

# Edinburgh Research Explorer

## Genomic epidemiology reveals multiple introductions of Zika virus into the United States

Citation for published version:

Citation for published version:
Grubaugh, ND, Ladner, JT, Kraemer, MUG, Dudas, G, Tan, AL, Gangavarapu, K, Wiley, MR, White, S, Theze, J, Magnani, DM, Prieto, K, Reyes, D, Bingham, A, Paul, LM, Robles-Sikisaka, R, Oliveira, G, Pronty, D, Barcellona, CM, Metsky, HC, Baniecki, ML, Barnes, KG, Chak, B, Freije, CA, Gladden-Young, A, Gnirke, A, Luo, C, MacInnis, B, Matranga, CB, Park, DJ, Qu, J, Schaffner, SF, Christopher, T-T, West, KL, Winnicki, SM, Wohl, S, Yozwiak, NL, Quick, J, Fauver, JR, Khan, K, Brent, SE, Reiner Jr, RC, Lichtenberger, PN, Ricciardi, M, Bailey, VK, Watkins, DI, Cone, MR, Kopp IV, EW, Hogan, KN, Cannons, AC, Jean, R, Monaghan, AJ, Garry, RF, Loman, NJ, Faria, NR, Porcelli, MC, Vasquez, C, Nagle, ER, Cummings, DAT, Stanek, D, Rambaut, A, Sanchez-Lockhart, M, Sabeti, PC, Gillis, LD, Michael, SF, Bedford, T, Pybus, OG, Isern, S, Palacios, GF & Andersen, KG 2017, 'Genomic epidemiology reveals multiple introductions of Zika virus into the United States', *Nature*, vol. 546, pp. 401–405. https://doi.org/10.1038/nature22400

### Digital Object Identifier (DOI):

10.1038/nature22400

#### Link:

Link to publication record in Edinburgh Research Explorer

#### **Document Version:**

Peer reviewed version

#### Published In:

Nature

**General rights** 

Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.

**ACCESS** 

Download date: 27. Apr. 2024

# Genomic epidemiology reveals multiple introductions of

## Zika virus into the United States 2 Zika virus into the United States Nathan D Grubaugh<sup>1\*</sup>, Jason T Ladner<sup>2,\*</sup>, Moritz UG Kraemer<sup>3,4,5,\*</sup>, Gytis Dudas<sup>6,\*</sup>, Amanda L Tan<sup>7,\*</sup>, Karthik Gangavarapu<sup>1\*</sup>, Michael R Wiley<sup>2,\*</sup>, Stephen White<sup>8,\*</sup>, Julien Thézé<sup>3,\*</sup>, Diogo M Magnani<sup>9</sup>, Karla Prieto<sup>2</sup>, Daniel Reyes<sup>2</sup>, Andrea Bingham<sup>10</sup>, Lauren M Paul<sup>7</sup>, Refugio Robles-Sikisaka<sup>1</sup>, Glenn Oliveira<sup>11</sup>, Darryl Pronty<sup>8</sup>, Carolyn M Barcellona<sup>7</sup>, Hayden C Metsky<sup>12</sup>, Mary Lynn Baniecki<sup>12</sup>, Kayla G Barnes<sup>12</sup>, Bridget Chak<sup>12</sup>, Catherine A Freije<sup>12</sup>, Adrianne Gladden-Young<sup>12</sup>, Andreas Gnirke<sup>12</sup>, Cynthia Luo<sup>12</sup>, Bronwyn MacInnis<sup>12</sup>, Christian B Matranga<sup>12</sup>, Daniel J Park<sup>12</sup>, James Qu<sup>12</sup>, Stephen F Schaffner<sup>12</sup>, Christopher Tomkins-Tinch<sup>12</sup>, Kendra L West<sup>12</sup>, Sarah M Winnicki<sup>12</sup>, Shirlee Wohl<sup>12</sup>, Nathan L Yozwiak<sup>12</sup>, Joshua Quick<sup>13</sup>, Joseph R Fauver<sup>14</sup>, Kamran Khan<sup>15,16</sup>, Shannon E Brent<sup>15</sup>, Robert C Reiner Jr.<sup>17</sup>, Paola N Lichtenberger<sup>18</sup>, Michael Ricciardi<sup>9</sup>, Varian K Bailey<sup>9</sup>, David I Watkins<sup>9</sup>, Marshall R Cone<sup>19</sup>, Edgar W Kopp IV<sup>19</sup>, Kelly N Hogan<sup>19</sup>, Andrew C Cannons<sup>19</sup>, Reynald Jean<sup>20</sup>, Andrew J Monaghan<sup>21</sup>, Robert F Garry<sup>22</sup>, Nicholas J Loman<sup>13</sup>, Nuno R Faria<sup>3</sup>, Mario C Porcelli<sup>23</sup>, Chalmers Vasquez<sup>23</sup>, Elyse R Nagle<sup>2</sup>, Derek AT Cummings<sup>24</sup>, Danielle Stanek<sup>10</sup>, Andrew Rambaut<sup>25,26</sup>, Mariano Sanchez-Lockhart<sup>2</sup>, Pardis C Sabeti<sup>12,27,28,29#</sup>, Leah D Gillis<sup>8,#</sup>, Scott F Michael<sup>7,#</sup>, Trevor Bedford<sup>6,#</sup>, Oliver G Pybus<sup>3,#</sup>, Sharon Isern<sup>7,#</sup>, Gustavo Palacios<sup>2,#,8</sup>, Kristian G Andersen<sup>1,11,30,#,8</sup> 8 9 10 11 12 13 14 15 16 <sup>1</sup>Department of Immunology and Microbial Science, The Scripps Research Institute, La Jolla, CA 92037, USA <sup>2</sup>Center for Genome Sciences, U.S. Army Medical Research Institute of Infectious Diseases, Fort Detrick, MD 21702, USA <sup>3</sup>Department of Zoology, University of Oxford, Oxford OX1 3PS, UK <sup>4</sup>Boston Children's Hospital, Boston, MA 02115, USA <sup>5</sup>Harvard Medical School, Boston, MA 02115, USA <sup>6</sup>Vaccine and Infectious Disease Division, Fred Hutchinson Cancer Research Center, Seattle, WA 98109, USA <sup>7</sup>Department of Biological Sciences, College of Arts and Sciences, Florida Gulf Coast University, Fort Myers, FL 33965, USA Bureau of Public Health Laboratories, Division of Disease Control and Health Protection, Florida Department of Health, Miami, FL 33125, USA <sup>9</sup>Department of Pathology, University of Miami Miller School of Medicine, Miami, FL 33136, USA 10 Bureau of Epidemiology, Division of Disease Control and Health Protection, Florida Department of Health, Tallahassee, FL 32399, USA <sup>11</sup>Scripps Translational Science Institute, La Jolla, CA 92037, USA <sup>12</sup>The Broad Institute of MIT and Harvard, Cambridge, MA 02142, USA <sup>13</sup>Institute of Microbiology and Infection, University of Birmingham, Birmingham B15 2TT, UK <sup>14</sup>Department of Microbiology, Immunology, and Pathology, Colorado State University, Fort Collins, CO 80523, USA <sup>15</sup>Li Ka Shing Knowledge Institute, St Michael's Hospital, Toronto, ON M5B 1T8, Canada <sup>16</sup>Division of Infectious Diseases, Department of Medicine, University of Toronto, Toronto, ON M5B 1T8, Canada <sup>17</sup>Institute for Health Metrics and Evaluation, University of Washington, Seattle, WA 98121, USA <sup>18</sup>Division of Infectious Diseases, University of Miami Miller School of Medicine, Miami, FL 33155, USA <sup>19</sup>Bureau of Public Health Laboratories, Division of Disease Control and Health Protection, Florida Department of Health, Tampa, FL 33612, <sup>20</sup>Florida Department of Health in Miami-Dade County, Miami, FL 33125, USA <sup>21</sup>National Center for Atmospheric Research, Boulder, CO 80307, USA <sup>22</sup>Department of Microbiology and Immunology, Tulane University School of Medicine, New Orleans, LA 70112, USA <sup>23</sup>Miami-Dade County Mosquito Control, Miami, FL 33178 USA <sup>24</sup>Department of Biology and Emerging Pathogens Institute, University of Florida, Gainesville, FL 32610, USA <sup>25</sup>Institute of Evolutionary Biology, University of Edinburgh, Edinburgh EH9 3FL, UK <sup>26</sup>Fogarty International Center, National Institutes of Health, Bethesda, MD 20892, USA <sup>27</sup>Center for Systems Biology, Department of Organismic and Evolutionary Biology, Harvard University, Cambridge MA 02138, USA <sup>28</sup>Department of Immunology and Infectious Diseases, Harvard T.H. Chan School of Public Health, Harvard University, Boston MA 02115, USA <sup>29</sup>Howard Hughes Medical Institute, Chevy Chase, MD 20815, USA <sup>30</sup>Department of Integrative Structural and Computational Biology, The Scripps Research Institute, La Jolla, CA 92037, USA

