



HAL
open science

Zika vector competence data reveals risks of outbreaks: the contribution of the European ZIKAlliance project

Thomas Obadia, Gladys Gutierrez-Bugallo, Veasna Duong, Ana Nuñez,
Rosilainy Fernandes, Basile Kamgang, Liza Hery, D. Jiolle, Célestine Atyame,
Yann Gomard, et al.

► To cite this version:

Thomas Obadia, Gladys Gutierrez-Bugallo, Veasna Duong, Ana Nuñez, Rosilainy Fernandes, et al..
Zika vector competence data reveals risks of outbreaks: the contribution of the European ZIKAlliance
project. *Nature Communications*, 2022, 13 (1), pp.4490. 10.1038/s41467-022-32234-y . pasteur-
03744692

HAL Id: pasteur-03744692

<https://pasteur.hal.science/pasteur-03744692>

Submitted on 3 Aug 2022

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



Distributed under a Creative Commons Attribution 4.0 International License

Zika vector competence data reveals risks of outbreaks: the contribution of the European ZIKAlliance project

Received: 5 April 2022

Accepted: 18 July 2022

Published online: 02 August 2022

 Check for updates

Thomas Obadia^{1,2}, Gladys Gutierrez-Bugallo ^{3,4}, Veasna Duong ⁵, Ana I. Nuñez⁶, Rosilainy S. Fernandes ⁷, Basile Kamgang ⁸, Liza Hery⁴, Yann Gomard ⁹, Sandra R. Abbo ¹⁰, Davy Jiolle¹¹, Uros Glavinic¹², Myrielle Dupont-Rouzeyrol¹³, Célestine M. Atyame⁹, Nicolas Pocquet¹⁴, Sébastien Boyer ¹⁵, Catherine Dauga¹⁶, Marie Vazeille ¹⁶, André Yébakima¹⁷, Michael T. White ², Constantianus J. M. Koenraadt¹⁸, Patrick Mavingui⁹, Anubis Vega-Rua⁴, Eva Veronesi¹², Gorben P. Pijlman ¹⁰, Christophe Paupy¹¹, Núria Busquets⁶, Ricardo Lourenço-de-Oliveira ⁷, Xavier De Lamballerie ¹⁹ & Anna-Bella Failloux ¹⁶ 

First identified in 1947, Zika virus took roughly 70 years to cause a pandemic unusually associated with virus-induced brain damage in newborns. Zika virus is transmitted by mosquitoes, mainly *Aedes aegypti*, and secondarily, *Aedes albopictus*, both colonizing a large strip encompassing tropical and temperate regions. As part of the international project ZIKAlliance initiated in 2016, 50 mosquito populations from six species collected in 12 countries were experimentally infected with different Zika viruses. Here, we show that *Ae. aegypti* is mainly responsible for Zika virus transmission having the highest susceptibility to viral infections. Other species play a secondary role in transmission while *Culex* mosquitoes are largely non-susceptible. Zika strain is expected to significantly modulate transmission efficiency with African strains being more likely to cause an outbreak. As the distribution of *Ae. aegypti* will doubtless expand with climate change and without new marketed vaccines, all the ingredients are in place to relive a new pandemic of Zika.

¹Institut Pasteur, Université Paris Cité, Bioinformatics and Biostatistics Hub, F-75015 Paris, France. ²Institut Pasteur, Université Paris Cité, G5 Infectious Disease Epidemiology and Analytics, F-75015 Paris, France. ³Department of Vector Control, Center for Research, Diagnostic, and Reference, Institute of Tropical Medicine Pedro Kouri, Havana, Cuba. ⁴Institut Pasteur de Guadeloupe, Laboratory of Vector Control Research, Unit Transmission Reservoir and Pathogens Diversity, Les Abymes, Guadeloupe. ⁵Institut Pasteur du Cambodge, Virology Unit, Phnom Penh, Cambodia. ⁶IRTA, Centre de Recerca en Sanitat Animal (CRESA, IRTA-UAB), Campus de la Universitat Autònoma de Barcelona, 08193 Bellaterra, Spain. ⁷Laboratório de Mosquitos Transmissores de Hematozoários, Instituto Oswaldo Cruz, Fiocruz, Rio de Janeiro, RJ, Brazil. ⁸Centre for Research in Infectious Diseases, Department of Medical Entomology, Yaoundé, Cameroon. ⁹UMR PIMIT (Processus Infectieux en Milieu Insulaire Tropical), Sainte-Clotilde, La Réunion, France. ¹⁰Laboratory of Virology, Wageningen University, Wageningen, The Netherlands. ¹¹IRD, MIVEGEC, University of Montpellier, IRD, CNRS, Montpellier, France. ¹²National Centre for Vector Entomology, Institute of Parasitology, Vetsuisse Faculty, University of Zürich, Zürich, Switzerland. ¹³Institut Pasteur de Nouvelle-Calédonie, URE Dengue et Arboviroses, Nouméa, New Caledonia. ¹⁴Institut Pasteur de Nouvelle-Calédonie, URE Entomologie Médicale, Nouméa, New Caledonia. ¹⁵Institut Pasteur du Cambodge, Medical Entomology Unit, Phnom Penh, Cambodia. ¹⁶Institut Pasteur, Université Paris Cité, Arboviruses and Insect Vectors, F-75015 Paris, France. ¹⁷VECCOTRA, Rivière Salée, Martinique. ¹⁸Laboratory of Entomology, Wageningen University & Research, Wageningen, The Netherlands. ¹⁹Unité des Virus Emergents (UVE), Aix Marseille Université, IHU Méditerranée Infection, Marseille, France. ✉e-mail: anna-bella.failloux@pasteur.fr

Viral pathogens with high epidemic potential have marked human history by massive losses of life and economic declines. Pandemics like the Spanish flu¹ have left traces and the fear of new viral emergences materializes today with the Severe Acute Respiratory Syndrome-Coronavirus-2 (SARS-CoV-2) pandemic. Over the past decades, efforts invested into vaccination programs and development of antiviral treatments have led to major medical progress resulting in measles decline, smallpox eradication, appropriate treatments of Human Immunodeficiency virus and Hepatitis C virus, and improved control of SARS-CoV-2. As major viral pathogens, arthropod-borne viruses (arboviruses) are also on the rise. Arboviruses have developed a strategy to accomplish their transmission cycle *via* replication in both vertebrate and arthropod hosts. Their RNA genome contributes to the generation of large populations of genetically distinct variants permitting adaptations to environmental changes including host switching and transmission². In the last 30 years, mosquito-borne viruses have dramatically expanded their distribution range leading to increasingly frequent and large epidemics³. In 2016, Zika was designated as a “Public Health Emergency of International Concern” by the World Health Organization, subsequently to the large spread of Zika virus (ZIKV) in the Pacific islands and the Americas with unusual notifications of microcephaly in newborns and different neurological disorders (e.g. Guillain-Barré syndrome); this happened 70 years after its first isolation from a human case in East Africa⁴. Local cases of Zika were detected in 2019 in Southern France additionally corroborating the role of the Asian tiger mosquito (*Aedes albopictus*) in ZIKV transmission in addition to chikungunya and dengue viruses⁵.

