Vector-borne transmission and evolution of Zika virus

Gladys Gutiérrez-Bugallo¹, Luis Augusto Piedra¹, Magdalena Rodriguez¹, Juan A. Bisset¹, Ricardo Lourenco-de-Oliveira², Scott C. Weaver³, Nikos Vasilakis ⁶ and Anubis Vega-Rúa ⁵*

Zika virus (ZIKV), discovered in the Zika Forest of Uganda in 1947, is a mosquito-borne flavivirus related to yellow fever, dengue and West Nile viruses. From its discovery until 2007, only sporadic ZIKV cases were reported, with mild clinical manifestations in patients. Therefore, little attention was given to this virus before epidemics in the South Pacific and the Americas that began in 2013. Despite a growing number of ZIKV studies in the past three years, many aspects of the virus remain poorly characterized, particularly the spectrum of species involved in its transmission cycles. Here, we review the mosquito and vertebrate host species potentially involved in ZIKV vector-borne transmission worldwide. We also provide an evidence-supported analysis regarding the possibility of ZIKV spillback from an urban cycle to a zoonotic cycle outside Africa, and we review hypotheses regarding recent emergence and evolution of ZIKV. Finally, we identify critical remaining gaps in the current knowledge of ZIKV vector-borne transmission.

ika virus (ZIKV) is an arthropod-borne flavivirus related to yellow fever, dengue and West Nile viruses¹. ZIKV was first reported in East Africa in 1947 and expanded from the ancestral enzootic cycle in Africa to Asia several decades ago (Fig. 1). At the beginning of the 21st century, the virus expanded into the South Pacific and the Americas, triggering a pandemic that led to 48 countries reporting active ZIKV transmission by 2017². Prior to this expansion, there was little scientific research on this virus³.

In this paper, we review the natural history of ZIKV and the current knowledge about ZIKV vector-borne transmission and the mosquito and vertebrate host species potentially involved world-wide. Furthermore, we discuss the possibility of ZIKV spillback into an enzootic cycle outside Africa and review hypotheses regarding ZIKV recent global emergence and evolution. Finally, we identify research priorities for filling remaining gaps and challenges in our understanding of ZIKV.

Zika virus natural history

The virus was first isolated from a sentinel rhesus macaque and from *Aedes (Stegomyia) africanus* mosquitoes in the same Ugandan location. Surveillance efforts identified immune people in at least 25 African countries from 1945 to 2014 (reviewed in ref. ⁴) and in 7 Asian countries or territories from 1952 to 1997 (Fig. 1). However, many of the serologic tests employed in this surveillance are cross-reactive among flaviviruses, thus these results must be interpreted with caution. Nevertheless, direct detection of ZIKV in countries including Senegal, the Ivory Coast, Burkina Faso, Nigeria, the Central African Republic, Uganda, Thailand, Malaysia, Indonesia and the Philippines (reviewed in ref. ⁵) confirmed its widespread distribution for many decades.

The first evidence of possible interhuman urban transmission came from Malaysia in 1966, when ZIKV was detected in the domesticated, anthropophilic mosquito, *Aedes (Stegomyia) aegypti aegypti*. However, between 1966 and 2013, there was no direct evidence of

ZIKV transmission by this species, which may reflect limited urban transmission or simply a lack of surveillance and diagnostics to detect endemic or epidemic circulation.

The first ZIKV outbreaks were documented in Gabon⁶ and Yap^{7,8} in 2007. In 2013, a major outbreak started in French Polynesia involving approximately 100,000 infected persons, providing the first evidence of a severe disease, Guillain Barré syndrome, associated with ZIKV infection⁹. This was followed by a spread throughout the South Pacific and eventually to the Americas, where a massive epidemic ensued and congenital microcephaly was first associated with maternal ZIKV infection² (Fig. 1). Millions of ZIKV cases have now been reported from 86 countries, including the expanded forms of congenital infection that comprise congenital Zika syndrome (CZS) and Guillain Barré syndrome¹⁰.

ZIKV vectors: evidence in the wild and from the laboratory

To properly incriminate a ZIKV vector, the virus must be isolated from field-collected mosquitoes, and laboratory studies must ascertain their ability to transmit the virus (vector competence; Supplementary Figure 1). Field surveys allow investigation of a large number of mosquito species, but do not deliver proof of transmission, because nondisseminated infections may occur or residual virus may be detected in a partially digested blood meal. Experimental vector-competence studies can demonstrate transmission potential, but practical constraints limit the number of species for which this can be performed, and competent species may not have proper contact with hosts to transmit the virus in nature. Consequently, combined information from both type of study is critical to the definition of the most plausible ZIKV vectors involved in sylvatic and urban transmission cycles (Fig. 2a).

Field investigations

Our literature search conducted in April 2018 reported 31 wild-caught mosquito species infected with ZIKV worldwide

¹Department of Vector Control, Center for Research, Diagnostic and Reference, Institute of Tropical Medicine Pedro Kourí, PAHO-WHO Collaborating Center for Dengue and its Control, Havana, Cuba. ²Fundação Oswaldo Cruz-Fiocruz, Instituto Oswaldo Cruz, Laboratório de Mosquitos Transmissores de Hematozoários, Rio de Janeiro, Brazil. ³Department of Microbiology and Immunology, University of Texas Medical Branch, Galveston, TX, USA. ⁴Department of Pathology, University of Texas Medical Branch, Galveston, TX, USA. ⁵Laboratory of Vector Control Research, Unit Transmission Reservoir and Pathogen Diversity, Institute Pasteur of Guadeloupe, Les Abymes, Guadeloupe, France. *e-mail: avega@pasteur.fr

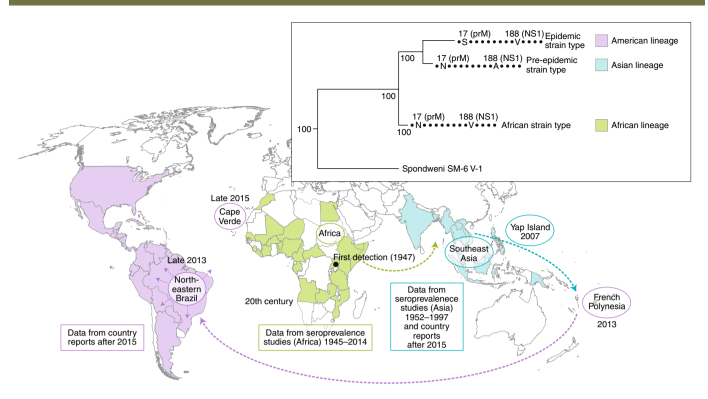


Fig. 1 | Emergence and evolution of Zika virus vector-borne transmission. Zika virus was first detected in Uganda in 1947. Seroprevalence data suggests the virus was present in Africa and Asia starting from the 1950s and spread to the Pacific and the Americas from 2013 onwards. There are three recognized lineages of Zika virus (African lineage, green; Asian lineage, blue and American lineage, purple), characterized by key pre-epidemic and post-epidemic mutations, shown on the phylogeny.