<sup># =</sup> co-senior

<sup>§ =</sup> co-corresponding

- Zika virus (ZIKV) is causing an unprecedented epidemic linked to severe congenital syndromes<sup>1,2</sup>. 56
- 57 In July 2016, mosquito-borne ZIKV transmission was reported in the continental United States and
- since then, hundreds of locally-acquired infections have been reported in Florida<sup>3,4</sup>. To gain insights 58
- 59 into the timing, source, and likely route(s) of ZIKV introduction, we tracked the virus from its first
- 60 detection in Florida by sequencing ZIKV genomes from infected patients and Aedes aegypti
- mosquitoes. We show that at least four introductions, but potentially as many as 40, contributed to 61
- 62 the outbreak in Florida and that local transmission likely started in the spring of 2016 - several
- 63 months before initial detection. By analyzing surveillance and genetic data, we discovered that
- 64 ZIKV moved among transmission zones in Miami. Our analyses show that most introductions are
- 65 linked to the Caribbean, a finding corroborated by the high incidence rates and traffic volumes
- 66 from the region into the Miami area. Our study provides an understanding of how ZIKV initiates
- 67 transmission in new regions.
- ZIKV transmission in the Americas was first reported in Brazil in May 2015<sup>5</sup>, though the virus was likely 68
- 69
- introduced 1-2 years prior to its detection<sup>6-8</sup>. By January 2016, ZIKV cases were reported from several South and Central American countries and most islands in the Caribbean<sup>9</sup>. Like dengue virus (DENV) and chikungunya virus (CHIKV), ZIKV is vectored primarily by *Aedes* mosquitoes<sup>10-13</sup>. The establishment of the peridomestic species *Ae. aegypti* in the Americas<sup>14</sup> has facilitated DENV, CHIKV, and now likely 70
- 71
- 72
- ZIKV to become endemic in this region<sup>15</sup>. In the continental United States, transient outbreaks of DENV and CHIKV have been reported in regions of Texas and Florida<sup>4,16–21</sup> with abundant seasonal *Ae. aegypti* 73
- 74
- populations<sup>14,22</sup>. 75
- 76 The 2016 ZIKV outbreak in Florida generated 256 confirmed ZIKV infections<sup>4</sup> (Fig. 1a). While
- 77 transmission was confirmed across four counties in Florida (Fig. 1b), the outbreak was most intense in
- 78 Miami-Dade County (241 infections). Although the case location could not always be determined, at least
- 79 114 (47%) infections were likely acquired in one of three distinct transmission zones: Wynwood, Miami
- 80 Beach, and Little River (Fig. 1c-d).
- 81 Using mosquito surveillance data, we determined the extent of mosquito-borne ZIKV transmission in
- 82 Miami. Of the 24,351 mosquitoes collected from June to November 2016, 99.8% were Ae. aegypti and 8
- 83 pools of ≤ 50 mosquitoes tested positive for ZIKV (Fig. 1c, Extended Data Fig. 1). From these pools, we
- 84 estimated that ~1 out of 1,600 Ae. aegypti mosquitoes were infected (0.061%, 95% CI: 0.028-0.115%,
- Extended Data Fig. 1a). This is similar to infection rates during DENV and CHIKV outbreaks<sup>23</sup>. Although 85
- 86 we did not detect ZIKV-infected mosquitoes outside Miami Beach (Fig. 1c), we found that the number of
- 87 human ZIKV cases correlated strongly with Ae. aegypti abundance within each transmission zone 88 (Spearman r = 0.61, Fig. 1d, Extended Data Fig. 1b). This suggests that Ae. aegypti mosquitoes were the
- 89 primary mode of transmission and that changes to vector abundance impacted human infection rates. We
- found that the application of insecticides<sup>3</sup> suppressed mosquito populations during periods of intensive 90
- 91 usage (Extended Data Fig. 1c), and therefore likely contributed to ZIKV clearance.
- 92 We sequenced 39 ZIKV genomes from clinical and mosquito samples without cell culture<sup>24</sup>
- 93 (Supplementary Table 1a). Our ZIKV dataset included 29 genomes from patients with locally-acquired
- 94 infections (Fig. 1d) and 7 from Ae. aegypti pools (Fig. 1c). We also sequenced 3 ZIKV genomes from
- 95 travel-associated cases from Florida. Our dataset included cases from all transmission zones in Miami
- 96 (Fig. 1d) and represented ~11% of all confirmed locally-acquired cases in Florida. We made all sequence
- 97 data openly available (PRJNA342539, PRJNA356429) immediately after data generation.
- 98 We reconstructed phylogenetic trees from our ZIKV genomes along with 65 published genomes from
- 99 other affected regions (Fig. 2, Extended Data Fig. 2 and 3). We found that the Florida ZIKV genomes
- 100 formed four distinct lineages (labeled F1-F4, Fig. 2a), three of which (F1-F3) belonged to the same clade
- 101 (labeled A, Fig. 2a). We only sampled a single human case each from the F3 and F4 lineages, consistent

- 102 with limited transmission (Fig. 2a). The other two Florida lineages (F1-F2) comprised ZIKV genomes
- 103 from human and mosquito samples within Miami-Dade County (Fig. 2b).
- Using time-structured phylogenies<sup>25</sup>, we estimated that at least four separate introductions were 104
- responsible for the locally-acquired cases observed in our dataset. The phylogenetic placement of lineage 105
- 106 F4 clearly indicates that it resulted from an independent introduction of a lineage distinct from those in
- 107 clade A (Fig. 2a). For the two well-supported nodes linking lineages F1-F2 (labeled B, Fig. 2a) and F1-F3
- 108 (A, Fig. 2a), we estimated the time of the most recent common ancestor (tMRCA) to be during the
- 109 summer of 2015 (95% highest posterior density [HPD]: June-September, 2015). Our data displayed a
- strong clock signal (Extended Data Fig. 2b) and tMRCA estimates were robust across a range of models 110
- (Extended Data Table 1a). Thus while F1-F3 belong to clade A, any fewer than three distinct 111
- 112 introductions leading to these lineages would have required undetected transmission of ZIKV in Florida
- 113 for approximately one year (Fig. 2a).
- 114 To estimate the likelihood of a single ZIKV transmission chain persisting for over a year, we modeled
- spread under different assumptions of the basic reproductive number  $(R_0)$ . Using the number of locally-115
- 116 acquired and travel-associated cases, along with the number of observed genetic lineages, we estimated an
- 117  $R_0$  between 0.5 and 0.8 in Miami-Dade County (Extended Data Fig. 4). Even at the upper end of this
- 118 range, the probability of a single transmission chain persisting for over a year is extremely low ( $\sim 0.5\%$ ,
- 119 Fig. 2c). This is especially true considering the low Ae. aegypti abundance during the winter months
- 120 (Extended Data Fig. 1d).
- 121 Given the low probability of long-term persistence, we expect that our ZIKV genomes (F1-F4) were the
- 122 result of at least four introductions. Differences in surveillance practices and a high number of travel-
- 123 associated cases (Fig. 1a), however, likely mean that unsampled ZIKV introductions also contributed to
- 124 the outbreak. To estimate the total number of ZIKV introductions, we modeled scenarios that resulted in
- 125 241 locally-acquired cases within Miami-Dade County, and found that with  $R_0$  values of 0.5-0.8, we
- expect 17-42 (95% CI 3-63) separate introductions to have contributed to the outbreak (Fig. 2d). The 126
- 127 majority of these introductions would likely have generated a single secondary case that was undetected
- 128 in our genetic sampling (Extended Data Fig. 4a). Incorporating under-reporting in a sensitivity analysis
- 129 increases  $R_0$  estimates slightly to 0.7-0.9 (Extended Data Fig. 4f-i).
- 130 The two main ZIKV lineages, F1 and F2, included the majority of genomes from Florida (92%, Fig. 2a).
- Assuming they represent two independent introductions, we estimated when each of these lineages 131
- 132 arrived in Florida. The probability densities for the tMRCAs of both F1 and F2 were centered around
- 133 March-April, 2016 (Fig. 2b, 95% HPD: January-May, 2016). The estimated timing for these introductions
- corresponds with suitable Ae. aegypti populations in Miami-Dade County<sup>26</sup> (Extended Data Fig. 1d) and 134
- suggests that ZIKV transmission could have started at least two months prior to its detection in July 2016 135
- 136 (Fig. 1a). The dates of the introductions could be more recent if multiple F1 or F2 lineage viruses arrived
- 137 independently. However, more than 2 introductions would be necessary to substantially change our
- 138 estimates for the timing of the earliest introduction.
- 139 To understand transmission dynamics within Miami, we analyzed our genomic data together with case
- 140 data from the Florida Department of Health (DOH, Supplementary Table 1a). While spatially distinct, the
- 141 three ZIKV transmission zones occurred within ~5 km of each other (Fig. 1c) and we found that the
- 142 ZIKV infections associated with each zone overlapped temporally (Fig. 1d). Our ZIKV genomes with
- 143 zone assignments all belonged to lineages F1 and F2, but neither of these lineages were confined to a
- 144 single zone (Fig. 2b). In fact, we detected both F1 and F2 lineage viruses from Ae. aegypti collected from
- 145 the same trap 26 days apart (mosquitoes 5 and 8, Fig. 2b). These findings suggest that ZIKV moved
- 146 among areas of Miami.

Determining the sources and routes of ZIKV introductions could help mitigate future outbreaks. We

found that lineages F1-F3 clustered with ZIKV genomes sequenced from the Dominican Republic and

Guadeloupe (Fig. 2, Extended Data Fig. 2 and 3). In contrast, F4 clustered with genomes from Central

America (Fig. 2, Extended Data Fig. 2 and 3). These findings suggest that while ZIKV outbreaks occurred

throughout the Americas, the Caribbean islands were the main source of establishing local ZIKV

transmission in Florida. Because of severe undersampling of ZIKV genomes, however, we cannot rule out

other source areas. Similarly, even though we found that the Florida ZIKV genomes clustered together

154 with sequences from the Dominican Republic, our results do not prove that ZIKV entered Florida from

this country.

173

156 We investigated ZIKV infection rates and travel patterns to corroborate our phylogenetic evidence for 157 Caribbean introductions. We found that the Caribbean islands bore the highest ZIKV incidence rates (Fig. 158 2b), despite Brazil and Colombia reporting the highest absolute number of cases (January to June, 2016, 159 Fig. 3a, Extended Data Fig. 5, Supplementary Table 1b). During the same time period, we estimated that ~3 million travelers arrived from the Caribbean, accounting for 54% of the total traffic into Miami, with 160 the vast majority (~2.4 million) arriving via cruise ships (Fig. 3b, Extended Data Fig. 6, Supplementary 161 162 Table 1b). Combining the infection rates with travel capacities, we estimated that ~60-70% of ZIKV 163 infected travelers arrived from the Caribbean (Fig. 3c and Extended Data Fig. 7a). We also found that the 164 number of travel-associated ZIKV cases correlated strongly with the expected number of importations 165 from the Caribbean (Spearman r = 0.8, Fig. 3d, Extended Data Fig. 7b). Finally, 67% of the travel-166 associated infections in Florida reported recent travel to the Caribbean (Fig. 3e); however, their mode of 167 travel is unknown. Taken together, these findings suggest that a high incidence of ZIKV in the Caribbean, 168 combined with frequent travel, could have played a key role in the establishment of ZIKV transmission in 169 Florida. These findings, however, do not indicate that cruise ships themselves are risk factors for human 170 ZIKV infection, but only that they served as a major mode of transportation from areas with active 171 transmission. In addition, ZIKV exposure may vary among individuals depending on their purpose of 172 travel and therefore we cannot determine the specific contribution of ZIKV-infected travelers arriving via

174 The majority of the Florida ZIKV outbreak occurred in Miami-Dade County (Fig. 1b). To determine if 175 there is a higher potential for ZIKV outbreaks in this area, we analyzed incoming passenger traffic from 176 regions with ZIKV transmission along with local Ae. aegypti abundance. We estimated that Miami and 177 nearby Fort Lauderdale received ~72% of traffic (Fig. 4) and Miami received more air and sea traffic 178 from ZIKV endemic areas than any other city in the United States (Extended Data Fig. 8). During January 179 to April 2016, we estimated that Ae. aegypti abundance was highest in southern Florida<sup>22</sup> (Fig. 4, Extended Data Fig. 1d, Extended Data Fig. 8). By June, most of Florida and several cities across the South likely supported high *Ae. aegypti* populations<sup>14,22</sup> (Extended Data Fig. 8); however, most of this 180 181 region has not reported local Ae. aegypti-borne virus transmission in at least 60 years 19. In fact, the only 182 region outside of Florida with local ZIKV transmission is southern Texas<sup>27</sup>, which is also the only other region with recent DENV outbreaks<sup>19–21</sup>. Therefore, the combination of travelers, mosquito ecology, and 183 184 185 human population density likely make Miami one of the few places in the continental United States at risk

186 for Ae. aegypti-borne virus outbreaks<sup>22,26,28</sup>.

airlines or cruise ships.

The extent of ZIKV transmission in Florida was unprecedented, with more reported ZIKV cases in 2016 (256) than DENV cases since 2009 (136)<sup>4,16,17</sup>. This case difference may be reflected by lower incidence of endemic DENV than epidemic ZIKV in source countries<sup>29,30</sup>, resulting in fewer DENV importations (reported travel cases since 2009: 654 DENV and 1,016 ZIKV)<sup>4</sup>. Given that the majority of ZIKV infections are asymptomatic<sup>2,31</sup>, the true number of ZIKV cases was likely much higher. Despite this, we estimated that the average  $R_0$  was less than 1 and therefore multiple introductions were necessary to give rise to the observed outbreak<sup>32</sup>. The high volume of traffic entering Florida from ZIKV-affected regions, especially the Caribbean, likely provided a substantial supply of ZIKV-infected individuals<sup>33</sup>. Because

Florida is unlikely to sustain long-term ZIKV transmission<sup>32</sup>, the potential for future ZIKV outbreaks in this region is dependent upon activity elsewhere. Therefore, we expect that outbreaks in Florida will cycle with the ZIKV transmission dynamics in the Americas<sup>7,8,15</sup>.

