	1	GCC	Α	GTG	V	GCG	Α	
NS1	188	GTC	V	GCT	Α	GTT	V	
NS5	872	GTG	V	ATG	М	GTG	V	

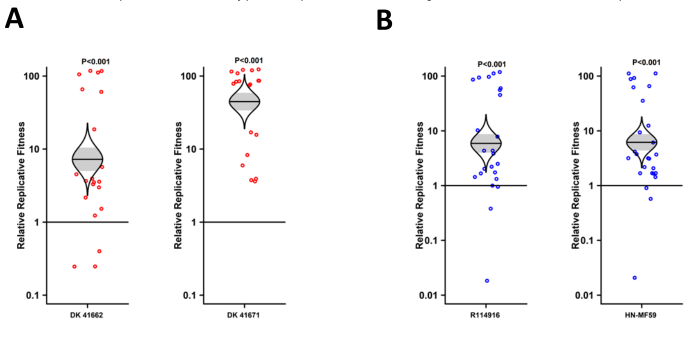


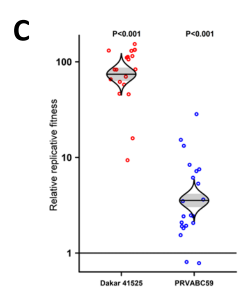
Extended Data Figure 2

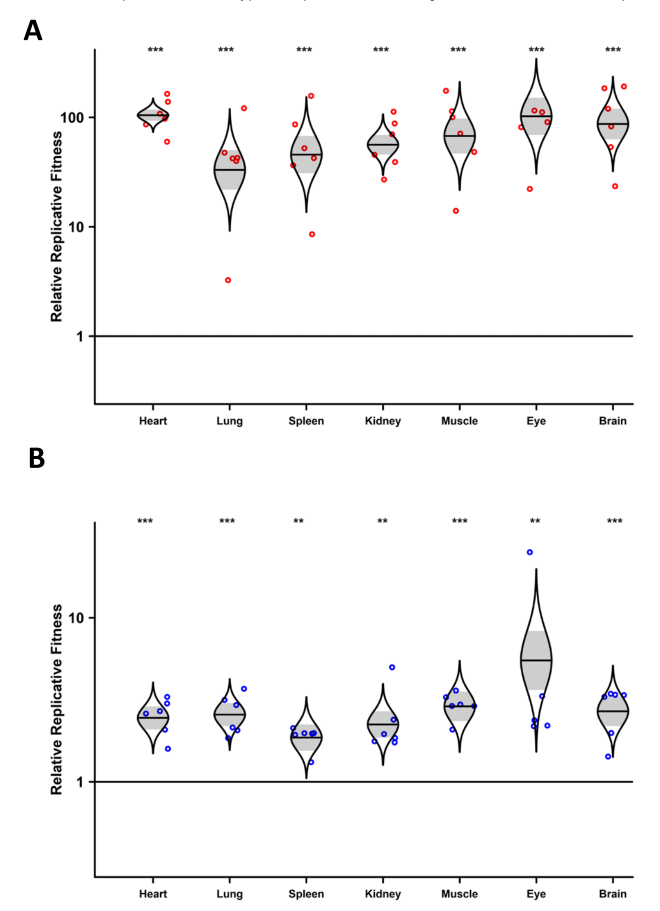
b