(Supplementary Table 1). These mosquitoes belong to five genera: Aedes (22 species), Culex (4 species), Anopheles and Eretmapodites (2 species each) and Mansonia (1 species) (Fig. 3). Among the Aedes mosquitoes, Stegomyia is the major subgenus from which ZIKV has been isolated (9 species). This distribution suggests the Aedes genus as the most important mosquito taxon involved in ZIKV transmission. Most ZIKV-infected mosquito species (25) have been caught in sylvatic settings, whereas only six species have been reported to be infected with ZIKV in urban settings: Ae. aegypti^{4,11-21}, Aedes (Stegomyia) albopictus^{6,12,22}, Culex (Culex) quinquefasciatus^{15,20,23}, Aedes (Aedimorphus) vexans²⁰, Culex (Culex) coronator²⁰ and Culex (Culex) tarsalis²⁰. With the exception of Ae. albopictus mosquitoes, ZIKV has been detected in salivary gland pools of all of these mosquito species caught in urban settings in Mexico²⁰. Nevertheless, the viral titers associated with these detections were often not compatible with transmission competence. To incriminate a vector, ZIKV titers consistent with transmission-competence are necessary; however, viral loads have only been measured by quantitative reverse transcription PCR (RT-qPCR) on five occasions, in Ae. aegypti, Ae. vexans, Ae. albopictus, Cx. tarsalis, Cx. coronator and Cx. quinquefasciatus mosquitoes 15,16,20,22,24.

Field collection of ZIKV-infected mosquitoes has been performed in 15 countries: 9 in epidemic regions and 6 in regions with enzootic ZIKV circulation (Fig. 3). The sylvatic species that have been more frequently found infected with ZIKV in the field are Aedes (Stegomyia) luteocephalus, Aedes (Diceromyia) furcifer, Aedes (Aedimorphus) dalzieli, Ae. aegypti, Ae. africanus, Aedes (Fredwardsius) vittatus and Aedes (Diceromyia) taylori. In urban settings, Ae. aegypti has been most frequently found ZIKV-infected and is the most widely distributed, with ZIKV detection in Africa, America, Asia and Oceania (Supplementary Table 1). Ae. africanus, Ae. furcifer and Ae. luteocephalus are also broadly

distributed in Africa. Finally, 14 species have been reported as being infected with ZIKV only once: Anopheles (Cellia) gambiae s.l., Aedes (Stegomyia) apicoargenteus, Aedes (Diceromyia) flavicollis, Aedes (Neomelaniconion) taeniarostris, Eretmapodites inornatus, Eretmapodites quinquevittatus, Ae. (Neomelaniconion) jamoti, Aedes (Mucidus) grahamii, Aedes (Aedimorphus) hirsutus, Culex (Culex) perfuscus, Aedes (Stegomyia) unilineatus, Ae. vexans, Cx. coronator and Cx. tarsalis (Supplementary Table 1). Although ZIKV detection in field-collected mosquitoes does not prove their role as vectors, the distribution and frequencies of infected mosquitoes suggest that Ae. africanus, Ae. furcifer, Ae. lutocephalus, Ae. vittatus, Ae. dalzielli and Ae. taylori are involved in sylvatic ZIKV transmission cycles, and Ae. aegypti is implicated in urban ZIKV cycles. The latter is supported by anthropophilic and anthropophagic behaviours of Ae. aegypti and the high abundance of this species in urban environments nearly throughout the tropics^{25,26}.

Vector competence

Our April 2018 literature survey identified 45 appropriate ZIKV vector-competence studies involving 25 mosquito species: 13 Aedes or Ae. (Ochlerotatus) spp. (Ochlerotatus is elevated to a genus by some authors), 8 Culex spp. and 4 Anopheles spp. (Supplementary Table 1). Eight of these species have been found to be able to transmit ZIKV in laboratory conditions, as infectious virus has been found in mosquito salivary glands or saliva: Ae. aegypti, Ae. albopictus, Ae. vexans, Aedes (Rampamyia) notoscriptus, Aedes (Ochlerotatus) camptorhynchus, Ae. vittatus, Ae. luteocephalus and Cx. quinquefasciatus.

Among the sylvatic mosquitoes examined, Ae. vittatus and Ae. luteocephalus were found to be competent ZIKV laboratory vectors, in agreement with the high frequency and distribution of ZIKV-infected specimens from these species found in the

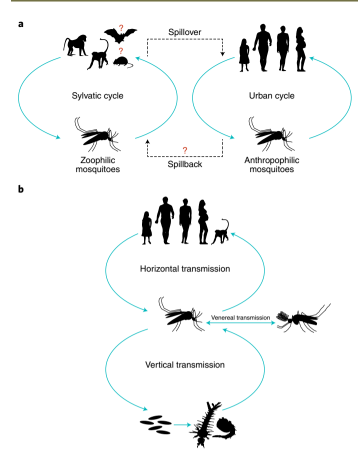


Fig. 2 | Zika virus vector-borne transmission. a, Horizontal transmission of ZIKV occurs in two distinct cycles: a sylvatic cycle in which the virus circulates between animal vertebrate hosts (i.e., probably nonhuman primates) and zoophilic mosquitoes, and an urban cycle in which the virus is transmitted to humans by highly anthropophilic mosquitoes (such as Ae. aegypti). b, Vector-borne transmission of ZIKV occurs via three main patterns: horizontal transmission between vertebrates and vectors, vertical transmission from and infected mosquito female to its progeny, and venereal transmission from an infected female to a mosquito male and vice versa.

field. This suggests their implication in sylvatic cycles in Africa. Outside Africa, many publications have recognized the roles of *Aedes (Stegomyia) hensilli* and *Aedes (Stegomyia) polynesiensis* in the Yap Island and French Polynesia ZIKV outbreaks, respectively, because of their high abundances in households^{7,8,27}, although these species have never been found to be infected in the field^{4,5,28,29}. Vector-competence assays have proved the ability of the two species to develop disseminated ZIKV infections, but have not confirmed transmission^{7,30}. Thus, further studies are needed in order to fully evaluate the role *Ae. hensilli* and *Ae. polynesiensis* in urban ZIKV cycles.

The Culex spp. controversy

Despite field reports of ZIKV infection in four *Culex* species (Fig. 3; Supplementary Table 1) and experimental oral infection assays highlighting that *Cx. quinquefasciatus* is susceptible to ZIKV with viral detection in saliva^{15,31,32}, there are so far no reports of naturally infected mosquitoes during outbreaks with viral loads consistent with transmission competence. Even if the infection rates of mosquito populations in nature are very low³³—meaning that ZIKV detection will be strongly dependent on the study sampling effort—the majority of field investigations in which *Cx. quinquefasciatus*

mosquitoes were captured did not find any ZIKV infection in this species^{4,6,7,14,16,21,34}.

Vector-borne transmission in natural conditions (known as vectorial capacity) is also determined by the abundance of the vector and its ecological traits, which are quantitatively more important than vector competence (although competence is a requirement for transmission). This may explain why some 'poor vectors' can mediate outbreaks regardless of limited vector competence³⁵. *Cx. quinquefasciatus* is a highly abundant species that may bite humans frequently in urban settings³⁶, which favours the human-vector contact that is critical to vectorial capacity³⁵ and thus, may occasionally transmit ZIKV. Nevertheless, efficient ZIKV transmission by *Cx. quinquefasciatus* would require sequential feedings on human hosts, and this species is more often associated with avian bloodmeals than human bloodmeals³⁷. Taken together, the collected evidence suggests that *Culex* species does not play an important role on ZIKV transmission.