Aedes mosquitoes are important vectors of arboviruses, acquiring the virus through a blood meal from an infected host. Subsequently, the virus replicates within the mosquito and is transmitted to a new host. The virus must overcome two anatomical barriers (midgut and salivary glands) and circumvent major antiviral pathways before being delivered in saliva⁶. Geographic populations of a same mosquito species may not share the same immunological background, leading to different vector competences⁷. The outcome of infection depends on specific pairings of mosquito and pathogen genotypes described under genotype-by-genotype (G x G) interactions⁸. These interactions could structure viral populations through adaptations to their local vector populations⁸. Vector competence data have been used to derive risk maps informed with distribution of mosquito species, to help focus and strengthen vector control measures⁹. Nevertheless, other factors, non-genetic, biotic and abiotic, should be considered in the functioning of the vectorial system making it even more complex than initially thought¹⁰. A global vector competence analysis of potential Zika vectors (*Aedes aegypti*, *Ae. albopictus*, *Aedes japonicus*, *Culex pipiens pipiens*, *Culex pipiens molestus*, and *Culex quinquefasciatus*) was performed to assess the risk of Zika disease outbreaks. Once the vector is well established in a region, associated arboviruses have the potential to emerge through imported human cases. To this aim, we analyzed the vector competence of 50 mosquito populations from 15 locations and 12 countries (Brazil, Cambodia, Cameroon, Congo, Cuba, France, Gabon, Guadeloupe, Haiti, La Reunion, Madeira, The Netherlands, New Caledonia, Spain, and Switzerland) which were challenged with different ZIKV genotypes (West Africa, Asia, America) using standardized experimental protocols (mosquito infections, viral titrations, parameters of vector competence). In this work, we elaborated a vector competence data-driven prediction for ZIKV transmission to inform on the current risk of Zika transmission at a global scale. We show that *Ae. aegypti* populations tested are highly competent to transmit the different ZIKV genotypes and more specifically, the African genotype. Moreover, other mosquito species such as *Ae. albopictus* or *Ae. japonicus* can play a secondary role in transmission while *Culex* mosquitoes are definitively non-susceptible to ZIKV.

Results

Culex spp. mosquitoes were resistant to most ZIKV strains, successfully blocking infection with an overall infection rate (IR) of 0.2% (2 mosquitoes out of 1089 tested) and complete absence of both dissemination efficiency (DE) and transmission efficiency (TE). As such, they were not included in more complex vector competence analyses. *Aedes* spp. showed different patterns depending on species and ZIKV strain used for infection. For *Ae. aegypti* and *Ae. albopictus*, the experimental design required nested, within-country random effects. In the case of *Ae. japonicus*, the country and mosquito population effects were fully colinear, therefore analyses only relied on logistic regressions.

Differential susceptibility of *Aedes* spp. mosquitoes to ZIKV

Depending on mosquito species and ZIKV strain, we observed different patterns of viral progression within the vector's organs. In *Ae. aegypti* infected with ZIKV strains, there were no notable differences between IR and DE at 21 days post-infection (dpi), with IR = 94.3% (95% CI: [92.1%–96.0%]) and 69.1% [64.0%–73.9%], and with DE = 89.4% [86.6%–91.8%] and 64.3% [59.1%–69.3%] for respectively, African and American strains. However, there was a large and significant decrease for TE = 71.9% [68.0%–75.6%] and 30.0% [25.3%–35.1%] for respectively, African and American strains as can be seen in Fig. 1. This is consistent with efficient viral progression through the midgut, and partial barriers to viral progression before reaching the salivary glands. In contrast, when infected with Asian ZIKV strains, both midguts and salivary glands mitigated progression of the virus: IR = 68.9% [64.3%–73.3%], DE = 56.7% [51.8%–61.5%], and TE = 28.9% [24.7%–33.5%].

In *Ae. albopictus* mosquitoes, viral progression was more efficient with the African ZIKV strain than with American or Asian ZIKV strains for all three outcomes. For the African ZIKV strain, a regular decrease was observed between IR (73.9%), DE (69.2%) and TE (55.8%), consistent with the absence of a marked barrier to viral progression through the midgut and into the salivary glands. In contrast, DE was significantly lower for American (15.6% [11.5–20.4%]) and Asian (11.2% [7.7–15.5%]) ZIKV strains, consistent with a lack of viral progression through the midgut.

Time since infection showed an increasing, approximately linear association with DE and TE in *Ae. aegypti* and *Ae. albopictus* mosquitoes, suggesting an accumulation pattern and an overall absence of viral clearance within the vector, while IR reached a peak value within 7 dpi and remained at comparable levels until 21 dpi. In contrast, *Ae. japonicus* infected by the African ZIKV strain showed that IR peaked at 14 dpi, with decreasing DE and TE values over time (TE_{7dpi} = 15.0% [7.1–26.6%], TE_{14dpi} = 15.0% [7.1–26.6%] and TE_{21dpi} = 6.7% [18.5–16.2%]).

Transmission efficiency is higher with the African ZIKV

Although all species of *Aedes* studied were competent to transmit ZIKV, infected *Ae. aegypti* species were on average more prone to successful dissemination than *Ae. albopictus* and *Ae. japonicus* (TE_{*aegypti*} = 34.0% [24.0–45.8%], TE_{*albopictus*} = 17.6% [11.4–26.0%], TE_{*japonicus*} = 7.2% [2.9–16.9%]). This was even more pronounced at 21 dpi when TE was significantly higher for *Ae. aegypti* than *Ae. albopictus* (46.7% [35.3–58.5%] vs. 25.3% [17.3–35.5%], $p < 0.0001$) and *Ae. japonicus* (46.7% [35.3–58.5%] vs. 10.8% [4.4–24.1%], $p = 0.0002$), the latter two not reaching statistical significance ($p = 0.09$). However, many combinations of ZIKV strains and mosquito populations were not available for *Ae. japonicus* for which results should be considered less robust.