198

200

201

#### Acknowledgements

202

203 We thank J. Weger-Lucarelli, G. Ebel, C. Moore, B. Alto, G. Donatti, and S. Taylor for discussions, E. 204 Spencer for IRB and logistics support, M. Pilcher for sequencing assistance, and G. Schroth and S. Gross 205 for designing and providing enrichment probes. N.D.G. is supported by NIH training grant 206 5T32AI007244-33. G.D. is supported by the Mahan Postdoctoral Fellowship from the Computational 207 Biology Program at Fred Hutch. K.G.B. is supported by the ASTMH Shope Fellowship. N.R.F. is funded 208 by a Sir Henry Dale Fellowship (Wellcome Trust/Royal Society Grant 204311/Z/16/Z). D.A.T.C. was 209 supported by US NIH MIDAS program (U54-GM088491) and CDC Cooperative Agreement 210 U01CK000510, A.R. is supported by EU Seventh Framework Programme (FP7/2007-2013) under Grant 211 278433-PREDEMICS, ERC Grant 260864, Horizon 2020 Grant 643476-COMPARE. T.B. is a Pew 212 Biomedical Scholar and is supported by NIH R35 GM119774-01. O.G.P. received funding from EU ERC 213 Seventh Framework Programme (FP7/2007-2013)/ERC number 614725-PATHPHYLODYN and the 214 USAID Emerging Pandemic Threats Program-2 PREDICT-2 (Cooperative Agreement No. AID-OAA-A-14-00102). S.I. and S.F.M. are supported by NIH NIAID 4R01AI099210-04. ZIKV sequencing at 215 216 USAMRIID was supported by DARPA (PI: C. Kane). K.G.A. is a Pew Biomedical Scholar, and is 217 supported by NIH NCATS CTSA UL1TR001114, NIAID contract HHSN272201400048C, and The Ray 218 Thomas Foundation. The content of this publication does not necessarily reflect the views or policies of 219 the US Army, the Department of Health and Human Services, the CDC, or the Florida DOH.

#### 220 **Author Contributions**

- 221 All contributions are listed in order of authorship. Designed the experiments: N.D.G., J.T.L., G.D.,
- 222 M.U.G.K., D.A.T.C., P.C.S, L.D.G., S.F.M., T.B., O.G.P., S.I., G.P., and K.G.A.; Collected samples:
- 223 A.L.T., S.W., D.M.M., A.B., L.M.P., D.P., P.N.L., M.R., V.K.B., D.I.W., M.R.C., E.W.K., K.N.H.,
- 224 A.C.C., R.J., M.C.P., C.V., D.S., L.D.G., S.F.M., and S.I.; Performed the sequencing: N.D.G., M.W.R.,
- 225 K.P., D.R., R.R.-S., G.O., and E.N.; Provided data, reagents, or protocols: N.D.G., J.T.L., G.D.,
- 226 M.U.G.K., K.G., M.R.W., R.R.-S., G.O., H.C.M., M.L.B., K.G.B., B.C., C.A.F., A.G.-Y., A.G., C.L.,
- 227 B.M., C.B.M., D.J.P., J.Q., S.F.S., C.T.-T., K.L.M., S.M.W., S.W., N.L.Y., J.Q., J.R.F., K.K., S.E.B.,
- 228 A.J.M., R.F.G., N.J.L., M.C.P., C.V., P.C.S., S.F.M., and S.I.; Analyzed the data: N.D.G., J.T.L., G.D.,
- 229
- M.U.G.K., K.G., J.T., J.R.F., R.C.R., N.R.F., D.A.T.C., A.K., M.S.-L., T.B., S.F.M, O.G.P., S.I., and
- 230 K.G.A.; Edited manuscript: G.D., M.U.G.K., J.T., S.F.S., A.R., T.B., O.G.P., S.I., and G.P.; Wrote
- 231 manuscript: N.D.G., J.T.L., and K.G.A.; All authors read and approved the manuscript.

#### 232 **Author Information**

- 233 The authors declare no competing financial interests. Correspondence and requests for materials should
- 234 be addressed to K.G.A (andersen@scripps.edu) or G.P. (gustavo.f.palacios.ctr@mail.mil).

#### References

- 281 1. Zika virus and complications. World Health Organization Available at:
- http://www.who.int/features/qa/zika/en/. (Accessed: 1st November 2016)
- 283 2. Lazear, H. M. & Diamond, M. S. Zika Virus: New Clinical Syndromes and Its Emergence in the
- 284 Western Hemisphere. J. Virol. **90**, 4864–4875 (2016).
- 285 3. Likos, A. et al. Local Mosquito-Borne Transmission of Zika Virus Miami-Dade and Broward
- 286 Counties, Florida, June-August 2016. MMWR Morb. Mortal. Wkly. Rep. 65, 1032–1038 (2016).
- 4. Mosquito-Borne Disease Surveillance. Florida Department of Health Available at:
- http://www.floridahealth.gov/diseases-and-conditions/mosquito-borne-diseases/surveillance.html.
- 289 (Accessed: 10th January 2017)
- 290 5. Hennessey, M., Fischer, M. & Staples, J. E. Zika Virus Spreads to New Areas Region of the
- 291 Americas, May 2015–January 2016. MMWR Morb. Mortal. Wkly. Rep. 65, 1–4 (2016).
- 292 6. Faria, N. R. et al. Zika virus in the Americas: Early epidemiological and genetic findings. Science
- **352**, 345–349 (2016).
- 7. Faria, N. R., Quick, J., Morales, I., Theze, J. & de Jesus, J. G. Epidemic establishment and cryptic
- transmission of Zika virus in Brazil and the Americas. *bioRxiv* (2017).
- 8. Metsky, H. C. *et al.* Genome sequencing reveals Zika virus diversity and spread in the Americas.
- 297 bioRxiv 109348 (2017). doi:10.1101/109348
- 298 9. Regional Zika Epidemiological Update (Americas). Pan American Health Organization (2016).
- 299 Available at:
- http://www.paho.org/hq/index.php?option=com\_content&view=article&id=11599&Itemid=41691&l
- ang=en. (Accessed: 1st December 2016)
- 302 10. Weger-Lucarelli, J. et al. Vector Competence of American Mosquitoes for Three Strains of Zika
- 303 Virus. *PLoS Negl. Trop. Dis.* **10**, e0005101 (2016).
- 304 11. 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.* (2016).

- 306 12. 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* 0 (2016).
- 308 13. Chouin-Carneiro, T. et al. Differential Susceptibilities of Aedes aegypti and Aedes albopictus from
- the Americas to Zika Virus. *PLoS Negl. Trop. Dis.* **10**, e0004543 (2016).
- 310 14. Kraemer, M. U. G. et al. The global distribution of the arbovirus vectors Aedes aegypti and Ae.
- 311 albopictus. *Elife* **4**, e08347 (2015).
- 312 15. Ferguson, N. M. et al. Countering the Zika epidemic in Latin America. Science 353, 353–354
- 313 (2016).
- 314 16. Teets, F. D. et al. Origin of the dengue virus outbreak in Martin County, Florida, USA 2013. Virol
- 315 Rep **1-2**, 2–8 (2014).
- 316 17. Graham, A. S. et al. Mosquito-associated dengue virus, Key West, Florida, USA, 2010. Emerg.
- 317 *Infect. Dis.* **17,** 2074–2075 (2011).
- 318 18. Kendrick, K. et al. Notes from the field: transmission of chikungunya virus in the continental United
- 319 States—Florida, 2014. MMWR Morb. Mortal. Wkly. Rep. 63, 1137 (2014).
- 320 19. Bouri, N. et al. Return of epidemic dengue in the United States: implications for the public health
- 321 practitioner. *Public Health Rep.* **127**, 259–266 (2012).
- 322 20. Ramos, M. M. et al. Epidemic Dengue and Dengue Hemorrhagic Fever at the Texas–Mexico Border:
- Results of a Household-based Seroepidemiologic Survey, December 2005. Am. J. Trop. Med. Hyg.
- **78,** 364–369 (2008).
- 325 21. Murray, K. O. et al. Identification of dengue fever cases in Houston, Texas, with evidence of
- autochthonous transmission between 2003 and 2005. Vector Borne Zoonotic Dis. 13, 835–845
- 327 (2013).
- 328 22. Monaghan, A. J. et al. On the Seasonal Occurrence and Abundance of the Zika Virus Vector
- Mosquito Aedes Aegypti in the Contiguous United States. *PLoS Curr.* **8**, (2016).
- 23. Dzul-Manzanilla, F. et al. Evidence of vertical transmission and co-circulation of chikungunya and
- dengue viruses in field populations of Aedes aegypti (L.) from Guerrero, Mexico. *Trans. R. Soc.*

- 332 *Trop. Med. Hyg.* **110**, 141–144 (2016).
- 333 24. Quick, J. et al. Multiplex PCR method for MinION and Illumina sequencing of Zika and other virus
- genomes directly from clinical samples. *bioRxiv* 098913 (2017).
- 25. Drummond, A. J., Suchard, M. A., Xie, D. & Rambaut, A. Bayesian phylogenetics with BEAUti and
- 336 the BEAST 1.7. *Mol. Biol. Evol.* **29**, 1969–1973 (2012).
- 337 26. Robert, M. A. et al. Modeling Mosquito-Borne Disease Spread in U.S. Urbanized Areas: The Case
- of Dengue in Miami. *PLoS One* **11**, e0161365 (2016).
- 27. McCarthy, M. First US case of Zika virus infection is identified in Texas. BMJ 352, i212 (2016).
- 340 28. Nelson, B. et al. Travel Volume to the United States from Countries and U.S. Territories with Local
- Zika Virus Transmission. *PLoS Curr.* **8,** (2016).
- 342 29. Dengue in Puerto Rico. The Centers for Disease Control and Prevention Available at:
- https://www.cdc.gov/dengue/about/inpuerto.html. (Accessed: 17th March 2017)
- 34. Zika-Epidemiological Report. Pan American Health Organization Available at:
- http://www.paho.org/hq/index.php?option=com\_content&view=article&id=11603&Itemid=41696&I
- ang=en. (Accessed: 1st December 2016)
- 31. Duffy, M. R. et al. Zika virus outbreak on Yap Island, Federated States of Micronesia. N. Engl. J.
- 348 *Med.* **360**, 2536–2543 (2009).