ZIKV vertical and venereal transmission reports

Vertical transmission of arboviruses in mosquitoes (from an infected female mosquito to her offspring) has been suggested as a mechanism that ensures maintenance of ZIKV during conditions adverse for horizontal transmission among vertebrate hosts (such as harsh winters, drought conditions and interepidemic stages limited by herd immunity), as well as a potential influence on the epidemiology of arbovirus infection³⁸. Vertical or venereal ZIKV transmission has been demonstrated for the mosquitoes Ae. furcifer^{34,39}, Ae. aegypti^{16,20} and Cx. quinquefasciatus²⁰ under natural conditions, as adult males infected with ZIKV have been caught in the field (Supplementary Table 2). In addition, ZIKV has been detected in Ae. albopictus eggs collected in Brazil22 and in adult Aedes (Stegomyia) bromeliae, Aedes unilineatus and Ae. vittatus reared from field-collected eggs (reviewed in ref. 5), suggesting the occurrence of natural vertical transmission in these species. Experimental studies have demonstrated vertical ZIKV transmission in Ae. aegypti⁴⁰⁻⁴² and Ae. albopictus⁴², whereas venereal transmission has been observed in both directions (male-to-female and vice versa) in Ae. aegypti^{43,44} (Supplementary Table 2). These findings reveal the complexity of vector-borne ZIKV transmission dynamics (Fig. 2b). Although these intervector transmission modes guarantee the spread and maintenance of the virus in vector populations, its contribution to the mosquito-human transmission chain and epidemiology continues to be a knowledge gap, because no infectious virus has been reported in saliva of vertically or venereally infected female mosquitoes.

ZIKV potential host species

Vertebrate animals are also major players in the ZIKV maintenance in nature, as ZIKV circulates in a sylvatic transmission cycle between nonhuman primate hosts and arboreal mosquitoes in tropical Africa⁴⁵ (Fig. 2a). Our literature review identified 42 publications reporting ZIKV-infected animals, in either natural or experimental conditions.

At least 79 animal species have been identified as naturally infected or experimentally susceptible to ZIKV infection (Supplementary Table 3). Mammals, and especially primates, are the most represented taxonomic group, although birds, reptiles and amphibians have also been identified, indicating the potential diversity of ZIKV hosts and the lack of a clear association between ZIKV and particular animal taxa (Fig. 4, Supplementary Table 3). Among these species, 63 have been found as naturally 'infected', whereas Macaca mulatta, Cercopithecus aethiops, Cercopithecus ascanius, Callithrix jacchus, Eidolon helvum, and Capra aegagrus have been recognized as being susceptible to ZIKV infection in both natural and experimental conditions. However, the serological assays generally used to assess ZIKV exposure and susceptibility are of lim-

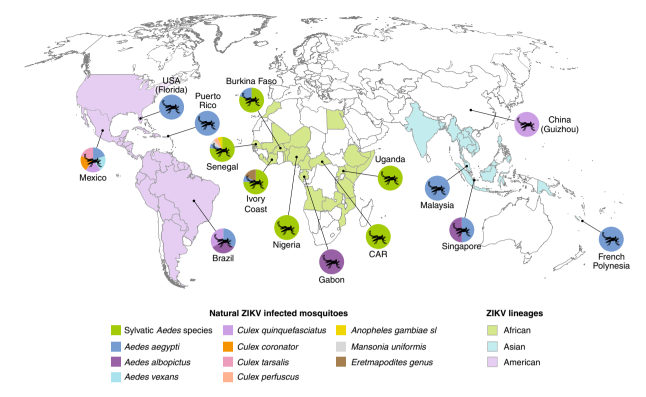


Fig. 3 | Distribution of vector species naturally infected with Zika virus. The colour code used for countries and territories represents the Zika virus lineages that have been epidemiologically associated with human infections in those regions.

ited specificity and could result in overestimation of the number of species susceptible to ZIKV infection.

The incrimination of such animals in ZIKV transmission chains (either sylvatic or urban) will depend on their viremia levels and duration, their spatial coincidence with competent ZIKV vectors and the feeding preferences of the latter. So far, viremia levels sufficient for transmission have only been detected in five monkey species (*M. mulatta*, *C. jacchus*, *Macaca fascicularis*, *Saimiri boliviensis boliviensis and Saimiri sciureus sciureus*) infected experimentally (Supplementary Table 3).

The ZIKV sylvatic transmission cycle between nonhuman hosts and arboreal mosquitoes has been most thoroughly studied in Africa (Fig. 4), and seroprevalence data suggest that nonhuman primates are important enzootic amplification hosts for ZIKV maintenance. The most intensively studied enzootic ZIKV focus is in eastern Senegal, where annual mosquito collections and periodic nonhuman primate sampling have revealed strong evidence of regular or continuous arboreal circulation. Interestingly, the same area also harbours dengue, chikungunya and yellow fever viruses with no apparent difference in nonhuman primate amplification hosts or mosquito vectors. However, whereas the latter three viruses are detected in mosquitoes only periodically in asynchronous 7-8year cycles, ZIKV is detected more frequently (ca. 4-year cycles)³⁹. This suggests that ZIKV may exploit additional, shorter-lived vertebrate hosts with faster turnover than the three monkey species (Chlorocebus aethiops, Erythrocebus patas and Papio papio) typically found in the Senegalese focus. Unfortunately, knowledge of animal reservoirs is lacking in regions outside Africa, as only four studies have been published on this topic: Brazil⁴⁶, Pakistan⁴⁷, Indonesia⁴⁸ and Thailand⁴⁹ (Fig. 4; Supplementary Table 3). Additional research using appropriate diagnostic techniques on potential zoonotic ZIKV hosts could offer important predictive data about the possibility of enzootic establishment outside Africa. Diagnostic methods allowing determination of viral loads in vertebrate hosts should be privileged, as this information is crucial to knowing which species

can generate sufficient viremia to infect mosquitoes and thus contribute to circulation.

Establishment of a ZIKV enzootic cycle in the Americas

The arboviruses yellow fever (YFV), chikungunya (CHIKV) and dengue (DENV) were introduced to the Americas following European discovery in the late 15th Century. An enzootic transmission cycle has established for YFV in the Americas⁵⁰ but has not been documented for CHIKV⁵¹ and DENV⁵².

The recent introduction and explosive spread of ZIKV in urban and rural settings throughout the Americas begs the question of whether spillback and establishment of an enzootic ZIKV transmission cycle could occur in the Americas. Brazil is home to multiple species of primates and mosquitoes potentially capable of ZIKV transmission, although direct assessment of host competence and vector competence of New World species is currently not known. The probability of enzootic establishment is dependent on host and vector population sizes, host birth rates and the ZIKV force of infection. We urgently need investigations of the host competence of New World monkeys or other mammals to ZIKV, of the vector competence of sylvatic New World Aedes, Sabethes, and Haemagogus mosquitoes for ZIKV and of the geographic range of these potential hosts and vectors. Moreover, independent enzootic circulation is difficult to distinguish from temporary spillback from interhuman transmission unless sufficient divergence between enzootic and human strains can be detected or epizootics in nonhuman primates suggest the presence of an enzootic cycle. Nonetheless, based on the experience acquired with the enzootic transmission cycle of YFV, the establishment of a ZIKV enzootic cycle in the Americas (or in Asia) could have important implications for the ultimate control of future epidemics, as these cycles are not amenable to human intervention and control.