The African ZIKV strain was the most efficient in infecting and disseminating, therefore posing a greater risk in case it was newly introduced in a region: the predicted expected marginal means 21 dpi were TE = 82.7% [78.5–86.1%] for *Ae. aegypti* and TE = 52.4% [45.4–59.3%] for *Ae. albopictus* (Fig. 2). American and Asian strains had comparable TE regardless of time since infection: (1) in Cameroon,

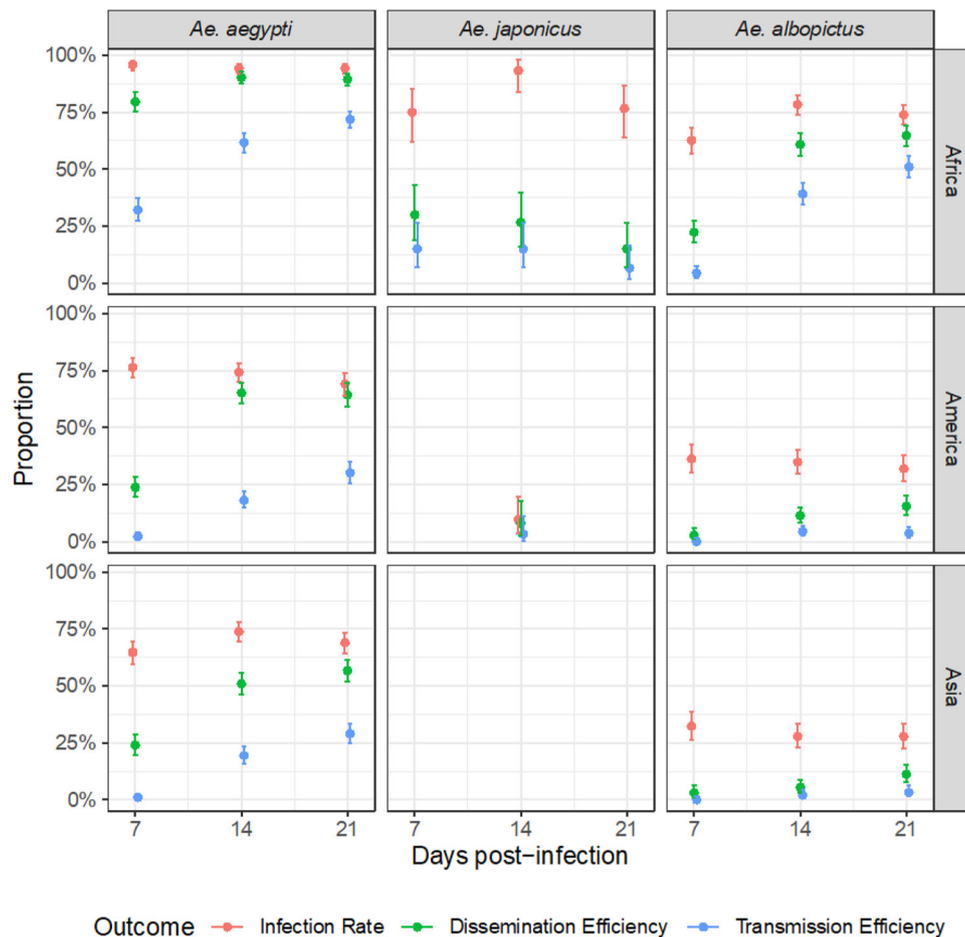


Fig. 1 | Infection rate, dissemination and transmission efficiencies of ZIKV strains (Africa, America, Asia) according to time since infection in three *Aedes* mosquito species. Panels show the rates achieved by averaging over all sampled mosquitoes, i.e., by pooling together mosquitoes from all countries used in the study. Error bars are exact 95% binomial confidence intervals centered on the

observed means of IR, DE and TE. ($n = 30$ biologically independent mosquitoes for each combination of species, country, mosquito population, days post-infection and ZIKV strain were studied, unless stated otherwise. See Supplementary Table 1 for complete break-down of sample sizes). Colors correspond to the studied outcome (IR, DE, TE).

Congo, and Gabon, TE was low (~10%), suggesting African mosquitoes are mostly susceptible to local ZIKV strains; (2) in Brazil, Haiti, Guadeloupe, Madeira, La Reunion and New Caledonia, TE reached moderate values ~20%. In Cambodia however, Asian and American ZIKV strains had much higher TE values, denoting mosquito populations highly susceptible to ZIKV.

Breaking down variable importance in ZIKV susceptibility

We evaluated the extent to which each covariate of interest contributed to explanatory value of fitted models. For both *Ae. aegypti* and *Ae. albopictus*, TE showed low variability among mosquito populations originating from the same country (Fig. 3). Another striking result from Fig. 3 confirmed the higher susceptibility to African ZIKV strains among all tested mosquitoes, both within and across countries. This suggests that within a given country, mosquito populations were equally able to disseminate ZIKV regardless of other extrinsic factors. We found that 51.7% of variance in *Ae. aegypti* TE could be explained by a statistical model accounting for mosquito strain, ZIKV strain, and dpi (Table 1). The relative importance of these factors was assessed by estimating the proportion of variance explained solely by these factors. For *Ae. aegypti* TE, ZIKV strain was the most important factor explaining 33.5% of variance, followed by dpi (30.1%). Of the factors considered, mosquito strain had the least impact on TE, with only 6.6% of variance explained by the country of a sampled mosquito strain. An additional 2.6% of variance was explainable when we considered

locations of strains within a country. For *Ae. japonicus*, no obvious pattern could be detected with only 5.2% of the observed variance explained by available covariates suggesting that mosquitoes were less susceptible to tested ZIKV strains.

Discussion

Our results confirmed that human-adapted *Aedes* mosquitoes prepare the field for new arbovirus emergences; in addition to other factors, some combinations of vector-virus are more prone to cause outbreaks, but globalization will smooth out differences leading most emergences to turn into pandemics owing to the large territories covered by vectors.

During the 2015–2016 Latin American and the Caribbean epidemic, the role of *Culex* as ZIKV vector has been the subject of major scientific debates which, if verified, would have revised the *modus operandi* of vector control¹¹. As ZIKV has also been detected in field-collected *Culex* mosquitoes^{12,13}, their susceptibility to ZIKV was then tested; from 36 studies on vector competence of *Culex* mosquitoes for ZIKV, seven studies showed that ZIKV was present in the salivary glands of *Cx. restuans*, *Cx. quinquefasciatus*, *Cx. tarsalis*, and *Cx. coronator*. Asian and American genotypes of ZIKV may occasionally infect *Culex* mosquitoes but not African strains¹⁴. From our dataset^{15–18}, we showed that ZIKV can accumulate in *Cx. pipiens* saliva only after forced infection by intrathoracic inoculation, short-circuiting the midgut barrier¹⁷; after oral infection, the virus was unlikely to disseminate in the mosquito ruling out the role of *Culex* mosquitoes in ZIKV