- 34. Dinh, L., Chowell, G., Mizumoto, K. & Nishiura, H. Estimating the subcritical transmissibility of the
- Zika outbreak in the State of Florida, USA, 2016. *Theor. Biol. Med. Model.* 13, 20 (2016).
- 35. Nunes, M. R. T. et al. Air travel is associated with intracontinental spread of dengue virus serotypes
- 352 1-3 in Brazil. *PLoS Negl. Trop. Dis.* **8,** e2769 (2014).
- 353 34. Basemaps, ESRI Available at: http://www.esri.com/data/basemaps. (Accessed: 1st October 2016)

## Figure Legends

**Figure 1 | Zika virus outbreak in Florida.** (a) Weekly counts of confirmed travel-associated and locally-acquired ZIKV cases in 2016. (b) Four counties reported locally-acquired ZIKV cases in 2016: Miami-Dade (241), Broward (5), Palm Beach (8), Pinellas (1), and unknown origin (1). (c) The locations of mosquito traps and collected *Ae. aegypti* mosquitoes found to contain ZIKV RNA (ZIKV+) in relation to the transmission zones within Miami. (d) Temporal distribution of weekly ZIKV cases (left y-axis), sequenced cases (bottom), and *Ae. aegypti* abundance per trap night (right y-axis) associated with the three described transmission zones. ZIKV cases and sequences are plotted in relation to symptom onset dates (n=18). Sequenced cases without onset dates or that occurred outside of the transmission zones are not shown (n=10). Human cases and *Ae. aegypti* abundance per week were positively correlated (Spearman r = 0.61, Extended Data Fig. 1b). The maps were generated using open source basemaps<sup>34</sup>.

Figure 2 | Multiple introductions of Zika virus into Florida. (a) Maximum clade credibility (MCC) tree of ZIKV genomes sequenced from outbreaks in the Pacific islands and the epidemic in the Americas. Tips are colored based on collection location. The five tips outlined in blue but filled with a different color indicate ZIKV cases in the United States associated with travel (fill color indicates the probable location of infection). Clade posterior probabilities are indicated by white circles filled with black relative to the level of support. The grey violin plot indicates the 95% highest posterior density (HPD) interval for the tMRCA for the epidemic in the Americas (AM). Lineage F4 contains two identical ZIKV genomes from the same patient. (b) A zoomed in version of the whole MCC tree showing the collection locations of Miami-Dade sequences and whether they were sequenced from mosquitoes (numbers correspond to trap locations in Fig. 1c). 95% HPD intervals are shown for the tMRCAs (c) The probability of ZIKV persistence after introduction for different  $R_0$ . Persistence is measured as the number of days from initial introduction of viral lineages until their extinction. Vertical dashed lines show the inferred mean persistence time for lineages F1, F2 and B based on their tMRCA. (d) Total number of introductions (mean with 95% CI) that contributed to the outbreak of 241 local cases in Miami-Dade County for different  $R_0$ .

Figure 3 | Frequent opportunities for Zika virus introductions into Miami from the Caribbean. (a) Reported ZIKV cases per country/territory from January to June, 2016 normalized by total population. (b) The number of estimated travelers entering Miami during January to June, 2016 by method of travel. (c) The number of travelers and the reported ZIKV incidence rate for the country/territory of origin were used to estimate the proportion of infected travelers coming from each region with ZIKV in the Americas. (d) The observed number of weekly travel-associated ZIKV cases in Florida were plotted with the expected number of ZIKV-infected travelers (as estimated in panel c) coming from all of the Americas (grey line) and the regional contributions (colored areas). (e) The countries visited by the 1,016 travel-associated ZIKV cases diagnosed in Florida.

**Figure 4 | Southern Florida has a high potential for** *Aedes aegypti***-borne virus outbreaks**. The estimated number of travelers per month (circles) entering Florida cities via flights and cruise ships were plotted with estimated relative *Ae. aegypti* abundance. Only cities receiving >10,000 passengers per month are shown. Relative *Ae. aegypti* abundance for every month is shown in Extended Data Fig. 1d.

Fig 1.

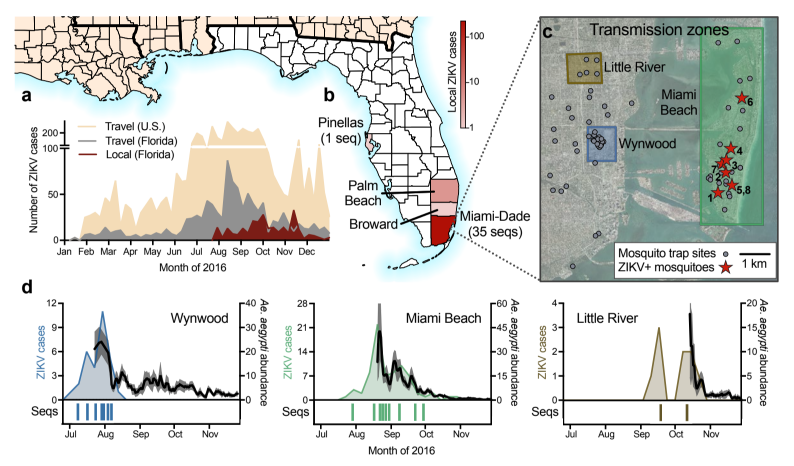


Fig 2.

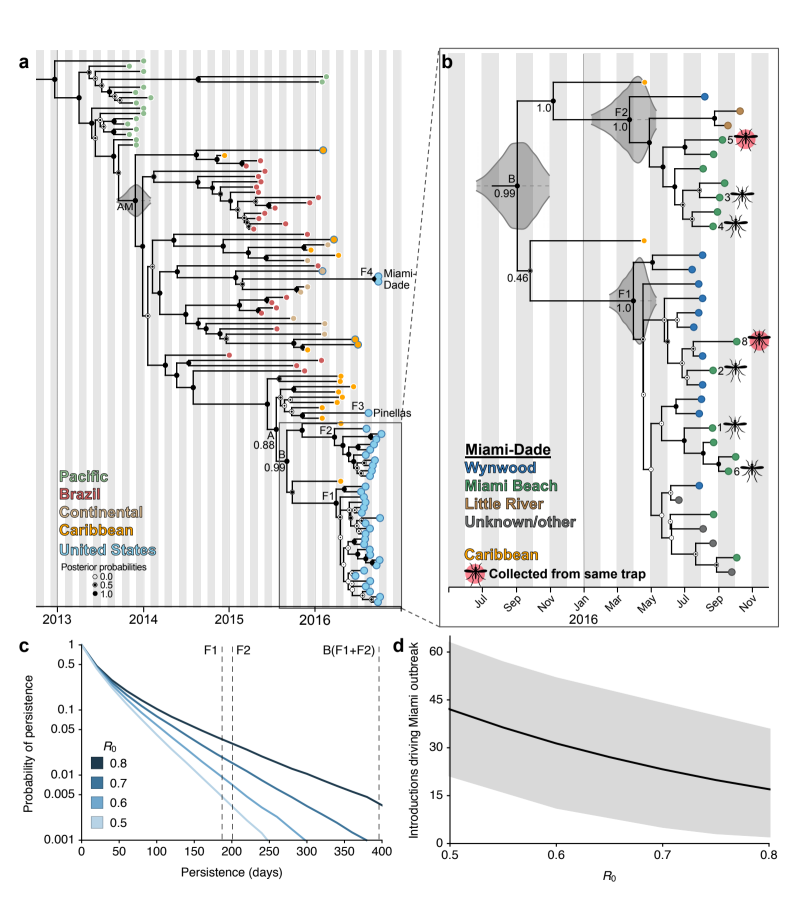


Fig 3.

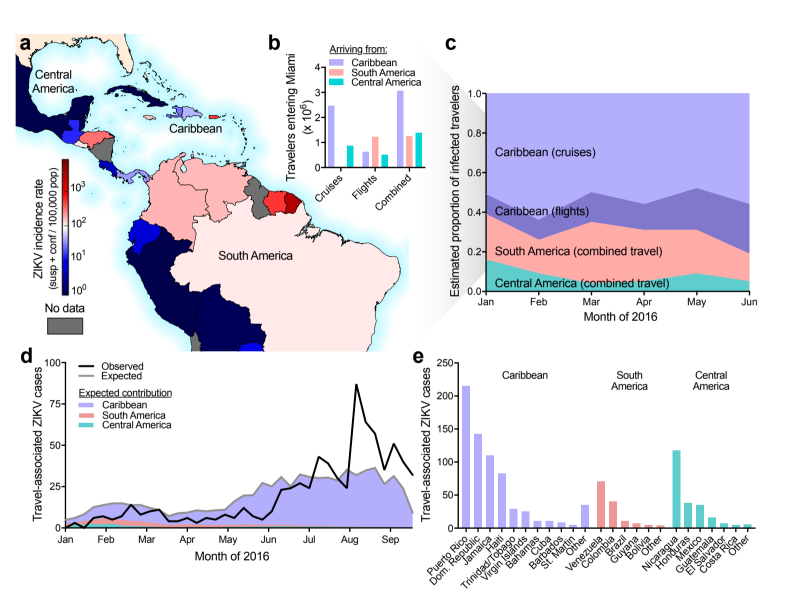
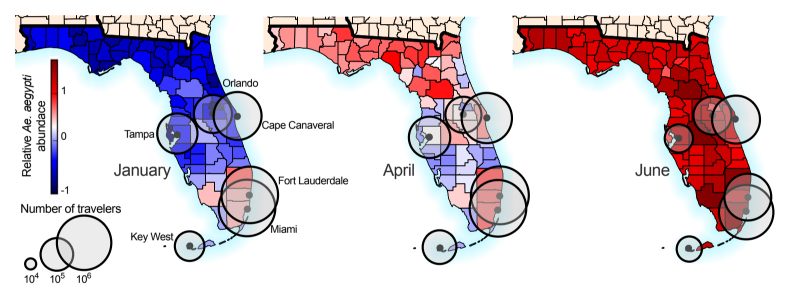


Fig 4.



#### Methods

355

365

377

388

#### 356 **Ethical statement**

- 357 This work was evaluated and approved by relevant Institutional Review Boards (IRB)/Ethics Review
- 358 Committees at The Scripps Research Institute (TSRI) and the US Army Medical Research Institute of
- 359 Infectious Diseases (USAMRIID) Office of Human Use and Ethics. This work was conducted as part of
- 360 the public health response in Florida and samples were collected under a waiver of consent granted by the
- Florida DOH Human Research Protection Program. The work received a non-human subjects research 361
- designation (category 4 exemption) by the Florida DOH since this research was performed with leftover 362
- 363 clinical diagnostic samples involving no more than minimal risk. All samples were deidentified prior to
- 364 receipt by the study investigators.

#### Florida Zika virus case data

- 366 Weekly reports of international travel-associated and locally-acquired ZIKV infections diagnosed in
- 367 Florida were obtained from the Florida DOH mosquito-borne disease surveillance system<sup>4</sup>. Dates of
- symptom onset from the Miami transmission zones (Wynwood, Miami Beach, and Little River) 368
- determined by the Florida DOH investigation process were obtained from the ZIKV resource website<sup>35</sup> 369
- and daily updates<sup>36</sup>. International travel-associated ZIKV case counts in the United States (outside of 370
- Florida) were obtained from the CDC<sup>37</sup>. The local and travel-associated ZIKV case numbers for Florida 371
- were obtained from the Florida DOH. The one local ZIKV infection diagnosed in Duval County was 372
- believed to have originated elsewhere in Florida. Therefore, this case is listed as "unknown origin" in Fig. 373
- 374 1b. In Fig. 3e, only the countries visited by 5 or more times by ZIKV-infected travelers diagnosed in
- Florida are shown. Countries with 5 or fewer visits were aggregated into an "other" category by region 375
- 376 (i.e., Caribbean, South America, or Central America).