Transmission in neotropical sylvatic mosquito species

Transmission of ZIKV in urban settings occurs mainly via Aedes spp. mosquitoes. However, because of the invasive nature and

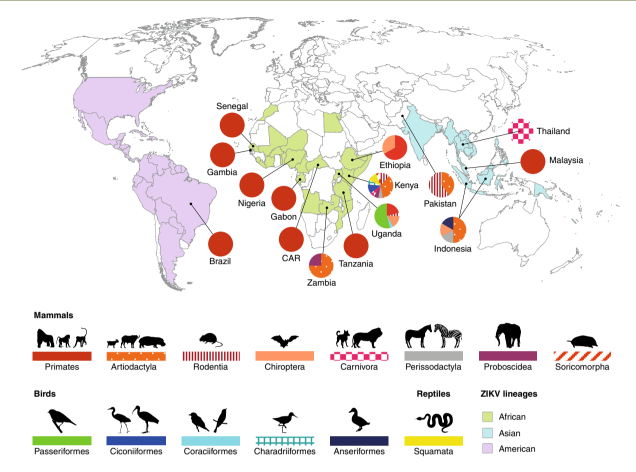


Fig. 4 | Distribution of host orders reported as susceptible to Zika virus in natural conditions. The colour code used for countries and territories represents the Zika virus lineages that have been epidemiologically associated with human infections in those regions (serological assays have been the tests used to assess ZIKV exposure and susceptibility in 87.1% of animals sampled in the field).

extensive geographic distribution of Ae. albopictus in tropical as well as temperate settings, this species has the potential to become a major ZIKV vector globally. Given that it has been implicated in Asia as a bridge vector for sylvatic DENV movement into the urban cycle (reviewed in refs. 52,53), could this vector serve as the bridge to facilitate reverse spillback and establishment of an enzootic ZIKV transmission cycle in the Americas? Subsequently, a number of New World mosquitoes involved in the enzootic transmission of YFV, including Haemagogus (Haemagogus) albomaculatus, Haemagogus (Haemagogus) spegazzini, Haemagogus (Haemagogus) janthinomys, Sabethes (Sabethoides) chloropterus, Sabethes (Sabethes) albiprivus, Sabethes (Sabethoides) glaucodaemon, Sabethes (Peytonulus) soperi, and Sabethes (Sabethes) cyaneus, Psorophora (Janthinosoma) ferox and Ae. (Ochlerotatus) serratus (reviewed in ref. 53) could serve as competent enzootic vectors of ZIKV and should be evaluated experimentally.

A recent mathematical dynamic transmission model to assess the probability of establishment of a sylvatic ZIKV transmission cycle in nonhuman primates and/or other mammals, via arboreal mosquito vectors in Brazil, demonstrated a high probability of establishment of enzootic ZIKV across a large range of biologically plausible parameters: mean mosquito lifespan, mean ZIKV recovery in nonhuman primates (NHPs), mosquito vertical transmission and rate of yearly ZIKV introduction⁴⁵. Whether ZIKV will emulate YFV (neotropical enzootic establishment) or DENV (lack of evidence for neotropical enzootic transmission) is an open and critical question. If ZIKV establishes a sylvatic cycle in the Americas (or in Asia), then mosquito control and even human herd immunity from vaccination (if a ZIKV vaccine becomes available) or limitation of

transmission from demographic turnover will not suffice to eradicate it from the region.

Transmission in neotropical NHPs

There are more than 190 species and subspecies of nonhuman primates in South America, of which a large number live in the Amazon Basin in proximity in some cases (e.g., Saimiri) to human activities⁵⁴. This diversity underscores the need for data on their susceptibility to ZIKV infection to predict potential future circulation in an enzootic cycle as well as re-emergence into the human transmission cycle.

In 2015, a serosurvey conducted in diverse ecotypes from Ceará State, Brazil, which at the time was a ZIKV-epidemic area, sampled sera and oral swabs from 15 marmosets (*C. jacchus*) and 9 capuchin monkeys (*Sapajus libidinosus*). Four ZIKV-positive samples from marmosets and three positive samples from capuchin monkeys were identified (Supplementary Table 3). Subsequent sequencing of amplified products indicated that they were identical to ZIKV strains circulating epidemically at the time, suggesting that they could serve as reservoirs for enzootic transmission. In addition, recent experimental ZIKV infections of marmosets reproduced key features of human ZIKV infection and viremia titers were near the necessary threshold for transmission ^{55,56}.

Another study characterized the dynamics of ZIKV infection in neotropical squirrel (*Saimiri* spp.) and owl monkeys (*Aotus* spp.)⁵⁷ (Supplementary Table 3). Viremia was observed in both species in the absence of detectible disease, and seroconversion occurred by day 28. This experimental study confirms the susceptibility to ZIKV infection of two New World NHP species living in close

proximity to humans in South America, suggesting that they could support a neotropical enzootic ZIKV cycle. However, it remains to be experimentally validated whether the observed viremia is sufficient to infect sympatric sylvatic mosquitoes (host competence) and whether the New World arboreal enzootic mosquitoes have the ability to become infected with ZIKV and transmit upon subsequent feedings (vector competence) to susceptible NHPs.

Zika virus recent emergence and evolution

Evolutionary studies of recent ZIKV spread. The first direct phylogenetic studies of ZIKV were reported in 2012, when Haddow et al. 58 analysed the genomic sequences of six ZIKV strains and found that they grouped by geographic origin, suggesting that African and Asian lineages diverged over long periods of isolation. Because the closest relative, Spondweni virus, is found only in Africa, the authors concluded that ZIKV originated on that continent. More extensive analyses of ZIKV strains dating from 1968 to 2002, collected in several locations of Africa, suggested at least two independent introductions into West Africa during the 20th century and the possibility of recombination there 18. However, the extremely limited ZIKV sampling, especially in East Africa since the 1950s, suggests that sampling bias affects such phylogeographic analyses.

Since the arrival of ZIKV in French Polynesia and the Americas, major phylogenetic analyses using much larger strain numbers have provided detailed understanding of patterns of its spread and evolution. ZIKV was introduced from somewhere in Southeast Asia into French Polynesia, which was followed by the spread both eastward and westward in the South Pacific, presumably by infected air travellers. Introduction into the Americas from French Polynesia is estimated to have occurred in late 2013⁵⁹, followed by a period of undetected circulation in Northeastern Brazil in early 2014. Dissemination to other parts of the Americas occurred subsequently before the first Brazilian outbreak was detected in early 201560 (Fig. 1). The rapid spread throughout the South American continent seems to have been enhanced by the warm temperatures associated with the 2015 appearance of El Niño, perhaps through stimulation of mosquito biting rates and shortening of the ZIKV extrinsic incubation period⁶¹. The subsequent explosive spread of ZIKV throughout the Americas facilitated its spread eastward, reaching Cape Verde in late 201562. In addition, ZIKV cases have been detected in Guinea Bissau⁶³, Angola⁶⁴, Senegal and Nigeria⁶⁵, but the lineage of the concerned strains has not been yet determined. The most detailed study of ZIKV introduction and spread into new territories comes from Florida in the United States, where an estimated 4-40 introductions (mainly from the Caribbean) seeded local transmission that probably began in the spring of 201666.