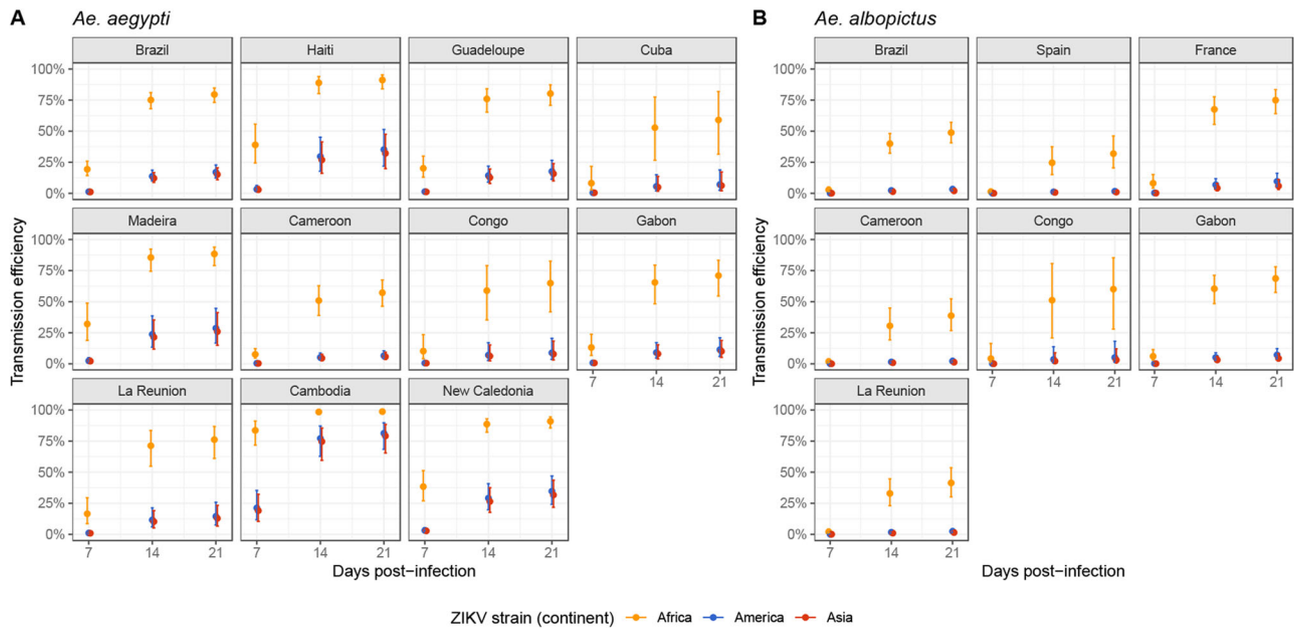


Fig. 2 | Model-predicted transmission efficiency of *Aedes* mosquitoes according to time since infection, split by country of mosquito sampling. TE predicted for (A) *Ae. aegypti* and (B) *Ae. albopictus*. Countries are ordered to reflect geographic proximity. Error bars show asymptotic 95% confidence interval from the mixed regression models centered on the average predictions of IR, DE and

TE. (n = 30 biologically independent mosquitoes for each combination of species, country, mosquito population, days post-infection and ZIKV strain were studied, unless stated otherwise. See Supplementary Table 1 for complete break-down of sample sizes). Colors correspond to continent-aggregated ZIKV strains.

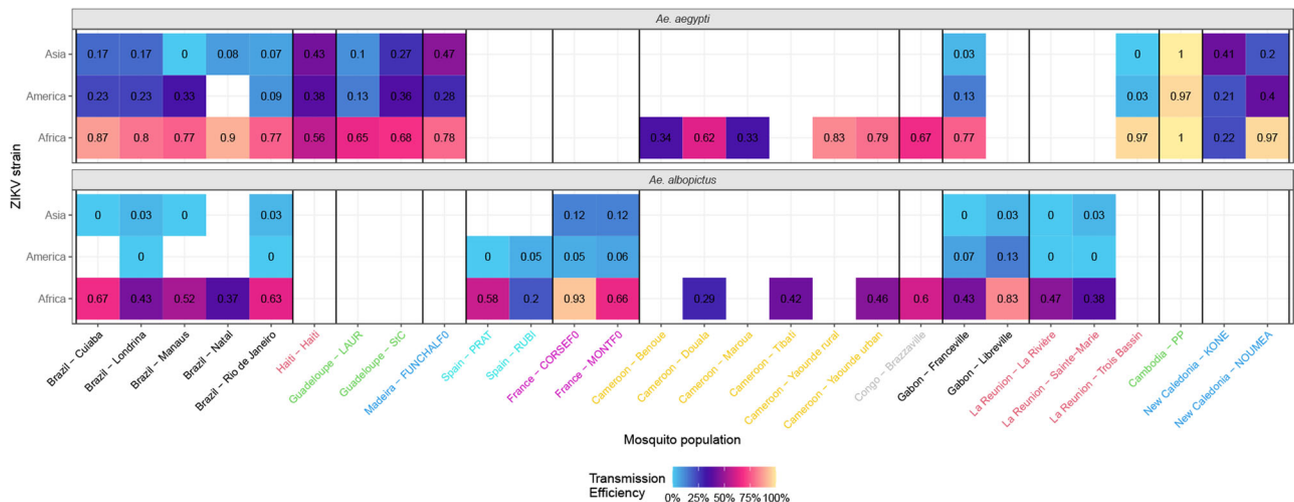


Fig. 3 | Transmission efficiencies of ZIKV strains at the regional level for *Ae. aegypti* and *Ae. albopictus* mosquitoes sampled at one or more locations in every studied country, at 21 dpi. ZIKV strains were pooled in a geographical

clustering that reflected their phylogeny. Vertical bars help identify different countries, shown on the x-axis with individual colors. Empty cells represent absence of data. Color gradients correspond to TE values (ranging 0–100%).

transmission. Viral replication was not detected in *Culex* which was unable to trigger the RNAi pathway as no viral replication signatures such as 21 nt viral siRNAs were detected¹⁹. Even though vectors of the flavivirus West-Nile virus, a phylogenetically ZIKV-related arbovirus, mosquitoes of the *Cx. pipiens* complex are not competent for ZIKV transmission and are very unlikely to play a role of vector during a Zika outbreak. Thus, to control Zika, targeted interventions should focus largely on *Aedes* mosquitoes^{11,20}.

Aedes aegypti is an African mosquito (named *Ae. aegypti formosus*) living primarily in rainforests colonizing natural breeding sites which in a subsequent step, has become adapted to living in the human environment (developing a human-biting behavior and colonizing human-made containers) before undergoing different waves of dissemination

outside Africa²¹. The “domestication” of this species (named *Ae. aegypti aegypti*) allowed its world-wide dispersion coinciding with the first twentieth century pandemics caused by arboviruses, dengue²², and then chikungunya²³ and Zika²⁴. In addition to increased vector-host contacts, *Ae. aegypti aegypti* has developed an enhanced permissiveness to some arboviruses⁷. At the mosquito level, successful transmission implies that the virus passes through two anatomical barriers, midgut and salivary glands²⁵. *Ae. aegypti* mosquitoes we tested were more prone to impose a strong bottleneck to the African ZIKV in the salivary glands leading to decrease ZIKV transmission. Nevertheless, all 11 *Ae. aegypti* populations transmitted at least 2–3 times more the African ZIKV than the American and Asian ZIKV strains underlining the high potential of African ZIKV to trigger an outbreak outside and inside

Table 1 | Contribution of random effects and fixed effects in species-wise regression models

Mosquito species	Total variance explained	Variance uniquely explained			
		Within-country (mosquito)	Between-country (mosquito)	ZIKV strain	Days post-infection
<i>Ae. aegypti</i>	51.7%	2.6%	6.6%	33.5%	30.1%
<i>Ae. japonicus</i>	5.2%		1.0%	0%	1.0%
<i>Ae. albopictus</i>	41.5%	1.7%	0%	29.0%	21.2%

The contribution of random effects was measured as the adjusted intra-class correlation coefficient (as defined in Nakagawa et al.⁵²), while the contribution of fixed effects was assessed by the semi-partial R^2 measure (from Stoffel et al.⁵³), i.e. the share of variance uniquely explained by a given covariate. The total variance explained is the R^2 measure of the saturated model.