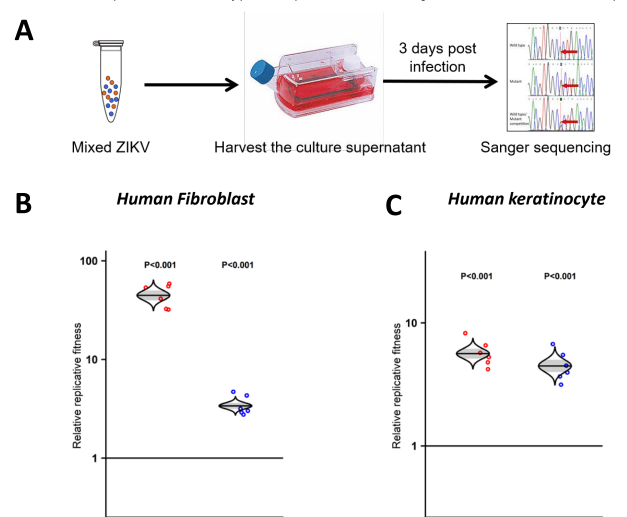












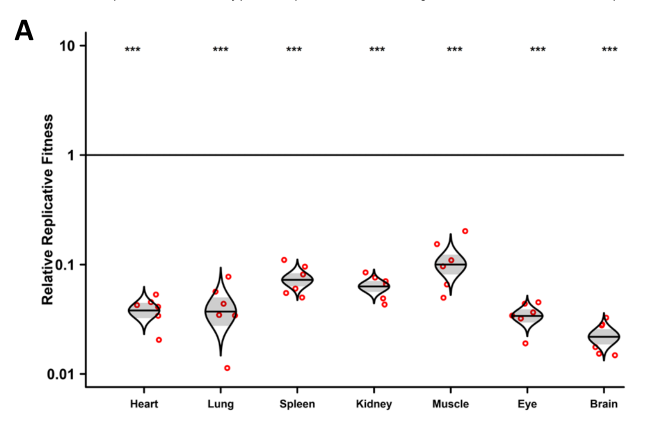
Dakar 41525

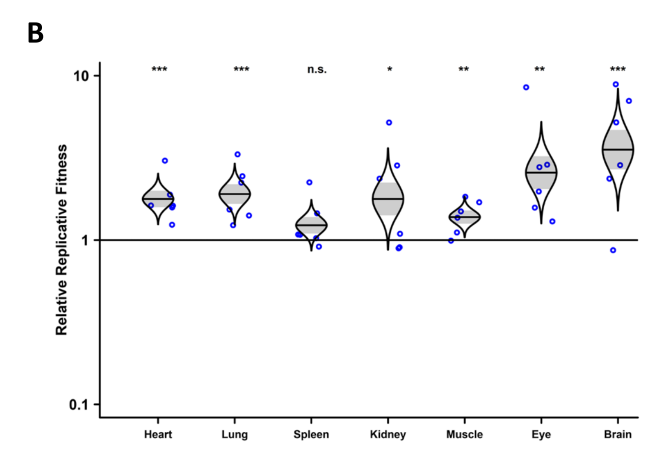
PRVABC59

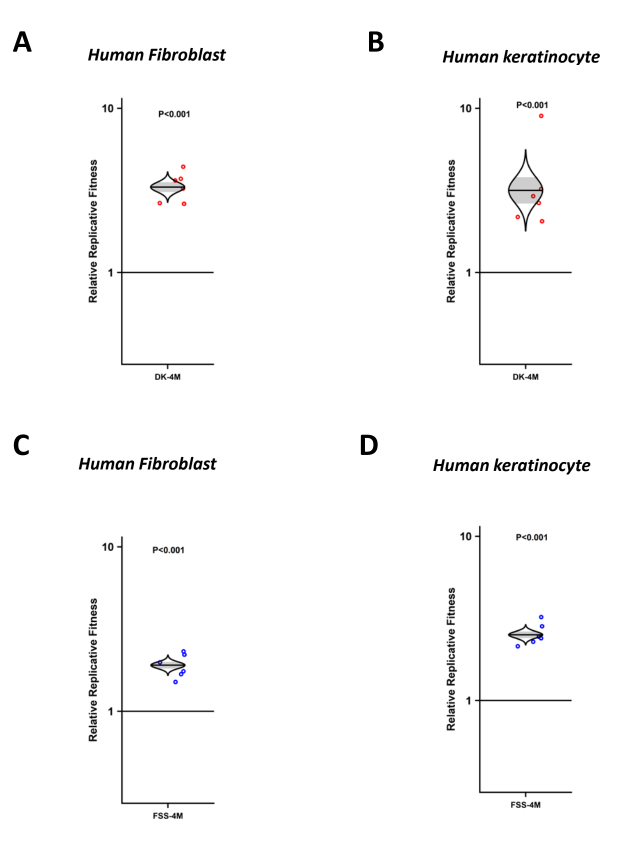
Extended Data Figure 6

Dakar 41525

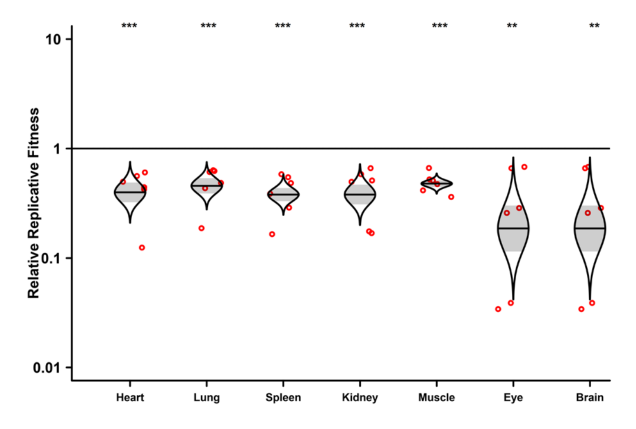