Emergence mechanisms of urban ZIKV transmission. The broad distribution and abundance of *Aedes* spp. vectors (especially domestic *Ae. aegypti*)²⁵, coupled with the inefficacy of traditional vector control measures, are key elements in ZIKV emergence⁶⁷. Poverty and inadequate or damaged infrastructure, deforestation and urbanization are factors that favour mosquito larval development sites, human–mosquito contacts and thereby ZIKV transmission^{68,69}. In addition, human movement and intercontinental air travel has expedited the spread of vector-borne diseases^{68,70–72}. Nevertheless, the impact of these socioeconomic and environmental factors on vector-borne transmission is a common feature for all emerging mosquito-borne viruses, and they do not explain the absence of ZIKV urban epidemics before 2007.

An intuitive explanation for the emergence of urban ZIKV transmission and congenital Zika syndrome was a change in ZIKV phenotypes coincident with its spread into the South Pacific and Americas. This hypothesis was suggested following the observed vector-adaptive evolution of the Indian Ocean lineage of chikungunya virus, which allowed, for the first time, exploitation of

Ae. albopictus as a major urban vector via a series of envelope glycoprotein substitutions that enhance transmission by this species⁷³. Could a similar phenomenon explain the massive urban amplification and spread of ZIKV? Alternatively, could the pathogenicity of ZIKV have changed coincident with this spread, for example, selected by increased viremia for transmission by peridomestic mosquitoes, which might also enhance fetal infection? Comprehensive sequencing and phylogenetic studies like those above have also been used to identify recent mutations in the ZIKV genome that might be associated with altered pathogenicity or urban vector transmission to explain the explosive nature of recent outbreaks and the lack of evidence for such epidemics in the past. Two approaches have been used to determine whether such mutations, several of which were traced to ZIKV spread into the South Pacific and the Americas⁷⁴, could have encoded critical phenotypes responsible for epidemic emergence and congenital disease.

Hypothetical emergence mechanism 1: adaptation for transmission by urban vectors. Urban vector-adaptive evolution has been identified as a major factor in the emergence of two other viruses with origins in nonhuman primate-arboreal mosquito vector enzootic cycles: dengue and chikungunya viruses. Although no overall increase in urban vector transmission efficiency has been linked to emergence of the human transmission cycle⁷⁵, the Asian lineage of dengue virus-2 (DENV-2) evolved for more efficient transmissibility by *Ae. aegypti*⁷⁶, and this phenotype has been linked to its replacement of the less transmissible and less human-virulent American DENV-2 genotype⁷⁷.

A mechanism for subgenomic flavivirus RNA (sfRNA)-mediated enhancement of flavivirus transmission by mosquitoes has also been suggested⁷⁸. Substitutions in the 3' UTR (untranslated region) from the DENV genome have led to higher amounts of sfRNA in *Ae. aegypti* salivary glands, which resulted in increased DENV infection rates and loads, as well as suppression of mosquito immune response in in this latter organ⁷⁹. Similar results have been obtained using the West Nile virus-*Culex pipiens* model⁸⁰, suggesting a potential role of sfRNA produced by the host metabolic machinery as a driver for ZIKV transmission.

The most striking example of vector-adaptive evolution by an arbovirus is CHIKV repeated emergence events and spread since 2004. One explanation for this dramatic spread is the adaptation via a series of mutations⁸¹ of one CHIKV strain, the Indian Ocean Lineage, for efficient midgut infection and transmission by the highly invasive mosquito *Ae. albopictus*, never before implicated as an important vector. Surprisingly, these mutations have little or no impact on infection of the primary vector *Ae. aegypti* in most locations and do not affect infection of *Ae. albopictus* in the genetic backbone of CHIKV strains circulating in the Americas⁷³.

The first evidence of potential Ae. aegypti-adaptive ZIKV evolution came from work examining the flavirirus nonstructural glycoprotein NS1 substitution that enhances infection of this species82. Starting with natural isolates that differ by an alanine versus a valine residue at position 188 showed that a strain encoding valine imported to China from the Americas generated higher levels of NS1 in the blood of infected mice as well as in the supernatant of infected cell lines. The strain encoding V188 also infects the Rockefeller colony of Ae. aegypti more efficiently, and NS1 spiked into artificial bloodmeals recapitulates this finding. However, other studies with ZIKV strains differing in these NS1 residues have not revealed comparable differences^{83,84}. Additional studies with wild mosquito populations (the Rockefeller colony is many decades old, and colonization is known to affect the susceptibility of mosquitoes to arboviruses) are needed to further evaluate the importance of NS1 change. Interestingly, the V188 residue is also present in African ZIKV strains, suggesting it was lost upon the introduction into Asia many decades ago, possibly the result of a founder effect.

Hypothetical emergence mechanism 2: ZIKV adaptive evolution in vertebrate hosts. Vertebrate-adaptive evolution has been identified as a possible factor in the emergence of DENV. Southeast Asian genotype of serotype 2 and genotype III of serotype 3 have been associated with higher viremias in humans, leading to severe dengue disease and higher transmission rates by susceptible Aedes spp. mosquitoes^{77,85-88}. Duplication of an adaptable RNA element has been suggested as a possible mechanism of sustaining high viral fitness for a virus that cycles between hosts89. However, when mechanisms of human DENV emergence due to host range expansion of sylvatic strains by adaptation to use humans as reservoirs (via increased magnitude of replication) were evaluated in two surrogate models of human infection, there were no significant differences, suggesting that the historical emergence of DENV from the ancestral sylvatic transmission cycle into human cycles may not have required adaptation to replicate in humans as reservoir hosts90. In a similar manner, the rate of evolutionary change and pattern of natural selection are similar among endemic and sylvatic DENV, suggesting that the dynamics of mutation, replication and selection are broadly equivalent for DENV across its host range⁹¹, and thus the potential for additional zoonotic virus emergence into the human transmission cycle is high.

The explosive global spread of ZIKV has been hypothesized to be attributed to adaptive evolution for efficient urban transmission in humans. A number of phylogenetic studies^{59,92} have suggested that adaptive evolution may have occurred in Southeast Asia. The Asian ZIKV lineage may have adapted to generate higher viremia levels in humans, leading to increased efficiencies in mosquito infection, transmission and spread. Higher viremia could then enhance transplacental transmission in humans, which could explain the dramatic emergence of microcephaly in the Americas. Some bioinformatic studies suggested an increase in the use of human codons by the virus, which may support the notion of adapted evolution in humans 18,93,94. In addition, one study 5 showed that a single serine-to-asparagine substitution (Ser17→Asn17 (S17N))in the ZIKV structural protein prM increased infectivity in both human and mouse neural progenitor cells, leading to more severe microcephaly in the mouse fetus. However, this potential link will require comprehensive longitudinal studies in humans and/or animal models, which will be difficult to test, because various animal models may not respond to ZIKV infection in the same manner as humans. To date, the lack of solid evidence of adaptive evolution in humans suggests that a combination of stochastic factors and selective evolution may have contributed to ZIKV's global emergence and spread.