Africa. It is now admitted that African ZIKV strains were more pathogenic than the Asian strains^{26,27}. African *Ae. aegypti* (Cameroon, Congo, Gabon) were more resistant to American and Asian ZIKV than to African ZIKV suggesting a local adaptation between a ZIKV strain and a mosquito population. There is evidence today that *Ae. aegypti formosus* is found both in urban and forest habitats in sub-Saharan Africa²⁸ and a mixture of *Ae. aegypti aegypti* and *Ae. aegypti formosus* is likely present in cities we sampled in Cameroon, Congo and Gabon²⁹. As *Ae. aegypti aegypti* is more susceptible to ZIKV than *Ae. aegypti formosus*, an intermediate level of susceptibility can be expected⁷. *Ae. aegypti formosus*, which is predominant in Benoué, was previously found less susceptible to ZIKV compared to the other populations from Cameroon³⁰. Nevertheless, mosquitoes were only identified according to morphological criteria³¹ and not based on their genotypes. As such, the observed local reduction in vector competence may also result from other inherited factors not detected through morphological examinations. Our experiments showed a decreased susceptibility within the country, with *Ae. aegypti* from Benoué having lower TE (34.4% [20.2–52.1%]) than those sampled elsewhere in Cameroon (ranging from 62.5 to 83.3%, Supplementary Table 2) except in Maroua, located close to Benoué. Intriguingly, human epidemics of ZIKV associated with *Ae. aegypti* have been observed in a limited number of countries in Asia or Africa following the 2015–2016 Latin American and the Caribbean epidemic²⁴. The Asian *Ae. aegypti* population from Cambodia we tested, transmitted the most efficiently all three ZIKV strains, African, American and Asian, corroborating its good vector competence to ZIKV. Recent systematic screenings of patients revealed that ZIKV was circulating in Singapore, Thailand, Cambodia, Myanmar and Vietnam suggesting that ZIKV has been present in Asia at a low but sustained level for years or even decades^{32–35}. Moreover, human population immunity could also modulate the risk of outbreaks³⁶; a large population exposed to dengue viruses may develop neutralizing antibodies against dengue viruses which cross-react with ZIKV adjusting the outcome of ZIKV infection³⁶. Thereby, Asian populations can be partially protected against ZIKV³⁷. At an intra-country level, mosquito populations (in our study, e.g. five in Brazil¹⁸) share the same profiles of transmission with a highest transmission level with African ZIKV. Once introduced, the African ZIKV has the potential to cause an outbreak whatever the origin of *Ae. aegypti* populations. Our assessments are based on vector competence which is only one aspect of vectorial capacity; it results from complex interactions of multiple factors influencing pathogen transmission by a vector, including internal factors such as genetic, evolution, immunity, or interactions with other microorganisms, modulated by external factors such as abiotic (like climate or topography) and biotic (like nutrition or hosts)³⁸.

In the tropical belt, *Ae. aegypti* and *Ae. albopictus* share the same ecological niche³⁹. *Ae. albopictus* mosquitoes we tested were more susceptible to African ZIKV suggesting that the species can contribute to an epidemic like it happened in Gabon in 2007⁴⁰. A lower transmission of American and Asian ZIKV was presumably related to a strong mosquito midgut barrier³⁰. Since its arrival in Central Africa in 2000, *Ae. albopictus* plays a major role in the transmission of different arboviruses⁴¹. *Ae. albopictus* tends to

predominate over *Ae. aegypti* in habitats where both species are sympatric⁴² and will certainly promote arbovirus spillovers at the edge of their natural sylvatic cycle owing to its generalist behavior (biting both humans and animals, and living in rural areas)⁴¹. *Ae. albopictus* from Central Africa are likely originated from South America which themselves were introduced from North America in 1986⁴³. European *Ae. albopictus* are presumably a genetic mixture of populations from United States and La Reunion⁴³, sharing similar profiles of ZIKV transmission with mosquitoes from La Reunion^{16,44,45}. American ZIKV predominant during the 2015–2016 epidemic, was efficiently transmitted by *Ae. albopictus*, in Europe⁵ and Brazil⁴⁶. In the absence of *Ae. aegypti*, *Ae. albopictus* is more likely to trigger ZIKV outbreaks when ZIKV comes from Africa.

Another invasive species, *Ae. japonicus* also experimentally transmits the African ZIKV⁴⁷. Native to East Asia, *Ae. japonicus* arrived in Europe in 2000⁴¹ and is established in 12 European countries up to now⁴⁸. Its role in pathogen transmission is still unclear even if arboviral ZIKV RNAs have been detected in field-collected mosquitoes⁴⁹. Its high ability to become infected, and to a lower extent, to ensure viral dissemination and transmission, concomitantly to its human-biting behavior gives *Ae. japonicus* the status of being a competent vector of ZIKV^{47,50}. These findings are however subject to caution, as the experimental design may not allow for generalization of our observations: *Ae. japonicus* could only be sampled and bred in European countries (France ($n = 90$), Switzerland ($n = 90$), the Netherlands ($n = 62$)). To add to this limitation, only two ZIKV strains were tested against these mosquitoes and fully collinear with the country of mosquito sampling: French- and Swiss-sampled *Ae. japonicus* were infected with an African ZIKV strain, whereas mosquitoes from the Netherlands were infected with an American ZIKV strain (see data structure in Supplementary Fig. 1). Therefore, we cannot distinguish between mosquito susceptibility and ZIKV strain potency.

We conclude that from our comprehensive assessment of vector competence, invasive *Aedes* mosquitoes set the bed for pandemic arboviruses; in the absence of the historical vector of human arboviruses, *Ae. aegypti*, other species such as *Ae. albopictus* or *Ae. japonicus* can take over as ZIKV vectors and transmit even more efficiently when infected with the African ZIKV.

Methods

Experimental procedures

All partners of the ZIKAlliance project (<https://zikalliance.tghn.org/>) performed their own mosquito experimental infections using a common standardized protocol (Supplementary Fig. 2). Some deviations from this protocol could not be avoided (e.g. availability of blood source, feeding via blood droplets instead of *via* membrane). Fifty populations (Fig. 4 and Supplementary Table 1) were collected as immature stages and reared in insectaries under controlled conditions. Mosquito females were exposed to an infectious blood meal containing one third of viral suspension and two third of washed rabbit erythrocytes (Supplementary Fig. 3) at a final titer of -10^7 TCID₅₀/ml. After 1 h, engorged females were isolated in boxes ($28^{\circ}\pm 1^{\circ}\text{C}$ and $80\pm 10\%$ humidity) and fed with 10% sucrose until analysis.