#### Clinical sample collection and RNA extraction

- 378 Clinical samples from locally-acquired ZIKV infections were collected from June 22 to October 11, 2016.
- 379 The Florida DOH identified persons with compatible illness and clinical samples were shipped to the
- 380 Bureau of Public Health Laboratories for confirmation by qRT-PCR and antibody tests following interim
- guidelines<sup>3,38-40</sup>. Clinical specimens (whole blood, serum, saliva, or urine) submitted for analysis were 381
- 382 refrigerated or frozen at ≤ -70°C until RNA was extracted. RNA was extracted using the RNAeasy kit
- 383 (QIAGEN), MagMAX for Microarrays Total RNA Isolation Kit (Ambion), or MagNA Pure LC 2.0 or 96
- Systems (Roche Diagnostics). Purified RNA was eluted into 50-100 µL using the supplied elution 384
- buffers, immediately frozen at ≤ -70°C, and transported on dry ice. The Florida DOH also provided 385
- 386 investigation data for these samples, including symptom onset dates and, when available, assignments to
- 387 the zone where infection likely occurred (Supplementary Table 1).

### Mosquito collection, RNA extraction, and entomological data analysis

- 389 24,351 Ae. aegypti and Ae. albopictus mosquitoes (sorted into 2,596 pools) were collected throughout
- 390 Miami-Dade County during June to November, 2016 using BG-Sentinel mosquito traps (Biogents AG).
- 391 Up to 50 mosquitoes of the same species and sex were pooled per trap. The pooled mosquitoes were
- 392 stored in RNAlater (Invitrogen), RNA was extracted using either the RNAeasy kit (QIAGEN) or
- 393 MagMAX for Microarrays Total RNA Isolation Kit (Ambion), and ZIKV RNA was detected by qRT-
- PCR targeting the envelope protein coding region<sup>40</sup> or the Trioplex qRT-PCR kit<sup>41</sup>. ZIKV infection rates 394
- were calculated per 1,000 female Ae. aegypti mosquitoes using the bias-corrected maximum likelihood 395
- estimate (MLE)<sup>42</sup>. Days of insecticide usage by the Miami-Dade Mosquito Control were inferred from the 396
- zone-specific ZIKV activities timelines published by the Florida DOH<sup>3</sup> 397

### Relative monthly Ae. aegypti abundance

For the purpose of this study we used Ae. aegypti suitability maps from Kraemer et al. And derived monthly estimates based on the statistical relationships between mosquito presence and environmental correlates Following Hwang et al. We used a simple mathematical formula to transform the probability of detection maps into mosquito abundance maps. In order to do so, we assumed P (Y=1) where Y is a binary variable (presence/absence). Using a Poisson distribution X() to govern the abundance of mosquitoes, the probability of not observing any mosquitoes can be related to the probability of absence as: P(X=0)=P(Y=0). We used the following transformation to generate abundance AO0 estimates per county in Florida:

$$e^{-\lambda} = P(Y = 0)$$
$$\lambda = -\log(P(Y = 0))$$
$$\lambda = -\log(1 - P(Y = 1))$$

We did not consider *Ae. albopictus* abundance in this study because 99.8% of mosquitoes collected in Miami-Dade County were *Ae. aegypti*. Relative Ae. aegypti abundance in major U.S. cities presented in Extended Data Fig. 8 was estimated as previously described<sup>22</sup>.

# 410 Zika virus quantification

398

434

411 ZIKV genome equivalents (GE) were quantified by qRT-PCR. At TSRI, ZIKV qRT-PCR was performed 412 as follows: ZIKV RNA standards were transcribed from the ZIKV NS5 region (8651-9498 nt) using the 413 T7 forward primer (5' - TAA TAC GAC TCA CTA TAG GGA GA TCA GGC TCC TGT CAA AAC CC - 3'), reverse primer (5' - AGT GAC AAC TTG TCC GCT CC - 3'), and the T7 Megascript kit 414 415 (Ambion). For qRT-PCR, primers and a probe targeting the NS5 region (9014-9123 nt) were designed 416 using the ZIKV isolate PRVABC59 (GenBank: KU501215); forward primer (5'- AGT GCC AGA GCT GTG TGT AC - 3'), reverse primer (5' - TCT AGC CCC TAG CCA CAT GT - 3'), and FAM-fluorescent 417 418 probe (5' - GGC AGC CGC GCC ATC TGG T - 3'). The qRT-PCR assays were performed in 25 µl 419 reactions using the iScript One-step RT-PCR Kit for probes (Bio-Rad Laboratories Inc.) and 2 µl of 420 sample RNA. Amplification was performed at 50°C for 20 min, 95°C for 3 min, and 40 cycles of 95°C 421 for 10 s and 57°C for 10 s. Fluorescence was read at the end of the 57°C annealing-extension step. 10-fold 422 dilutions of the ZIKV RNA transcripts (2 µl/reaction) were used to create a standard curve for 423 quantification of ZIKV GE/µl of RNA. The lower limits of quantification are 4 GE/µl RNA, or at a cycle 424 threshold of ~36.

ZIKV GE were quantified at USAMRIID using the University of Bonn ZIKV envelope protein (Bonn E) 425 qRT-PCR assay<sup>45</sup>. RNA standards were transcribed using an amplicon generated from a ZIKV plasmid 426 427 containing T7 promoter at the start of the 5' untranslated region (UTR). The plasmid was designed using 428 the ZIKV isolate BeH819015 (GenBank: KU365778.1) and the amplicon included nts 1-4348, which 429 covers the 5' UTR, C, prM, M, E, NS1, and NS2 regions. The qRT-PCR assays were performed in 25 ul reactions using the SuperScript III platinum One-step qRT-PCR Kit (ThermoFisher) and 2 ul of sample 430 RNA was used. Amplification was performed following conditions as previously described<sup>45</sup>. 10-fold 431 432 dilutions of the ZIKV RNA transcripts (5 µl/reaction) were used to create a standard curve for 433 quantification of ZIKV GE/µl of RNA.

#### Amplicon-based Zika virus sequencing

2IKV sequencing at TSRI was performed using an amplicon-based approach using the ZikaAsian V1 scheme, as described<sup>24</sup>. This approach is similar to "RNA jackhammering" to sequence low-quality viral samples developed by Worobey *et al.*<sup>46</sup>. Briefly, cDNA was reverse transcribed from 5 μl of RNA using SuperScript IV (Invitrogen). ZIKV cDNA (2.5 μl/reaction) was amplified in 35 × 400 bp fragments from two multiplexed PCR reactions using O5 DNA High-fidelity Polymerase (New England Biolabs). The

- amplified ZIKV cDNA fragments (50 ng) were prepared for sequencing using the Kapa Hyper prep kit 440
- 441 (Kapa Biosystems) and SureSelect XT2 indexes (Agilent). Agencourt AMPure XP beads (Beckman
- 442 Coulter) were used for all purification steps. Paired-end 251 nt reads were generated on the MiSeq using
- 443 the V2 500 cycle or V3 600 cycle kits (Illumina).
- 444 Trimmomatic was used to remove primer sequences (first 22 nt from the 5' end of the reads, which is the
- 445 maximum length of the primers used for the multiplexed PCR) and bases at both ends with Phred quality
- score < 20<sup>47</sup>. The reads were then aligned to the complete genome of a ZIKV isolate from the Dominican 446
- 447 Republic, 2016 (GenBank: KU853012) using Novoalign v3.04.04 (www.novocraft.com). Samtools was
- used to sort the aligned BAM files and to generate alignment statistics<sup>48</sup>. Snakemake was used as the 448
- workflow management system<sup>49</sup>. The code and reference indexes for the pipeline can be found at 449
- https://github.com/andersen-lab/zika-pipeline. ZIKV-aligned reads were visually inspected using 450
- Geneious v9.1.5<sup>50</sup> before generating consensus sequences. A minimum of 3× read-depth coverage, in 451
- 452 support of the consensus, was required to make a base call.

#### **Enrichment-based Zika virus sequencing**

453

472

- 454 ZIKV sequencing at USAMRIID was performed using a targeted enrichment approach. Sequencing
- 455 libraries were prepared using the TruSeq RNA Access Library Prep kit (Illumina) with custom ZIKV
- 456 probes. The set included 866 unique probes each of which was 80 nt in length (Supplementary Table 2a).
- The probes were designed to cover the entire ZIKV genome and to encompass the genetic diversity 457
- 458 present on GenBank on January 14, 2016. In total, 26 ZIKV sequences were used during probe design
- 459 (Supplementary Table 2b). Extracted RNA was fragmented at 94 °C for 0-60 s and each sample was
- 460 enriched separately using a quarter of the reagents specified in the manufacturer's protocol. Samples were
- 461 barcoded, pooled and sequenced using the MiSeq Reagent kit v3 (Illumina) on an Illumina MiSeq with a
- minimum of  $2 \times 151$  bp reads. Dual indexing, with no overlapping indices, was used. 462
- The random hexamer associated with read one and the Illumina adaptors were removed from the 463
- sequencing reads using Cutadapt v1.9.dev1<sup>51</sup>, and low-quality reads/bases were filtered using Prinseq-lite 464
- 465 v0.20.3<sup>52</sup>. Reads were aligned to a reference genome (GenBank: KX197192.1) using Bowtie2 v2.0.6<sup>53</sup>,
- duplicates were removed with Picard (http://broadinstitute.github.io/picard), and a new consensus was 466
- $v0.1.18^{48}$ 467 generated using combination of Samtools custom scripts
- 468 (https://github.com/itladner/Scripts/blob/master/reference-based assembly/consensus fasta.py).
- 469 bases with Phred quality score ≥ 20 were utilized in consensus calling, and a minimum of 3× read-depth
- coverage, in support of the consensus, was required to make a call; positions lacking this depth of 470
- 471 coverage were treated as missing (i.e. called as "N").