Conclusion

Ae. aegypti is the principal vector in urban ZIKV transmission worldwide, whereas other broadly distributed Aedes species may act as ZIKV vectors in specific environments where their abundance is important. However, information about ZIKV titers in mosquitoes consistent with transmission competence is still scarce, resulting in difficulty to assess their specific roles in transmission. Another important knowledge gap that deserves more investigation concerns the contribution of vertical and venereal transmission in mosquito–human transmission.

The NHPs are the most important enzootic amplification hosts for ZIKV maintenance in the sylvatic cycle, although other orders of animal species appear to be susceptible to ZIKV infection. However, there is scarce information on the viremia levels that these potential hosts can develop, which limits complete comprehension of their contribution to ZIKV transmission. In addition, the knowledge of sylvatic species competence (both vectors and hosts) to transmit ZIKV in the Americas and Asia is still limited. Despite these gaps, the evidence collected in the wild and in the lab suggest a high probability of establishment of enzootic ZIKV in the Americas, which could make impossible its eradication from the region. Finally, the

conducted studies regarding ZIKV-adaptive evolution in vectors and vertebrate hosts suggest that a combination of stochastic factors and Darwinian evolution may have contributed to ZIKV's global emergence and spread.

Received: 5 June 2018; Accepted: 4 February 2019; Published online: 18 March 2019

References

- Dick, G. W. A. Zika virus. II. Pathogenicity and physical properties. Trans. R. Soc. Trop. Med. Hyg. 46, 521–534 (1952).
- Pan American Health Organization. Regional Zika Epidemiological Update (Americas) August 25, 2017 (2017).
- Martinez-Pulgarin, D. F., Acevedo-Mendoza, W. F., Cardona-Ospina, J. A., Rodríguez-Morales, A. J. & Paniz-Mondolfi, A. E. A bibliometric analysis of global Zika research. *Travel Med. Infect. Dis.* 14, 55–57 (2016).
- Musso, D. & Gubler, D. J. Zika Virus. Clin. Microbiol. Rev. 29, 487–524 (2016).
- Boyer, S., Calvez, E., Chouin-Carneiro, T., Diallo, D. & Failloux, A. B. An overview of mosquito vectors of Zika virus. *Microbes Infect.* 20, 646–660 (2018).
- Grard, G. et al. Zika virus in Gabon (Central Africa) 2007: a new threat from Aedes albopictus? PLoS Negl. Trop. Dis. 8, e2681 (2014).
- Ledermann, J. P. et al. Aedes hensilli as a potential vector of Chikungunya and Zika viruses. PLoS Negl. Trop. Dis. 8, e3188 (2014).
- Duffy, M. R. et al. Zika virus outbreak on Yap Island, Federated States of Micronesia. N. Engl. J. Med. 360, 2536–2543 (2009).
- Oehler, E. et al. Zika virus infection complicated by Guillain-Barre syndrome-case report, French Polynesia, December 2013. Euro Surveill. 19, 7–9 (2014).
- World Health Organisation. Zika Virus (ZIKV) Classification Table 15th February. Who 1–2 (2018).
- Marchette, N. J., Garcia, R. & Rudnick, A. Isolation of Zika virus from *Aedes aegypti* mosquitoes in Malaysia. *Am. J. Trop. Med. Hyg.* 18, 411–415 (1969).
- Singapore Zika Study Group. Outbreak of Zika virus infection in Singapore: an epidemiological, entomological, virological, and clinical analysis. *Lancet Infect. Dis.* 17, 813–821 (2017).
- Mutebi, J. P. et al. Zika virus MB16-23 in mosquitoes, Miami-Dade County, Florida, USA, 2016. Emerg. Infect. Dis. 24, 808-810 (2018).
- Guerbois, M. et al. Outbreak of Zika virus infection, Chiapas State, Mexico, 2015, and first confirmed transmission by Aedes aegypti mosquitoes in the Americas. J. Infect. Dis. 214, 1349–1356 (2016).
- Guedes, D. R. D. et al. Zika virus replication in the mosquito Culex quinquefasciatus in Brazil. Emerg. Microbes Infect. 6, e69 (2017).
- Ferreira-de-Brito, A. et al. First detection of natural infection of Aedes aegypti with Zika virus in Brazil and throughout South America. Mem. Inst. Oswaldo Cruz 111, 655-658 (2016).
- Ayllón, T. et al. Early evidence for zika virus circulation among Aedes aegypti mosquitoes, Rio de Janeiro, Brazil. Emerg. Infect. Dis. 23, 1411–1412 (2017).
- Faye, O. et al. Molecular evolution of Zika virus during its emergence in the 20th century. PLoS Negl. Trop. Dis. 8, e2636 (2014).
- Akoua-Koffi, C. et al. Investigation autour d'un cas mortel de fièvre jaune en Côte d'Ivoire en 1999. Bull Soc. Pathol. Exot. 94, 227–230 (2001).
- Elizondo-Quiroga, D. et al. Zika virus in salivary glands of five different species of wild-caught mosquitoes from Mexico. Sci. Rep. 8, 809 (2018).
- Yee, D. A., Dejesus-Crespo, R., Hunter, F. F. & Bai, F. Assessing natural infection with Zika virus in the southern house mosquito, *Culex quinquefasciatus*, during 2016 in Puerto Rico. *Med. Vet. Entomol.* 32, 255–258 (2017).
- Smartt, C. T. et al. Evidence of Zika virus RNA fragments in Aedes albopictus (Diptera: Culicidae) field-collected eggs from Camaçari, Bahia, Brazil. J. Med. Entomol. 54, 1085–1087 (2017).
- 23. Song, S. et al. Could Zika virus emerge in Mainland China? Virus isolation from nature in *Culex quinquefasciatus*, 2016. *Emerg. Microbes Infect.* **6**, e93 (2017)
- Díaz-Quiñonez, J. A. et al. Evidence of the presence of the Zika virus in Mexico since early 2015. Virus Genes 52, 855–857 (2016).
- Kraemer, M. U. G. et al. The global distribution of the arbovirus vectors *Aedes aegypti* and *Ae. albopictus. eLife* 4, e08347 (2015).
- Ritchie, S. A. In *Dengue and Dengue Hemorrhagic Fever*, 2nd Edition. (eds Gubler, D. J., Ooi, E. E., Vasudevan, S. & Farrar, J.) 455–480 (CAB International, 2014).
- 27. Musso, D., Nilles, E. J. & Cao-Lormeau, V. M. Rapid spread of emerging Zika virus in the Pacific area. *Clin. Microbiol. Infect.* **20**, O595–O596 (2014).
- Magalhaes, T., Foy, B. D., Marques, E. T. A., Ebel, G. D. & Weger-Lucarelli, J. Mosquito-borne and sexual transmission of Zika virus: recent developments and future directions. Virus Res. 254, 1–9 (2018).