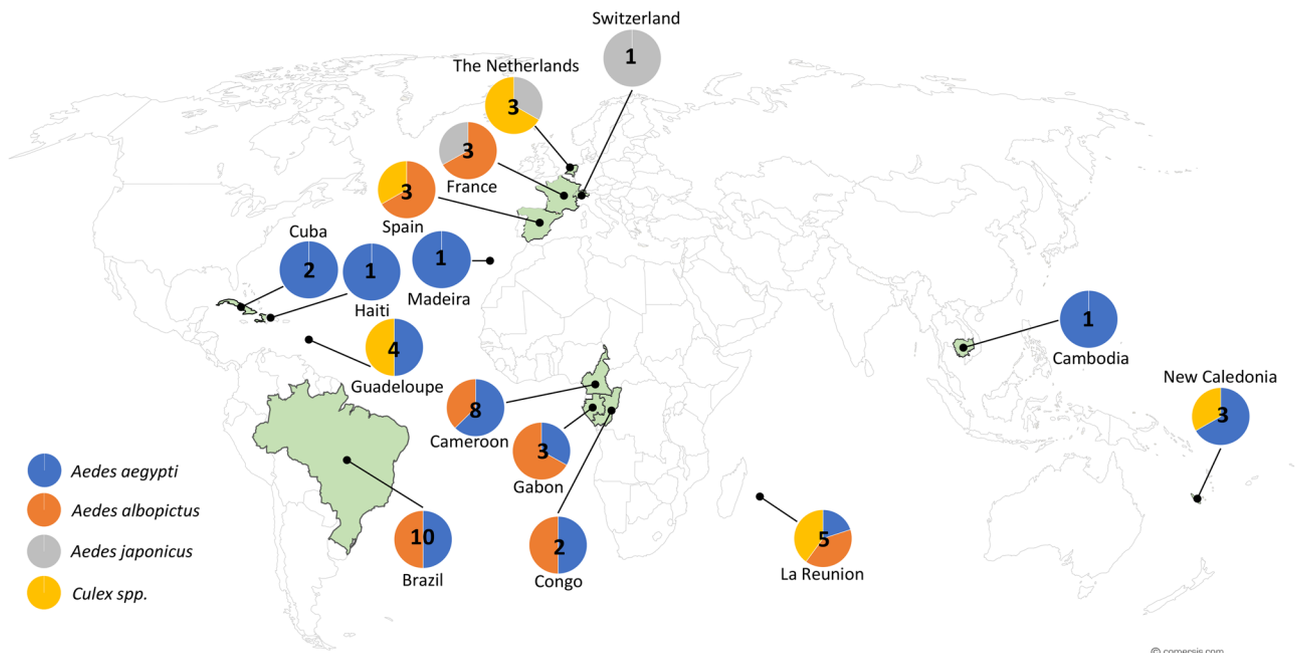


Fig. 4 | Distribution of mosquito species experimentally infected with Zika viruses. The color code used represents the different mosquito species and numbers refer to the number of mosquito populations tested. In green, are the

countries sampled. Colors correspond to mosquito species. The map was built using the open source map site <https://cmap.comersis.com/cartes-Monde-WORLD.html>.

Different geographic ZIKV strains were used (Supplementary Fig. 4): (1) Africa (Dakar; KU955592; isolated in 1984 from *Aedes taylori* in Senegal and passaged four times on BHK21 cells), (2) Asia (Cambodia; KU955593; isolated from a human case in 2010 and passaged three times on Vero cells and Malaysia; KX694533; isolated in 1966 from *Aedes aegypti* in Malaysia and passaged four times on Vero cells), and (3) America (Guadeloupe; LR792671.1; isolated from a human case in Guadeloupe in 2016 and passaged two times on Vero cells, Martinique; KU647676; isolated from a human case in 2015 and passaged three times on Vero cells, and Suriname; KU937936; isolated from a human serum in Suriname in 2016 and passaged 5–6 times on Vero cells). ZIKV Dakar, Malaysia, and Martinique provided by EVAg (<https://www.european-virus-archive.com/>) were used in most infections. ZIKV Cambodia was used for the mosquito populations Haiti, Funchal, Mont, Corse, ZIKV Suriname for Best, Amsterdam and Lelystad, and ZIKV Guadeloupe for HAV-PT, HAV-PRG.

At 7, 14, and 21 days post-infection (dpi), a batch of mosquitoes was processed to estimate if viral particles were present in the body (thorax and abdomen) as the marker for infection, head for dissemination and saliva for transmission; 0 refers to no viral particles and 1 to at least one viral particle detected. Presence of viral particles was asserted by cytopathic effects observed on a monolayer of Vero CCL-81 cells in 96-well plates⁴⁵. Three parameters were measured: (1) IR corresponding to the proportion of mosquitoes with infected body (thorax and abdomen) among examined mosquitoes, (2) DE referring to the proportion of mosquitoes with virus detected in the head among examined mosquitoes, and (3) TE representing the proportion of mosquitoes with virus detected in saliva among examined mosquitoes. TE was used as the primary endpoint to evaluate vector competence with respect to ZIKV strain. It is important to note that, by definition, $TE \leq DE \leq IR$, the denominator of each being the total number of mosquitoes dissected.

Statistical analysis

Success for transmission, dissemination and infection were investigated as binomial outcomes to assess how their population-level counterpart (TE, DE, IR) were associated with available

combinations of mosquito populations and ZIKV strains, according to time since infection. Time since infection was considered a categorical covariate. Prior to any modeling approach, all outcomes were visualized with heat maps similar to those presented in Supplementary Figs. 6, 8 and 9, for all combinations of ZIKV strain and mosquito population and at every value of time since infection. The resulting experimental design, consisting of combinations of country of study, mosquito species and populations as well as time since infection, was used to guide the development of generalized linear (mixed) models.

Owing to complex hierarchical experimental design, each mosquito species was modeled separately. When several mosquito populations were sampled at different locations within a given country, the variations resulting from the population effect was modeled by a random effect nested within the country, in a mixed regression framework. The contribution of each fixed-effect term was assessed by refitting univariable models and measuring the semi-partial R-squared value (as defined by Stoffel et al.⁵¹). Likewise, the contribution of random effects was measured as the adjusted intra-class correlation coefficient (as defined by Nakagawa et al.⁵²). Analyses were conducted using the R software (v. 4.2.0)⁵³ with the following packages: lme4 (v. 1.1-27.1)⁵⁴, emmeans (v. 1.7.2), performance (v.0.8.0)⁵⁵ and partR2 (v. 0.9.1).