#### Validation and comparison of sequencing methods

- 473 The consensus ZIKV sequences from FL01M and FL03M generated by sequencing 35 × 400 bp
- 474 amplicons on the MiSeq were validated using the following approaches: 1) sequencing the 35 × 400 bp
- amplicons on the Ion S5 platform (ThermoFisher), 2) sequencing amplicons generated using an Ion 475
- AmpliSeq® (ThermoFisher) panel customly targeted towards ZIKV on the Ion S5 platform, and 3) 476
- 477 sequencing 5 × 2,150-2,400 bp ZIKV amplicons on the MiSeq. For Ion library preparation, cDNA was
- 478 synthesized using the SuperScript VILO kit (ThermoFisher). ThermoFisher designed 875 custom ZIKV
- 479 primers to produce 75 amplicons of ~200 bp in two PCR reactions for use with their Ion AmpliSeq
- 480 Library Kit 2.0. The reagent FuPa was used to digest the modified primer sequences after amplification.
- 481 The DNA templates were loaded onto Ion 520 chips using the Ion Chef and sequenced on the Ion S5 with
- the 200 bp output (ThermoFisher). The  $35 \times 400$  bp amplicons generated for the MiSeq as described 482
- 483 above were introduced into the Ion workflow using the Ion AmpliSeq Library Kit 2.0, but without
- 484 fragmentation. Primers to amplify 2,150-2,400 bp ZIKV fragments (Supplementary Table 2c) were kindly
- 485 provided by Shelby O'Connor, Dawn Dudly, Dave O'Connor, and Dane Gellerup (AIDS Vaccine
- 486 Research Laboratory, University of Wisconsin, Madison). Each fragment was amplified individually by

- PCR using the cDNA generated above, Q5 DNA High-fidelity Polymerase, and the following 487
- thermocycle conditions: 55 °C for 30 m, 94 °C for 2 m, 35 cycles of 94 °C for 15 s, 56 °C for 30 s, and 68 488
- °C for 3.5 m, 68 °C for 10 m, and held at 4 °C until use. Each PCR product was purified using Agencourt 489
- AMPure XP beads, sheared to 300 to 400 nt fragments using the Covaris S2 sonicator, indexed and 490
- prepared for sequencing as described above, and sequenced using the MiSeq V2 500 cycle kit (paired-end 491
- 492 251 nt reads). Compared to the consensus sequences generated using 35 × 400 bp amplicons on the
- 493 MiSeq, there were no consensus-level mismatches in the coding sequence using any of the other three
- 494 approaches (Extended Data Table 2). There were, however, some mismatches in the 5' and 3' UTRs
- 495 (where the genomic RNA is heavily structured), likely a result of PCR bias and decreased coverage depth.
- 496 At least 95% of the ZIKV genome was covered from samples with as low as 4 and 9 GE/µl RNA from the
- 497 amplicon and enrichment approaches, respectively. These results are similar to our previously determined
- clinical range of 10-16 ZIKV GE/µl RNA to achieve at least 95% genome coverage using our amplicon-498
- based approach<sup>24</sup>. On average, the amplicon-based sequencing approach covered 97% of the ZIKV 499
- 500 genome (>3× read-depth) and the targeted enrichment approach covered 82% of the ZIKV genome from
- 501 clinical samples (Supplementary Table 2d).

#### Phylogenetic analyses

502

527

- 503 All published and available complete ZIKV genomes of the Asian genotype from the Pacific and the
- 504 Americas were retrieved from GenBank public database as of December 2016. Public sequences (n=65)
- were codon-aligned together with ZIKV genomes generated in this study (n=39) using MAFFT<sup>54</sup> and 505
- 506 inspected manually. The multiple alignment contained 104 ZIKV sequences collected between 2013 and
- 507 2016, from the Pacific (American Samoa, French Polynesia, and Tonga), Brazil, other South and Central
- 508 Americas (Guatemala, Mexico, Suriname, and Venezuela), the Caribbean (Dominican Republic,
- 509 Guadeloupe, Haiti, Martinique, and Puerto Rico), and the United States (Supplementary File 1).
- In order to determine the temporal signal of the sequence dataset, a maximum likelihood (ML) phylogeny 510
- was first reconstructed with PhyML<sup>55</sup> using the general time-reversible (GTR) nucleotide substitution model and gamma distributed rates amongst sites<sup>56</sup> (Supplementary File 1), which was identified as the 511
- 512
- best fitting model for ML inference by iModelTest2<sup>57</sup>. Then, a correlation between root-to-tip genetic 513
- divergence and date of sampling was conducted in TempEst<sup>58</sup>. 514
- Bayesian phylogenetic analyses were performed using BEAST v.1.8.425 to infer time-structured 515
- phylogenies. We used an SDR06 nucleotide substitution model<sup>59</sup> with a non-informative continuous time 516
- Markov chain reference prior (CTMC)<sup>60</sup> on the molecular clock rate. Replicate analyses using multiple 517
- combinations of molecular clock and coalescent models were explored to select the best fitting model by 518
- marginal likelihood comparison using path-sampling and stepping-stone estimation approaches 61-63 519
- (Extended Data Table 1b). The best fit model was a relaxed molecular clock along with a Bayesian 520
- Skyline model<sup>64</sup>. All the Bayesian analyses were run for 30 million Markov chain Monte Carlo steps, 521
- sampling parameters and trees every 3000 generations (BEAST XML file and MCC tree available in 522
- 523 Supplementary File 1). Support values for all nodes are embedded in the phylogenetic tree files
- (Supplementary File 1). Tree visualizations were generated with baltic (github.com/blab/baltic). 524
- 525 The travel-associated ZIKV genomes add to the Caribbean dataset, but do not directly influence our
- 526 conclusions about the source of ZIKV introductions into Florida.

## Expected number and distribution of local cases from Zika virus importations

- We used branching process theory<sup>65,66</sup> to generate the offspring distribution (subsequent local cases) that 528
- is expected from a single introduction. The offspring distribution L is modelled with a negative binomial 529
- distribution with mean  $R_0$  and over-dispersion parameter k. The total number of cases j that is caused by a 530
- single importation (including the index case) after an infinite time<sup>67</sup> has the following form: 531

$$L = \frac{\Gamma(kj+j-1)}{\Gamma(kj)\,\Gamma(j+1)}\,\frac{(\frac{R_0}{k})^{j-1}}{(1+\frac{R_0}{k})^{kj+j-1}}$$

- 532 The parameter k represents the variation in the number of secondary cases generated by each case of
- 533 ZIKV<sup>65</sup>. In the case of vector borne diseases, local heterogeneity is high due to a variety of factors such as
- mosquito population abundance, human to mosquito interaction, and control interventions<sup>68–73</sup>. Here, we 534
- assumed high heterogeneity (k=0.1) following previous estimates for vector borne diseases<sup>66</sup>. This 535
- distribution L is plotted in Extended Data Fig. 4a. For the following, we took a forward simulation 536
- 537 approach, drawing random samples from this distribution. All estimates were based on 100,000 random
- 538 simulations.
- 539 We used this formula to estimate the probability of observing 241 local cases in Miami-Dade County
- 540 alongside 320 travel-associated cases. We approached this by sampling 320 introduction events from L
- 541 and calculating the total number of local cases in the resulting outbreak (Extended Data Fig. 4b). We also
- 542 calculated the likelihood of observing 241 local cases in the total outbreak (Extended Data Fig. 4c),
- 543 finding that the MLE of  $R_0$  lies between 0.35 and 0.55. As a sensitivity analysis, we additionally modelled
- introductions with the assumption that only 50% of travelers were infectious at time of arrival into 544
- 545 Miami-Dade County, resulting in an MLE of  $R_0$  of 0.45–0.8.
- 546 We further used this formula to address the probability of observing 3 distinct genetic clusters (F1, F2 and
- 547 F3) representing 3 introduction events in a sample of 27 ZIKV genomes from Miami-Dade County. We
- 548 approached this by sampling introduction events until we accumulated 241 local cases according to L,
- arriving at N introduction events with case counts  $(j_1, j_2, ..., j_N)$ . We then sampled 27 cases without replacement from  $(j_1, j_2, ..., j_N)$  following a hypergeometric distribution and recorded the number of 549
- 550
- distinct clusters drawn in the sample. We found that higher values of  $R_0$  resulted in fewer distinct clusters 551
- within the sample of 27 genomes (Extended Data Fig. 4d). We additionally calculated the likelihood of 552
- 553 sampling 3 distinct genetic clusters in 27 genomes (Extended Data Fig. 4e), finding an MLE estimate of
- 554  $R_0$  of 0.7–0.9. Additionally, as a sensitivity analysis we modelled a preferential sampling process in which
- larger clusters are more likely to be drawn from than smaller clusters. Here, we used a parameter  $\alpha$  that enriches the hypergeometric distribution following  $(j_1^{\alpha}, j_2^{\alpha}, \dots j_N^{\alpha})$ . In this case, we found an MLE 555
- 556
- 557 estimate of  $R_0$  of 0.5–0.9.
- 558 Using the overlap of estimates of  $R_0$  from local case counts (0.35–0.8) and genetic clusters (0.5–0.9), we
- 559 arrived at a 95% uncertainty range of R<sub>0</sub> of 0.5–0.8. As an additional sensitivity analysis, we incorporated
- 560 under-reporting in which either 50% of travel-associated cases and 25% of local cases are reported or in
- 561 which 10% of travel-associated cases and 5% of local cases are reported. We find differential reporting of
- 562 travel and local cases results in increased mean R<sub>0</sub> estimates when comparing counts of travel-associated
- to local cases (Extended Data Figure 4f-g). Additionally, we find that under-reporting increases estimates 563
- of  $R_0$  from the sampling analysis (Extended Data Figure 4h-i). Thus, moderate under-reporting is 564
- 565 consistent with  $R_0$  estimates of  $\sim 0.8$ .
- We additionally perform birth-death stochastic simulations assuming a serial interval with mean 20 566
- 567 days<sup>15</sup>. We record the number of stochastic simulations still persisting after a particular number of days
- for different values of  $R_0$  (Fig. 2c). 568

#### Zika virus incidence rates

- 570 Weekly suspected and confirmed ZIKV case counts from countries and territories within the Americas
- 571 with local transmission (January 1 to September 18, 2016) were obtained from the Pan American Health
- Organization (PAHO)<sup>30</sup>. In most cases, the weekly case numbers per country were only reported in bar 572
- 573 graphs. We contacted PAHO multiple times with the hope of gaining access to the raw data included in

- 574 the bar graphs, but our requests were unfortunately denied. Therefore we used WebPlotDigitizer v3.10
- 575 (http://arohatgi.info/WebPlotDigitizer) to estimate the numbers. We compared the actual ZIKV case
- numbers reported in Ecuador<sup>74</sup> (only country with available raw data and reported cases > 10 per week) to 576
- 577 our estimates from the PAHO bar graphs and found that the WebPlotDigitizer was ~99% accurate
- 578 (Extended Data Fig. 5a-b).