PRVABC59

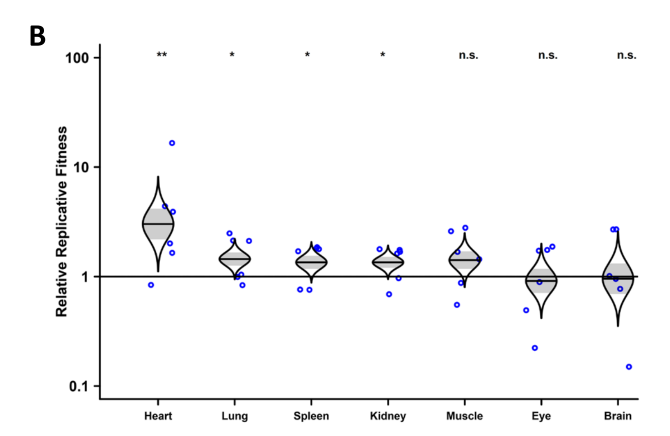


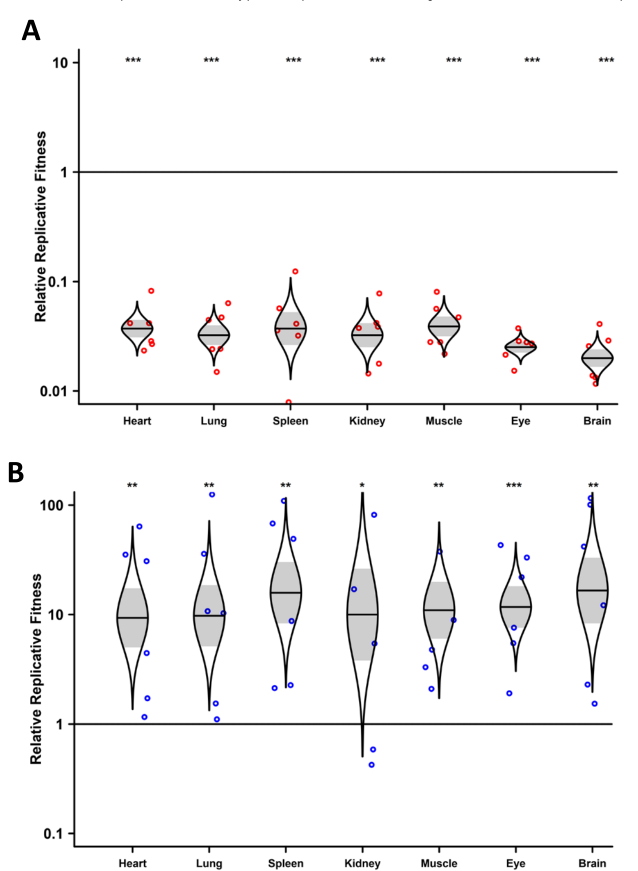












Extended Data Table 1. The relative replicative fitness^a of ZIKV mutant strains in mosquitoes, mice and human primary cells.