The robustness of results was assessed using sensitivity analyses, where the same regression models were applied with data aggregated at larger geographical scales for covariates of interest. Three variations of the same analyses were conducted: (1) by aggregating country of study by continent (therefore assuming random variations across countries) and preserving individual ZIKV strains, (2) by aggregating ZIKV strains by continent (corresponding to relatedness between strains in the phylogenetic tree, see Supplementary Fig. 4) and preserving individual countries of study, and (3) aggregating both country and virus strain by continent. The results from aggregation detailed in point (2) are presented in the main text as they provide the most comprehensive set of results. The results corresponding to aggregation levels from points (1), (3) and (4) are presented in Supplementary Figs. 5 through 10.

Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The data that support the findings of this study are fully available in an electronic Supplementary table (Microsoft Excel XLSX format) at <https://zenodo.org/record/6788898#.Ysgu0oTP2Uk> as well as in the GitLab repository found in the Code availability section.

Code availability

All analyses conducted in this manuscript and associated code have been uploaded to a Git repository at <https://gitlab.pasteur.fr/tobadia/17289-zikalliance> and are readily available. The results of the present manuscript are based on commit 1477a8be1554c91d2a5ea038964407e88499ddf9.

References

- Patterson, K. D. & Pyle, G. F. The geography and mortality of the 1918 influenza pandemic. *Bull. Hist. Med.* **65**, 4–21 (1991).
- Poirier, E. Z. & Vignuzzi, M. Virus population dynamics during infection. *Curr. Opin. Virol.* **23**, 82–87 (2017).
- Huang, Y.-J. S., Higgs, S. & Vanlandingham, D. L. Emergence and re-emergence of mosquito-borne arboviruses. *Curr. Opin. Virol.* **34**, 104–109 (2019).
- Dick, G. W. A., Kitchen, S. F. & Haddow, A. J. Zika Virus (I). Isolations and serological specificity. *Trans. R. Soc. Tropical Med. Hyg.* **46**, 509–520 (1952).
- Giron, S. et al. Vector-borne transmission of Zika virus in Europe, southern France, August 2019. *Eurosurveillance* **24**, 1900655 (2019).
- Sim, S., Jupatanakul, N. & Dimopoulos, G. Mosquito immunity against arboviruses. *Viruses* **6**, 4479–4504 (2014).
- Aubry, F. et al. Enhanced Zika virus susceptibility of globally invasive *Aedes aegypti* populations. *Science* **370**, 991–996 (2020).
- Lambrechts, L. et al. Genetic specificity and potential for local adaptation between dengue viruses and mosquito vectors. *BMC Evol. Biol.* **9**, 160 (2009).
- Mariconti, M. et al. Estimating the risk of arbovirus transmission in Southern Europe using vector competence data. *Sci. Rep.* **9**, 17852 (2019).
- Viglietta, M., Bellone, R., Blisnick, A. A. & Failloux, A.-B. Vector specificity of arbovirus transmission. *Front. Microbiol.* **12**, 773211 (2021).
- Roundy, C. M. et al. Lack of evidence for Zika virus transmission by *Culex* mosquitoes. *Emerg. Microbes Infect.* **6**, 1–2 (2017).
- Guo, X. et al. *Culex pipiens quinquefasciatus*: a potential vector to transmit Zika virus. *Emerg. Microbes Infect.* **5**, 1–5 (2016).
- Guedes, D. R. et al. Zika virus replication in the mosquito *Culex quinquefasciatus* in Brazil. *Emerg. Microbes Infect.* **6**, 1–11 (2017).
- Viveiros-Rosa, S. G., Regis, E. G. & Santos, W. C. Vector competence of *Culex* mosquitoes (Diptera: Culicidae) in Zika virus transmission: an integrative review. *Rev. Panam. Salud Publica* **44**, e7 (2020).
- Hery, L., Boullis, A., Delannay, C. & Vega-Rúa, A. Transmission potential of African, Asian and American Zika virus strains by *Aedes aegypti* and *Culex quinquefasciatus* from Guadeloupe (French West Indies). *Emerg. Microbes Infect.* **8**, 699–706 (2019).
- Gomard, Y., Lebon, C., Mavingui, P. & Atyame, C. M. Contrasted transmission efficiency of Zika virus strains by mosquito species *Aedes aegypti*, *Aedes albopictus* and *Culex quinquefasciatus* from Reunion Island. *Parasites Vectors* **13**, 398 (2020).
- Abbo, S. R. et al. Forced Zika virus infection of *Culex pipiens* leads to limited virus accumulation in mosquito saliva. *Viruses* **12**, 659 (2020).
- Fernandes, R. S. et al. Vector competence of *Aedes aegypti*, *Aedes albopictus* and *Culex quinquefasciatus* from Brazil and New Caledonia for three Zika virus lineages. *Pathogens* **9**, 575 (2020).
- Lourenço-de-Oliveira, R. et al. *Culex quinquefasciatus* mosquitoes do not support replication of Zika virus. *J. Gen. Virol.* **99**, 258–264 (2018).
- Lourenço-de-Oliveira, R. & Failloux, A.-B. Lessons learned on Zika virus vectors. *PLoS Negl. Trop. Dis.* **11**, e0005511 (2017).
- Gloria-Soria, A. et al. Global genetic diversity of *Aedes aegypti*. *Mol. Ecol.* **25**, 5377–5395 (2016).
- Guzman, M. G. & Harris, E. Dengue. *Lancet* **385**, 453–465 (2015).
- Weaver, S. C. & Lecuit, M. Chikungunya virus and the global spread of a mosquito-borne disease. *N. Engl. J. Med.* **372**, 1231–1239 (2015).
- Weaver, S. C., Charlier, C., Vasilakis, N. & Lecuit, M. Zika, Chikungunya, and other emerging vector-borne viral diseases. *Annu. Rev. Med.* **69**, 395–408 (2018).
- Kramer, L. D. & Ciota, A. T. Dissecting vectorial capacity for mosquito-borne viruses. *Curr. Opin. Virol.* **15**, 112–118 (2015).
- Aubry, F. et al. Recent African strains of Zika virus display higher transmissibility and fetal pathogenicity than Asian strains. *Nat. Commun.* **12**, 916 (2021).
- Smith, D. R. et al. African and Asian Zika virus isolates display phenotypic differences both in vitro and in vivo. *Am. J. Trop. Med. Hyg.* **98**, 432–444 (2017).
- Powell, J. R. Mosquitoes on the move. *Science* **354**, 971–972 (2016).
- Weetman, D. et al. *Aedes* mosquitoes and *Aedes*-borne arboviruses in Africa: current and future threats. *Int. J. Environ. Res. Public Health* **15**, 220 (2018).
- Kamgang, B. et al. Different populations of *Aedes aegypti* and *Aedes albopictus* (Diptera: Culicidae) from Central Africa are susceptible to Zika virus infection. *PLoS Negl. Trop. Dis.* **14**, e0008163 (2020).
- McClelland, G. A. H. A worldwide survey of variation in scale pattern of the abdominal tergum of *Aedes aegypti* (L.) (Diptera: Culicidae). *Trans. R. Entomol. Soc. Lond.* **126**, 239–259 (1974).
- Ho, Z. J. M. et al. Outbreak of Zika virus infection in Singapore: an epidemiological, entomological, virological, and clinical analysis. *Lancet Infect. Dis.* **17**, 813–821 (2017).
- Ruchatsawat, K. et al. Long-term circulation of Zika virus in Thailand: an observational study. *Lancet Infect. Dis.* **19**, 439–446 (2019).
- Tun, M. M. N. et al. Detection of Zika virus infection in Myanmar. *Am. J. Trop. Med. Hyg.* **98**, 868–871 (2018).
- Quyen, N. T. H. et al. Chikungunya and Zika virus cases detected against a backdrop of endemic Dengue transmission in Vietnam. *Am. J. Trop. Med. Hyg.* **97**, 146–150 (2017).
- Wen, J. & Shresta, S. Antigenic cross-reactivity between Zika and dengue viruses: is it time to develop a universal vaccine? *Curr. Opin. Immunol.* **59**, 1–8 (2019).
- Luo, X. S., Imai, N. & Dorigatti, I. Quantifying the risk of Zika virus spread in Asia during the 2015–16 epidemic in Latin America and the Caribbean: a modeling study. *Travel Med. Infect. Dis.* **33**, 101562 (2020).
- Lefèvre, T., Vantaux, A., Dabiré, K. R., Mouline, K. & Cohuet, A. Non-genetic determinants of mosquito competence for malaria parasites. *PLoS Pathog.* **9**, e1003365 (2013).
- Lwande, O. W. et al. Globe-Trotting *Aedes aegypti* and *Aedes albopictus*: risk factors for arbovirus pandemics. *Vector-Borne Zoonotic Dis.* **20**, 71–81 (2020).
- Grard, G. et al. Zika Virus in Gabon (Central Africa)—2007: a new threat from *Aedes albopictus*? *PLoS Negl. Trop. Dis.* **8**, e2681 (2014).
- Ngoagouni, C., Kamgang, B., Nakouné, E., Paupy, C. & Kazanji, M. Invasion of *Aedes albopictus* (Diptera: Culicidae) into central