583

606

- 579 Country and territory total population sizes to calculate weekly and monthly ZIKV incidence rates were
- also obtained from PAHO<sup>75</sup>. Incidence rates calculated from countries and territories in the Americas 580
- 581 during January to June, 2016 (based on the earliest introduction time estimates until the first known cases)
- were used as an estimate for infection likelihood to investigate sources of ZIKV introductions. 582

#### Airline and cruise ship traffic

- 584 To investigate whether the transmission of ZIKV in Florida coincides with travel patterns from ZIKV
- 585 endemic regions, we obtained the number of passengers arriving at airports in Florida via commercial air
- 586 travel. We collated flight data from countries and territories in the Americas with local ZIKV
- 587 transmission between January and June, 2016 (based on the earliest introduction time estimates until the
- 588 first known cases, Supplementary Table 1b), arriving at all commercial airports in Florida. The data were
- 589 obtained from the International Air Transportation Association, which collects data on an estimated 90%
- of all passenger trips worldwide. Nelson et al.<sup>28</sup> previously reported flight data from 33 countries with 590
- ZIKV transmission entering major United States airports during October 2014 through September 2015, 591
- 592 which we used to assess the potential for ZIKV introductions outside of Florida.
- 593 Schedules for cruise ships visiting Miami, Port Canaveral, Port Everglades, Fort Lauderdale, Key West,
- 594 Jacksonville (all in Florida), Houston, Galveston (both in Texas), Charleston (South Carolina) and New
- 595 Orleans (Louisiana) ports in the year 2016 were collated from www.cruisett.com and confirmed by cross-
- 596 referencing ship logs reported by Port of Miami and reported ship schedules
- 597 Scheduled www.miamidade.gov/portmiami/. cruise ship capacities
- 598 www.cruisemapper.com. Every country/territory with ZIKV transmission visited by a cruise ship 10 days
- 599 (the approximate mean time to ZIKV clearance in human blood [i.e., the infectious period])<sup>76</sup> prior to
- arrival was counted as contributing the ship's capacity worth of passengers to Miami to the month of 600
- arrival (Supplementary Table 1b). While the air traffic was based on the reported number of travelers, we 601
- estimated the sea traffic by ship capacity. Lee and Ramdeen<sup>77</sup> reported that the average occupancy of 602
- 603
- cruise ships traveling to the Caribbean Islands exceeded 100% in 2011, and according to the Florida-
- Caribbean Cruise Association<sup>78</sup>, it remained >100% in 2015. Occupancy data for 2016 was not available 604
- at the time of publication, but we assumed that it was also near 100%. 605

#### Expected number of travelers infected with Zika virus

- 607 We estimated the expected number of travelers entering Miami who were infected with ZIKV ( $\lambda$ ) by
- using the total travel capacity (C) and the likelihood of ZIKV infection (infections (I) per person (N)) 608
- 609 from each country/territory (i):

$$\lambda = \sum_{i} C_{i} \frac{I_{i}}{N_{i}}$$

- 610 We summed the number of expected infected travelers from each country/territory with ZIKV
- 611 transmission by region and travel method (flights or cruises). The number of ZIKV cases reported by each
- 612 country are likely under-estimates in part because the majority of ZIKV infections are asymptomatic<sup>2,31</sup>.
- 613 We normalized some of the potential reporting variances between countries by reporting the data as the
- 614 relative proportion of infected travelers (Fig. 3c, Extended Data Fig. 7a) and as the absolute number of
- infected travelers (Fig. 3d, Extended Data Fig. 7b, Supplementary Table 1b) from each region. We also 615
- 616 accounted for potential reporting biases with incidence rates by using ZIKV attack rates (i.e., proportion

617 618 619 620 621 622	infected before epidemic burnout) to estimate peak transmission intensity. Attack rates were calculated using a susceptible–infected–recovered (SIR) transmission model derived from seroprevalence studies and environmental factors as described <sup>79</sup> . Using attack rates as an estimate of infection likelihood, we predict that ~60% of the infected travelers entering Miami came from the Caribbean (Extended Data 7b), which is in agreement with our methods using incidence rates of ~60-70% (Fig. 3c). A list of countries and territories used in these analyses can be found in Supplementary Table 1b.
623	Maps
624 625 626	The maps presented in our figures were generated using Matplotlib <sup>80</sup> and ESRI basemaps (www.esri.com/data/basemaps). The software and basemaps are open source and "freely available to anyone".
627	Data availability
628 629 630	All ZIKV sequencing data is available under the NCBI BioProjects PRJNA342539 and PRJNA356429. Individual sample GenBank access numbers are listed in Supplementary Table 1a. All other data is available in the Extended Data, Supplemental Information, or upon request.
631	
632	

#### **Extended References**

- 756 35. Zika virus. Florida Department of Health Available at: http://www.floridahealth.gov/diseases-and-
- 757 conditions/zika-virus/index.html?utm\_source=flhealthIndex. (Accessed: 10th January 2017)
- 758 36. Daily Zika update. Florida Department of Health Available at:
- http://www.floridahealth.gov/newsroom/all-articles.html. (Accessed: 10th January 2017)
- 760 37. Zika virus case counts in the US. The Centers for Disease Control and Prevention (2016). Available
- at: http://www.cdc.gov/zika/geo/united-states.html. (Accessed: 10th January 2017)
- 762 38. Rabe, I. B. et al. Interim Guidance for Interpretation of Zika Virus Antibody Test Results. MMWR
- 763 *Morb. Mortal. Wkly. Rep.* **65**, 543–546 (2016).
- 764 39. Waggoner, J. J. & Pinsky, B. A. Zika Virus: Diagnostics for an Emerging Pandemic Threat. J. Clin.
- 765 *Microbiol.* **54,** 860–867 (2016).
- 766 40. Lanciotti, R. S. et al. Genetic and serologic properties of Zika virus associated with an epidemic,
- 767 Yap State, Micronesia, 2007. *Emerg. Infect. Dis.* **14**, 1232–1239 (2008).
- 768 41. Interim Guidance for Zika Virus Testing of Urine United States, 2016. MMWR Morb. Mortal.
- 769 Wkly. Rep. **65**, 474 (2016).
- 42. Biggerstaff, B. PooledInfRate, version 4.0. An Excel Add-In to Compute Infection Rates from Pooled
- 771 Data. Centers for Disease Control, Fort Collins, CO (2009).
- 772 43. 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.* (2016).
- 774 doi:10.1016/S1473-3099(16)30270-5
- 775 44. Hwang, W.-H. & He, F. Estimating abundance from presence/absence maps. *Methods Ecol. Evol.* 2,
- 776 550–559 (2011).
- 45. Corman, V. M. et al. Clinical comparison, standardization and optimization of Zika virus molecular
- detection. Bull. World Health Organ. (2016).
- 46. Worobey, M. et al. 1970s and 'Patient 0' HIV-1 genomes illuminate early HIV/AIDS history in
- 780 North America. *Nature* (2016). doi:10.1038/nature19827

- 781 47. Bolger, A. M., Lohse, M. & Usadel, B. Trimmomatic: a flexible trimmer for Illumina sequence data.
- 782 *Bioinformatics* **30,** 2114–2120 (2014).
- 48. Li, H. et al. The Sequence Alignment/Map format and SAMtools. Bioinformatics 25, 2078–2079
- 784 (2009).
- 785 49. Köster, J. & Rahmann, S. Building and documenting workflows with python-based snakemake. in
- 786 OASIcs-OpenAccess Series in Informatics 26, (Schloss Dagstuhl-Leibniz-Zentrum fuer Informatik,
- 787 2012).
- 788 50. Kearse, M. et al. Geneious Basic: an integrated and extendable desktop software platform for the
- organization and analysis of sequence data. *Bioinformatics* **28**, 1647–1649 (2012).
- 790 51. Martin, M. Cutadapt removes adapter sequences from high-throughput sequencing reads.
- 791 *EMBnet.journal* **17**, 10–12 (2011).
- 792 52. Schmieder, R. & Edwards, R. Quality control and preprocessing of metagenomic datasets.
- 793 *Bioinformatics* **27**, 863–864 (2011).
- 794 53. Langmead, B. & Salzberg, S. L. Fast gapped-read alignment with Bowtie 2. Nat. Methods 9, 357–
- 795 359 (2012).
- 796 54. Katoh, K. & Standley, D. M. MAFFT multiple sequence alignment software version 7:
- improvements in performance and usability. *Mol. Biol. Evol.* **30**, 772–780 (2013).
- 798 55. Guindon, S. & Gascuel, O. A simple, fast, and accurate algorithm to estimate large phylogenies by
- 799 maximum likelihood. *Syst. Biol.* **52**, 696–704 (2003).
- 800 56. Yang, Z. Maximum likelihood phylogenetic estimation from DNA sequences with variable rates
- 801 over sites: approximate methods. *J. Mol. Evol.* **39**, 306–314 (1994).
- 802 57. Darriba, D., Taboada, G. L., Doallo, R. & Posada, D. ModelTest 2: more models, new heuristics
- and parallel computing. *Nat. Methods* **9,** 772 (2012).
- 804 58. Rambaut, A., Lam, T. T., Max Carvalho, L. & Pybus, O. G. Exploring the temporal structure of
- heterochronous sequences using TempEst (formerly Path-O-Gen). Virus Evol 2, vew007 (2016).
- 806 59. Shapiro, B., Rambaut, A. & Drummond, A. J. Choosing appropriate substitution models for the

- phylogenetic analysis of protein-coding sequences. *Mol. Biol. Evol.* **23**, 7–9 (2006).
- 808 60. Ferreira, M. A. R. & Suchard, M. A. Bayesian analysis of elapsed times in continuous-time Markov
- chains. The Canadian Journal of Statistics / La Revue Canadienne de Statistique 36, 355–368
- 810 (2008).
- 811 61. Baele, G. et al. Improving the accuracy of demographic and molecular clock model comparison
- while accommodating phylogenetic uncertainty. *Mol. Biol. Evol.* **29**, 2157–2167 (2012).
- 813 62. Xie, W., Lewis, P. O., Fan, Y., Kuo, L. & Chen, M.-H. Improving marginal likelihood estimation for
- Bayesian phylogenetic model selection. Syst. Biol. **60**, 150–160 (2011).
- 815 63. Gelman, A. & Meng, X.-L. Simulating Normalizing Constants: From Importance Sampling to
- 816 Bridge Sampling to Path Sampling. *Stat. Sci.* **13**, 163–185 (1998).
- 817 64. Drummond, A. J., Rambaut, A., Shapiro, B. & Pybus, O. G. Bayesian coalescent inference of past
- population dynamics from molecular sequences. *Mol. Biol. Evol.* **22**, 1185–1192 (2005).
- 819 65. Churcher, T. S. et al. Public health. Measuring the path toward malaria elimination. Science 344,
- 820 1230–1232 (2014).
- 821 66. Lloyd-Smith, J. O., Schreiber, S. J., Kopp, P. E. & Getz, W. M. Superspreading and the effect of
- individual variation on disease emergence. *Nature* **438**, 355–359 (2005).
- 823 67. Nishiura, H., Yan, P., Sleeman, C. K. & Mode, C. J. Estimating the transmission potential of
- 824 supercritical processes based on the final size distribution of minor outbreaks. J. Theor. Biol. 294,
- 825 48–55 (2012).
- 826 68. Perkins, T. A., Scott, T. W., Le Menach, A. & Smith, D. L. Heterogeneity, mixing, and the spatial
- 827 scales of mosquito-borne pathogen transmission. *PLoS Comput. Biol.* **9**, e1003327 (2013).
- 828 69. Kraemer, M. U. G. et al. Big city, small world: density, contact rates, and transmission of dengue
- 829 across Pakistan. J. R. Soc. Interface 12, 20150468 (2015).
- 830 70. Kraemer, M. U. G. et al. Spread of yellow fever virus outbreak in Angola and the Democratic
- Republic of the Congo 2015–16: a modelling study. *Lancet Infect. Dis.* **17**, 330–338 (2017/3).
- 71. Struchiner, C. J., Rocklöv, J., Wilder-Smith, A. & Massad, E. Increasing Dengue Incidence in

- Singapore over the Past 40 Years: Population Growth, Climate and Mobility. *PLoS One* **10**,
- 834 e0136286 (2015).
- 72. Fauver, J. R. et al. Temporal and Spatial Variability of Entomological Risk Indices for West Nile
- 836 Virus Infection in Northern Colorado: 2006-2013. J. Med. Entomol. **53**, 425–434 (2016).
- 837 73. Rocklöv, J. et al. Assessing Seasonal Risks for the Introduction and Mosquito-borne Spread of Zika
- 838 Virus in Europe. *EBioMedicine* **9,** 250–256 (2016).
- 74. Ministry of Public Health, Ecuador. National Direction of Epidemiological Surveillance for Vector
- Transmitted Diseases (Spanish) Available at: http://www.salud.gob.ec/wp-
- content/uploads/2015/12/GACETA-ZIKA SE5corregido.pdf. (Accessed: 19th March 2017)
- 842 75. Zika Cumulative Cases. Pan American Health Organization Available at:
- http://www.paho.org/hq/index.php?option=com\_content&view=article&id=12390&Itemid=42090&I
- ang=en. (Accessed: 1st December 2016)
- 845 76. Lessler, J. T. et al. Times to key events in the course of Zika infection and their implications: a
- systematic review and pooled analysis. *Bull. World Health Organ.* (2016).
- 77. Lee, S. & Ramdeen, C. Cruise ship itineraries and occupancy rates. *Tourism Manage*. **34,** 236–237
- 848 (2013/2).
- 78. FCCA Research & Statistics. Florida-Caribbean Cruise Association Available at: http://www.f-
- cca.com/research.html. (Accessed: 1st March 2017)
- 79. Perkins, T. A., Siraj, A. S., Ruktanonchai, C. W., Kraemer, M. U. G. & Tatem, A. J. Model-based
- projections of Zika virus infections in childbearing women in the Americas. *Nat Microbiol* **1,** 16126
- 853 (2016).
- 854 80. Hunter, J. D. Matplotlib: A 2D Graphics Environment. Comput. Sci. Eng. 9, 90–95 (2007).