Virus backbone	Virus competition	Mosquito body	Mosquito leg	Mosquito saliva	Mouse serum	Human Keratinocyte	Human Fibroblast
Dakar virus backbone	FSS13025/Dakar 41525	0.4*			0.1***	0.1***	0.02***
	DK-4M/DK-WT	0.4***	0.3***	0.3***	0.2**	0.3***	0.3***
	DK-A106T/DK-WT	0.4*	0.5*	0.3**			
	DK-A1V/DK-WT	1.0	1.1	0.8			
	DK-V188A/DK-WT	0.4*	0.2**	0.4*	0.5*		
	DK-V872M/DK-WT	0.9	0.7	0.8			
FSS13025 virus backbone	PRVABC59/FSS13025	2.0*			2.7***	3.54***	3.3***
	FSS-4M/FSS-WT	3.68***	2.8***	2.1*	1.3	2.74**	1.9*
	FSS-T106A/FSS-WT	1.0					
	FSS-V1A/FSS-WT	1.1					
	FSS-A188V/FSS-WT	1.5*			1.3*		
	FSS-M872V/FSS-WT	1.0					

^a Defined as final competitor ratio / inoculum ratio, as described previously to compare adaptive chikungunya virus mutations³⁵.

^{*} P<0.05; ** P<0.01; *** P<0.001.

AGRCAAGCAGTCTCCCGGATG

FAM-TGCCGCCACCAAGATGAACTGATTGGCCG-TAMRA

Extended Data Table 2. Primers and probes for gene cloning and qPCR

ZIKV envelope gene

Primers for cloning ZIKV full-length mutants	Upper primer	Lower primer
FSS-T106A	GAAGAAGAGACGAGGCgCAGATACTAGTGTCG	CGACACTAGTATCTGcGCCTCGTCTCTTCTTC
FSS-VIA	CCACAGCCATGGCAGcGGAGGTCACTAGACG	CGTCTAGTGACCTCCgCTGCCATGGCTGTGG
FSS-A188V	GTCATTGGAACAGCCGtTAAGGGAAAGGAGGCT	TAGCCTCCTTTCCCTTAaCGGCTGTTCCAATGAC
FSS-M872V	CACAGTCAACATGgTcCGTAGGATCATAGGTG	CACCTATGATCCTACGgAcCATGTTGACTGTG
DK-A106T	GGAAGAGACGTGGCaCTGACACCAGCATCG	CGATGCTGGTGTCAGtGCCACGTCTCTTCC
DK-AIV	CACAGCCATGGCAGtCGAGATCACTAGACG	CGTCTAGTGATCTCGaCTGCCATGGCTGTG
DK-VI88A	CCGTCATAGGAACAGCTGcCAAGGGAAAGG	CCTTTCCCTTGgCAGCTGTTCCTATGACGG
DK-V872M	AGACACAGTCAACATGaTGCGTAGGATCAT	ATGATCCTACGCAtCATGTTGACTGTGTCT
Primers for one-step RT-PCR	Upper primer	Lower primer
FSS-T106A	GGCGATTCTAGCCTTTTTG	TACCCCCTCATCCAGCATA
FSS-VIA	GGCGATTCTAGCCTTTTTG	TACCCCCTCATCCAGCATA
FSS-A188V	TGACACACTGAAGGAATGC	CACTATGCCATGGCCCTTT
FSS-M872V	TGGAACAGAGTGTGGATTG	ATGGCGTTCTCGGCCTGACTA
DK-A106T	GGCGATACTAGCCTTCTTG	CAGGTTCCGTACACACCCA
DK-AIV	GGCGATACTAGCCTTCTTG	CAGGTTCCGTACACACCCA
DK-V188A	GCCAGTGCCTGTGAATGA	CACTATGCCATGGCCCTTT
DK-V872M	AGAGTGTGGATTGAGGAGAA	ACTCAGTGTCCTCTGAGGGG
Primers for sanger sequencing	Upper primer	
FSS-T106A	GGCGATTCTAGCCTTTTTG	
FSS-VIA	GGCGATTCTAGCCTTTTTG	
FSS-A188V	TGACACACTGAAGGAATGC	
FSS-M872V	TGGAACAGAGTGTGGATTG	
DK-A106T	GGCGATACTAGCCTTCTTG	
DK-AIV	GGCGATACTAGCCTTCTTG	
DK-V188A	GCCAGTGCCTGTGAATGA	
DK-V872M	A CA CTCTCC ATTCA CCA CA A	
	AGAGTGTGGATTGAGGAGAA	

AGGACGGGAGRTCCATTGTGG