- Africa: what consequences for emerging diseases? *Parasites Vectors* **8**, 191 (2015).
42. Kamgang, B. et al. Temporal patterns of abundance of *Aedes aegypti* and *Aedes albopictus* (Diptera: Culicidae) and mitochondrial DNA analysis of *Ae. albopictus* in the Central African Republic. *PLoS Negl. Trop. Dis.* **7**, e2590 (2013).
 43. Vega-Rúa, A. et al. Vector competence of *Aedes albopictus* populations for chikungunya virus is shaped by their demographic history. *Commun. Biol.* **3**, 1–13 (2020).
 44. Nuñez, A. I. et al. Evidence of Zika virus horizontal and vertical transmission in *Aedes albopictus* from Spain but not infectious virus in saliva of the progeny. *Emerg. Microbes Infect.* **9**, 2236–2244 (2020).
 45. Vazeille, M. et al. Zika virus threshold determines transmission by European *Aedes albopictus* mosquitoes. *Emerg. Microbes Infect.* **8**, 1668–1678 (2019).
 46. Parra, M. C. P. et al. Detection of Zika RNA virus in *Aedes aegypti* and *Aedes albopictus* mosquitoes, São Paulo, Brazil. *Infect., Genet. Evolution* **98**, 105226 (2022).
 47. Glavinic, U. et al. Assessing the role of two populations of *Aedes japonicus japonicus* for Zika virus transmission under a constant and a fluctuating temperature regime. *Parasites Vectors* **13**, 479 (2020).
 48. Koban, M. B. et al. The Asian bush mosquito *Aedes japonicus japonicus* (Diptera: Culicidae) in Europe, 17 years after its first detection, with a focus on monitoring methods. *Parasites Vectors* **12**, 109 (2019).
 49. Yang, F. et al. Cache valley virus in *Aedes japonicus japonicus* Mosquitoes, Appalachian Region, United States. *Emerg. Infect. Dis.* **24**, <https://doi.org/10.3201/eid2403.161275> (2018).
 50. Abbo, S. R. et al. The invasive Asian bush mosquito *Aedes japonicus* found in the Netherlands can experimentally transmit Zika virus and Usutu virus. *PLoS Negl. Trop. Dis.* **14**, e0008217 (2020).
 51. Stoffel, M. A., Nakagawa, S. & Schielzeth, H. partR2: partitioning R² in generalized linear mixed models. *PeerJ* **9**, e11414 (2021).
 52. Nakagawa, S., Johnson, P. C. D. & Schielzeth, H. The coefficient of determination R² and intra-class correlation coefficient from generalized linear mixed-effects models revisited and expanded. *J. R. Soc. Interface* **14**, 20170213 (2017).
 53. R Core Team. *R: A Language and Environment for Statistical Computing* (R Foundation for Statistical Computing, 2022).
 54. Bates, D., Mächler, M., Bolker, B. & Walker, S. Fitting linear mixed-effects models using lme4. *J. Stat. Softw.* **67**, 1–48 (2015).
 55. Lüdtke, D. et al. see: An R Package for Visualizing Statistical Models. <https://doi.org/10.31234/osf.io/m4uax> (2021).

Acknowledgements

We thank Richard Paul (Institut Pasteur) for correcting English. This study was supported by the European Union's Horizon 2020 research and innovation program under ZIKAlliance grant agreement no. 734548.

The funders had no role in study design, data collection and interpretation, or decision to submit the work for publication.

Author contributions

T.O. performed the writing and editing, data analysis and interpretation. G.G.B., A.I.N., R.S.F., B.K., L.H., Y.G., S.R.A., D.J., U.G., M.V., A.Y., and C.J.M.K. did the investigation and data acquisition. V.D. did the investigation, methodology, and data acquisition. M.D.R., C.M.A., N.P., S.B., P.M., A.V.R., E.V., G.P.P., C.P., N.B., and R.L.O. contributed to supervision, data analysis and interpretation. C.D. and M.T.W. did the data analysis and interpretation. X.D.L. participated in funding acquisition and project administration. A.B.F. intervened in conceptualization, project administration, supervision, and writing original draft. All authors have read and agreed with the published version of the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information The online version contains supplementary material available at <https://doi.org/10.1038/s41467-022-32234-y>.

Correspondence and requests for materials should be addressed to Anna-Bella Failloux.

Peer review information *Nature Communications* thanks Sally Paulson and the other, anonymous, reviewer(s) for their contribution to the peer review of this work.

Reprints and permission information is available at <http://www.nature.com/reprints>

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

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit <http://creativecommons.org/licenses/by/4.0/>.

© The Author(s) 2022