#### Extended Data

Extended Data Fig. 1 | Miami-Dade mosquito surveillance and relative Aedes aegypti abundance. (a) Mosquito surveillance data reported from June 21 to November 28, 2016 was used to evaluate the risk of ZIKV infection from mosquito-borne transmission in Miami. A total of 24,306 Ae. aegypti and 45 Ae. albopictus were collected. Trap nights are the total number of times each trap site was used and the trap locations are shown in Fig. 1d (some "Other Miami" trap sites are located outside of mapped region). Up to 50 mosquitoes of the same species and trap night were pooled together for ZIKV RNA testing. The infection rates were calculated using a maximum likelihood estimate (MLE). None of the Ae. albopictus pools contained ZIKV RNA. (b) The number of weekly ZIKV cases (based on symptoms onset) was correlated with mean Ae. aegypti abundance per trap night determined from the same week and zone (Spearman r = 0.61). This suggests that when the virus is present, mosquito abundance numbers alone could be used to target control efforts. (c) Insecticide usage, including truck and aerial adulticides and larvacides, by the Miami-Dade Mosquito Control in Wynwood (left) and Miami Beach (right) was overlaid with Ae. aegypti abundance per trap night to demonstrate that intense usage of insecticides may have helped to reduce local mosquito populations. (d) Relative Ae. aegypti abundance for each Florida county and month was estimated using a multivariate regression model, demonstrating spatial and temporal heterogeneity for the risk of ZIKV infection.

Extended Data Fig. 2 | Maximum likelihood tree and root-to-tip regression of Zika virus genomes from Pacific islands and the epidemic in Americas. (a) Maximum likelihood tree of publicly available ZIKV sequences and sequences generated in this study (n=104). tips are coloured by location, labels in bold indicate sequences generated in this study, Florida clusters F1-F4 are indicated by vertical lines to the right of the tree. Bootstrap support values are shown at key nodes. All other support values can be found in Supplementary File 1. (b) Linear regression of sample tip dates against divergence from root based on sequences with known collection dates estimates an evolutionary rate for the ZIKV phylogeny of 1.10×10<sup>-3</sup> nucleotide substitutions/site/year (subs/site/yr). This is consistent with BEAST analyses using a relaxed molecular clock and a Bayesian Skyline tree prior, the best-performing combination of clock and demographic model according to marginal likelihood estimates (Extended Data Table 1c), which estimated an evolutionary rate of 1.21×10<sup>-3</sup> (95% highest posterior density: 1.01 - 1.43×10<sup>-3</sup>) subs/site/yr (Extended Data Table 1a). These values are in agreement with previous estimates calculated based on ZIKV genomes from Brazil<sup>6</sup>.

**Extended Data Fig. 3 I Molecular clock dating of Zika virus clades.** Maximum clade credibility (MCC) tree of ZIKV genomes collected from Pacific islands and the epidemic in Americas (n=104). Circles at the tips are colored based on origin location. Clade posterior probabilities are indicated by white circles filled with black relative to the support. A posterior probability of 1 fills the entire circle black. The grey violin plot indicates the 95% highest posterior density (HPD) interval for the tMRCA of the American epidemic. We estimated that the tMRCA for the ongoing epidemic in the Americas occurred during October, 2013 (node AM, Extended Table 1, 95% HPD: August, 2013-January, 2014), which is consistent with previous analysis based on ZIKV genomes from Brazil<sup>6</sup>.

 Extended Data Fig. 4 l Estimation of basic reproductive number and number of introductions in Miami-Dade County. (a) Probability distribution of estimated total number of cases caused by a single introduction (excluding the index case) for different values of  $R_0$ . (b) Mean and 95% CI for total number of local cases caused by 320 introduction events (*i.e.*, travel-associated cases diagnosed in Miami-Dade County) for different values of  $R_0$  and for different assumptions of proportion of infectious travelers. (c) Log likelihood of observing 241 local cases in Miami-Dade County with 320 introduction events for different values of  $R_0$  along with 95% maximum likelihood estimate (MLE) bounds on  $R_0$ . (d) Mean and 95% uncertainty interval for total number of distinct phylogenetic clusters observed in 27 sequenced ZIKV genomes from human cases diagnosed in Miami-Dade County for different values of  $R_0$  and for

different assumptions of sampling bias, from  $\alpha$ =1 (no sampling bias) to  $\alpha$ =2 (skewed toward preferentially sampling larger clusters). (e) Log likelihood of observing 3 clusters (*i.e.*, ZIKV lineages F1, F2, and F4, Fig. 2a) in 27 sequenced cases for different values of  $R_0$  along with 95% MLE bounds on  $R_0$ . (f) Mean and 95% CI for total number of local cases caused by 320 observed travel-associated cases with travel-associated vs local reporting rates of 50%/25% and 10%/5%. This assumes 50% of travelers are infectious. (g) Log likelihood of observing 241 local cases with 320 introduction events for different values of  $R_0$  along with 95% MLE bounds on  $R_0$  with travel-associated vs local reporting rates of 50%/25% and 10%/5%. (h) Mean and 95% uncertainty interval for total number of distinct phylogenetic clusters observed in 27 sequenced ZIKV genomes for different values of  $R_0$  and for assumptions of local reporting rate of 5% and 25%. This assumes preferential sampling ( $\alpha$ =2). (i) Log likelihood of observing 3 clusters in 27 sequenced cases for different values of  $R_0$  along with 95% MLE bounds on  $R_0$  with local reporting rate of 5% and 25%. At 5% local reporting rate, 0 of the 100,000 replicates for all  $R_0$  values showed 3 clusters.

Extended Data Fig. 5 I Weekly reported Zika virus case numbers and incidence rates in the Americas. (a) Most ZIKV case numbers reported by PAHO<sup>30</sup> were only available as bar graphs (raw data was not made available to us at the time of request). Therefore we used the WebPlotDigitizer to estimate the weekly case numbers from the PAHO bar graphs. ZIKV cases reported from Ecuador was the only data set to include a link to the actual case numbers that also had >10 cases per week<sup>74</sup>. To validate the WebPlotDigitizer, we compared the weekly reported case numbers from Ecuador to our estimates. (b) The reported and estimated case numbers were strongly correlated (Spearman r = 0.9981). The WebPlotDigitizer was used to estimate the ZIKV case numbers for all subsequent analysis. (c) ZIKV cases (suspected and confirmed) and (d) incidence rates (normalized per 100,000 population) are shown for each country or territory with available data per epidemiological week from January 1 to September 18, 2016. (e) Each country or territory with available data is colored by its reported ZIKV incidence rate from January to June, 2016 (the time frame for analysis of ZIKV introductions into Florida).

**Extended Data Fig. 6 | Cruise and flight traffic entering Miami from regions with Zika virus transmission.** The estimated number of passengers entering Miami, by either (a) cruises or (b) flights, from each country or territory in the Americas with ZIKV transmission per month (left panel). The center map and inset show the cumulative numbers of travelers entering Miami during January to June, 2016 (the time frame for analysis of ZIKV introductions into Florida) from each country or territory per method of travel. (c) The total traffic (*i.e.* cruises and flights) is shown entering Miami per month.

Extended Data Fig. 7 l Expected number of Zika virus infected travelers from the Caribbean is correlated with the total observed number of travel-associated infections. (a) In order to account for potential biases in ZIKV reporting accuracies, we also estimated the proportion of infected travelers using projected ZIKV attack rates<sup>79</sup> (*i.e.* predicted proportion of population infected before epidemic burnout). About 60% of the infected travelers are expected to have arrived from the Caribbean, similar to our results using incidence rates (Fig. 3c). (b) The expected number of travel-associated ZIKV cases were estimated by the number of travelers coming into Miami from each country/territory (travel capacity) and the in-country/territory infection likelihood (incidence rate per person) per week. The expected travel cases were summed from all of the Americas (left), Caribbean (left center), South America (right center), and Central America (right) and plotted with the observed travel-associated ZIKV cases. Numbers in each plot indicate Spearman correlation coefficients. Negative Spearman r coefficients indicated a negative correlation between the number of expected and observed travel cases.

**Extended Data Fig. 8 I Greater early season potential for Zika virus introductions into Miami.** The monthly cruise ship and airline<sup>28</sup> capacity from countries/territories with ZIKV transmission for the major United States travel hubs (shown as circle diameter) with monthly potential *Ae. aegypti* abundance (circle color), as previously estimated<sup>22</sup>. The abundance ranges were chosen with respect to the May-Oct Miami

735	mean: "None to low" (<2%), "Low to moderate" (2-25%), "Moderate to high (25-75%), and "High"
736	(>75%). Mosquito-borne transmission is unlikely in the "None to low" range. Cruise capacities from
737	Houston and Galveston, Texas were combined.
738	
739	Extended Data Table 1   (a) Time of the most recent common ancestor and evolutionary rate and
740	(b) Model selection to infer time-structured phylogenies.
741	HPD, highest posterior density. Dates listed as proportion of days elapsed with a year. Clades refer to Fig.
742	2a.
743	<del></del>
744	Extended Data Table 2   Validation of sequencing results.
745	<sup>a</sup> Compared to the consensus genomes generated by sequencing 35 × 400 bp amplicons on the MiSeq.
746	b Amplicons produced using Ion AmpliSeq and 875 custom ZIKV primers.
747	NGS, next-generation sequencing; UTR, untranslated region; CDS, coding sequence.
	1700, new generation sequencing, o 11t, and answer region, 020, coding sequence.
748	
749	
750	
751	
731	
752	
753	
754	

а	Miami-Dade mosquito surveillence					Ae. aegypti	No. per	Pools	ZIKV+	MLE
	Location	Start date	End date	Trap sites	Trap nights	collected	trap night	tested	pools	infection rate
	Wynwood	Jul 23	Nov 25	29	824	4,972	6.0	738	0	0
	Miami Beach	Aug 21	Nov 28	29	2,180	13,265	6.1	1,473	8	0.61 (0.28-1.15)
	Little River	Oct 15	Nov 28	5	217	485	2.2	136	0	0
	Other Miami	Jun 21	Sep 23	52	235	5,584	23.8	249	0	0