- Ioos, S. et al. Current Zika virus epidemiology and recent epidemics. Med. Mal. Infect. 44, 302–307 (2014).
- Richard, V., Paoaafaite, T. & Cao-Lormeau, V. M. Vector competence of French Polynesian Aedes aegypti and Aedes polynesiensis for Zika virus. PLoS Negl. Trop. Dis. 10, e0005024 (2016).
- 31. Guo, X. X. et al. Culex pipiens quinquefasciatus: a potential vector to transmit Zika virus. Emerg. Microbes Infect. 5, e102 (2016).
- Smartt, C. T., Shin, D., Kang, S. & Tabachnick, W. J. Culex quinquefasciatus (Diptera: Culicidae) From Florida transmitted Zika virus. Front. Microbiol. 9, 768 (2018).
- Day, J. F. Predicting St. Louis encephalitis virus epidemics: lessons from recent, and not so recent, outbreaks. Annu. Rev. Entomol. 46, 111–138 (2001).
- Diallo, D. et al. Zika virus emergence in mosquitoes in southeastern Senegal, 2011. PLoS One 9, e109442 (2014).
- Miller, B. R., Monath, T. P., Tabachnick, W. J. & Ezike, V. I. Epidemic yellow fever caused by an incompetent mosquito vector. *Trop. Med. Parasitol.* 40, 396–399 (1989).
- Fernandes, R. S. et al. Culex quinquefasciatus from Rio de Janeiro is not competent to transmit the local Zika virus. PLoS Negl. Trop. Dis. 10, e0004993 (2016).
- Garcia-Rejon, J. E. et al. Host-feeding preference of the mosquito, Culex quinquefasciatus, in Yucatan State, Mexico. J. Insect Sci. 10, 32 (2010).
- Lequime, S., Paul, R. E. & Lambrechts, L. Determinants of arbovirus vertical transmission in mosquitoes. *PLoS Pathog.* 12, e1005548 (2016).
- Althouse, B. M. et al. Impact of climate and mosquito vector abundance on sylvatic arbovirus circulation dynamics in Senegal. Am. J. Trop. Med. Hyg. 92, 88–97 (2015).
- Thangamani, S., Huang, J., Hart, C. E., Guzman, H. & Tesh, R. B. Vertical transmission of Zika virus in *Aedes aegypti* mosquitoes. *Am. J. Trop. Med. Hyg.* 95, 1169–1173 (2016).
- Li, C. X. et al. Vector competence and transovarial transmission of two Aedes aegypti strains to Zika virus. Emerg. Microbes Infect. 6, e23 (2017).
- Ciota, A. T., Bialosuknia, S. M., Ehrbar, D. J. & Kramer, L. D. Vertical transmission of Zika virus by Aedes aegypti and Ae. albopictus mosquitoes. Emerg. Infect. Dis. 23, 880–882 (2017).
- 43. Campos, S. S. et al. Zika virus can be venereally transmitted between *Aedes aegypti* mosquitoes. *Parasit. Vectors* **10**, 605 (2017).
- Pereira-Silva, J. W. et al. First evidence of Zika virus venereal transmission in Aedes aegypti mosquitoes. Mem. Inst. Oswaldo Cruz 113, 56–61 (2018).
- Althouse, B. M. et al. Potential for Zika virus to establish a sylvatic transmission cycle in the Americas. PLoS Negl. Trop. Dis. 10, e0005055 (2016).
- Favoretto, S. et al. First detection of Zika virus in neotropical primates in Brazil: a possible new reservoir. bioRxiv https://doi.org/10.1101/049395 (2016).
- Darwish, M. A., Hoogstraal, H., Roberts, T. J., Ahmed, I. P. & Omar, F. A sero-epidemiological survey for certain arboviruses (Togaviridae) in Pakistan. *Trans. R. Soc. Trop. Med. Hyg.* 77, 442–445 (1983).
- Olson, J. G. et al. A survey for arboviral antibodies in sera of humans and animals in Lombok, Republic of Indonesia. *Ann. Trop. Med. Parasitol.* 77, 131–137 (1983).
- Joob, B. & Wiwanitkit, V. Immunological reactive rate to Zika virus in canine sera: a report from a tropical area and concern on pet, zoonosis and reservoir host. Asian Pac. J. Trop. Med. 10, 208–209 (2017).
- Bryant, J. E., Holmes, E. C. & Barrett, A. D. T. Out of Africa: a molecular perspective on the introduction of yellow fever virus into the Americas. *PLoS Pathog.* 3, e75 (2007).
- Volk, S. M. et al. Genome-scale phylogenetic analyses of chikungunya virus reveal independent emergences of recent epidemics and various evolutionary rates. J. Virol. 84, 6497–6504 (2010).
- Vasilakis, N., Cardosa, J., Hanley, K. A., Holmes, E. C. & Weaver, S. C. Fever from the forest: prospects for the continued emergence of sylvatic dengue virus and its impact on public health. *Nat. Rev. Microbiol.* 9, 532–541 (2011).
- 53. Hanley, K. A. et al. Fever versus fever: the role of host and vector susceptibility and interspecific competition in shaping the current and future distributions of the sylvatic cycles of dengue virus and yellow fever virus. *Infect. Genet. Evol.* 19, 292–311 (2013).
- 54. Rylands, A. B. & Mittermeier, R. A. in South American Primates, Developments in Primatology: Progress and Prospects (eds Garber, P.A., Estrada, A., Bicca-Marques, J. C., Hetmann, E.W. & Strier, K.B.) 23–55 (Springer, 2009).
- 55. Seferovic, M. et al. Experimental Zika virus infection in the pregnant common marmoset induces spontaneous fetal loss and neurodevelopmental abnormalities. *Sci. Rep.* **8**, 6851 (2018).
- Chiu, C. Y. et al. Experimental Zika virus inoculation in a new world monkey model reproduces key features of the human infection. Sci. Rep. 7, 17126 (2017).
- Vanchiere, J. A. et al. Experimental Zika virus infection of neotropical primates. Am. J. Trop. Med. Hyg. 98, 173–177 (2018).

- Haddow, A. D. et al. Genetic characterization of Zika virus strains: geographic expansion of the Asian lineage. PLoS Negl. Trop. Dis. 6, e1477 (2012).
- Faria, N. R. et al. Zika virus in the Americas: early epidemiological and genetic findings. Science 352, 345–349 (2016).
- Faria, N. R. et al. Establishment and cryptic transmission of Zika virus in Brazil and the Americas. *Nature* 546, 406–410 (2017).
- Caminade, C. et al. Global risk model for vector-borne transmission of Zika virus reveals the role of El Niño 2015. *Proc. Natl Acad. Sci. USA* 114, 119–124 (2017).
- Lourenço, J. et al. Epidemiology of the Zika virus outbreak in the Cabo Verde Islands, West Africa. PLoS Curr. https://doi.org/10.1371/currents.outbreaks.19 433b1e4d007451c691f138e1e67e8c (2018)
- 63. Rosenstierne, M. W. et al. Zika virus IgG in infants with microcephaly, Guinea-Bissau, 2016. *Emerg. Infect. Dis.* 24, 948–950 (2018).
- Kraemer, M. U. G. et al. Zika virus transmission in Angola and the potential for further spread to other African settings. *Trans. R. Soc. Trop. Med. Hyg.* 111, 527–529 (2017).
- Herrera, B. B. et al. Continued transmission of Zika virus in Humans in West Africa, 1992–2016. J. Infect. Dis. 215, 1546–1550 (2017).
- Grubaugh, N. D. et al. Genomic epidemiology reveals multiple introductions of Zika virus into the United States. *Nature* 546, 401–405 (2017).
- Fernández-Salas, I. et al. Historical inability to control Aedes aegypti as a main contributor of fast dispersal of chikungunya outbreaks in Latin America. Antiviral Res. 124, 30–42 (2015).
- Ali, S. et al. Environmental and social change drive the explosive emergence of Zika virus in the Americas. PLoS Negl. Trop. Dis. 11, e0005135 (2017).
- 69. Reina Ortiz, M. et al. Post-earthquake Zika virus surge: disaster and public health threat amid climatic conduciveness. Sci. Rep. 7, 15408 (2017).
- Harrington, L. C. et al. Dispersal of the dengue vector Aedes aegypti within and between rural communities. Am. J. Trop. Med. Hyg. 72, 209–220 (2005).
- 71. Stoddard, S. T. et al. House-to-house human movement drives dengue virus transmission. *Proc. Natl Acad. Sci. USA* 110, 994–999 (2013).
- Bogoch, I. I. et al. Potential for Zika virus introduction and transmission in resource-limited countries in Africa and the Asia-Pacific region: a modelling study. *Lancet Infect. Dis.* 16, 1237–1245 (2016).
- 73. Tsetsarkin, K. A., Chen, R. & Weaver, S. C. Interspecies transmission and chikungunya virus emergence. *Curr. Opin. Virol.* 16, 143–150 (2016).
- 74. Pettersson, J. H. O. et al. How did zika virus emerge in the Pacific Islands and Latin America? *MBio* 7, 1–6 (2016).
- Vasilakis, N. et al. Sylvatic dengue viruses share the pathogenic potential of urban/endemic dengue viruses. J. Virol. 84, 3726–3728 (2010).
- Armstrong, P. M. & Rico-Hesse, R. Differential susceptibility of *Aedes aegypti* to infection by the American and Southeast Asian genotypes of dengue type 2 virus. *Vector Borne Zoonotic Dis.* 1, 159–168 (2001).
- Cologna, R., Armstrong, P. M. & Rico-Hesse, R. Selection for virulent dengue viruses occurs in humans and mosquitoes. J. Virol. 79, 853–859 (2005).
- 78. Yeh, S. C. & Pompon, J. Flaviviruses produce a subgenomic flaviviral RNA that enhances mosquito transmission. *DNA Cell Biol.* **37**, 154–159 (2017).
- Pompon, J. et al. Dengue subgenomic flaviviral RNA disrupts immunity in mosquito salivary glands to increase virus transmission. *PLoS Pathog.* 13, e1006535 (2017).
- Göertz, G. P. et al. Noncoding subgenomic flavivirus RNA is processed by the mosquito RNA interference machinery and determines West Nile Virus transmission by *Culex pipiens* mosquitoes. *J. Virol.* 90, 10145–10159 (2016).
- Tsetsarkin, K. A. et al. Multi-peaked adaptive landscape for chikungunya virus evolution predicts continued fitness optimization in *Aedes albopictus* mosquitoes. *Nat. Commun.* 5, 4084 (2014).
- 82. Liu, Y. et al. Evolutionary enhancement of Zika virus infectivity in *Aedes aegypti* mosquitoes. *Nature* **545**, 482–486 (2017).
- Weger-Lucarelli, J. et al. Vector competence of American mosquitoes for three strains of Zika virus. PLoS Negl. Trop. Dis. 10, e0005101 (2016).
- Roundy, C. M. et al. Variation in Aedes aegypti mosquito competence for Zika virus transmission. Emerg. Infect. Dis. 23, 625–632 (2017).
- Libraty, D. H., Pichyangkul, S., Ajariyakhajorn, C., Endy, T. P. & Ennis, F. A. Human dendritic cells are activated by dengue virus infection: enhancement by gamma interferon and implications for disease pathogenesis. *J. Virol.* 75, 3501–3508 (2001).
- Vaughn, D. W. et al. Dengue viremia titer, antibody response pattern, and virus serotype correlate with disease severity. J. Infect. Dis. 181, 2–9 (2000).
- 87. Messer, W. B., Gubler, D. J., Harris, E., Sivananthan, K. & de Silva, A. M. Emergence and global spread of a dengue serotype 3, subtype III virus. *Emerg. Infect. Dis.* 9, 800–809 (2003).
- Wang, W. K. et al. High levels of plasma dengue viral load during defervescence in patients with dengue hemorrhagic fever: implications for pathogenesis. *Virology* 305, 330–338 (2003).
- Villordo, S. M., Filomatori, C. V., Sánchez-Vargas, I., Blair, C. D. & Gamarnik, A. V. Dengue virus RNA structure specialization facilitates host adaptation. *PLoS Pathog.* 11, e1004604 (2015).

- 90. Vasilakis, N. et al. Potential of ancestral sylvatic dengue-2 viruses to re-emerge. *Virology* **358**, 402–412 (2007).
- 91. Vasilakis, N. et al. Evolutionary processes among sylvatic dengue type 2 viruses. *J. Virol.* **81**, 9591–9595 (2007).
- 92. Weaver, S. C. et al. Zika virus: history, emergence, biology, and prospects for control. *Antiviral Res.* 130, 69–80 (2016).
- 93. Wang, H., Liu, S., Zhang, B. & Wei, W. Analysis of synonymous codon usage bias of zika virus and its adaption to the hosts. *PLoS One* 11, e0166260 (2016).
- Butt, A. M., Nasrullah, I., Qamar, R. & Tong, Y. Evolution of codon usage in Zika virus genomes is host and vector specific. *Emerg. Microbes Infect.* 5, e107 (2016).
- 95. Yuan, L. et al. A single mutation in the prM protein of Zika virus contributes to fetal microcephaly. *Science* **358**, 933–936 (2017).

Acknowledgements

This work has been notably supported by the Special Programme for Research and Training in Tropical Diseases (TDR) from the World Health Organization (Grant 2017/713047-0). The European Union's Horizon 2020 Research and innovation programme under "ZIKALLIANCE" (Grant Agreement no. 734548) partially supported A.V.-R. and R.L.d.O. The Programme Opérationnel FEDER-Guadeloupe-Conseil

Régional 2014- 2020 (Grant 2015-FED-192) partially supported A.V.-R. NIH grants R24 AI120942 and U01 AI115577 provided a partial support for N.V. and S.C.W.

Author contributions

G.G.-B., L.A.P., M.R., J.A.B., R.L.d.O., S.C.W., N.V. and A.V.-R. participated in data acquisition, analysis and interpretation. G.G.-B., A.V.-R., S.C.W. and N.V. wrote the paper. A.V.-R. conceived the study. All authors revised and approved the final manuscript.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information is available for this paper at https://doi.org/10.1038/s41559-019-0836-z.

Reprints and permissions information is available at www.nature.com/reprints.

Correspondence should be addressed to A.V.-R.

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

© Springer Nature Limited 2019