Distinct Zika Virus Lineage in Salvador, Bahia, Brazil

Zika virus is an arthropodborne RNA virus primarily transmitted by mosquitoes of the species Aedes (1). The virus has 2 genotypes: African, found only in the continent of Africa; and Asian, associated with outbreaks in Southeast Asia, several Pacific islands, and, recently, the Americas (2). In May 2015, Brazil reported its first autochthonous cases of Zika virus infection, which occurred in northeast Brazil (3,4). As of June 30, 2016, all 27 federal states in Brazil had confirmed Zika virus transmission (http://www.paho.org/hq/index.php?option=com_docman&task=doc_view&Itemid=270&gid=35262&lang=en). The rapid geographic expansion of Zika virus transmission and the virus’s association with microcephaly and congenital abnormalities (5) demand a rapid increase in molecular surveillance in areas that are most affected. Molecular surveillance is particularly relevant for regions where other mosquito-borne viruses, particularly dengue and chikungunya viruses, co-circulate with Zika virus (2); surveillance on the basis of clinical symptoms alone is highly inaccurate. Genetic characterization of circulating Zika virus strains can help determine the origin and potential spread of infection in travelers returning from Zika virus–endemic countries. Previous analyses have suggested that Zika virus was introduced in the Americas at least 1 year before the virus’s initial detection in Brazil (1). The state of Bahia, Brazil, reported most (93%) suspected Zika virus infections in Brazil during 2015 (2), including cases of Zika virus–associated fetal microcephaly (6); however, except for 1 complete genome, no genetic information from the region has been available (2,7). We report molecular epidemiologic findings resulting from 11 new complete and partial Zika virus genomes recovered from serum samples from patients at the Hospital Aliança in the city of Salvador in Bahia, Brazil.

The Study
Symptomatic patients with suspected Zika virus infection were enrolled in a research study approved by the Brazilian Ministry of Health (Certificado de Apresentação para Apreciação Eética 45483115.0.0000.0046, no. 1159.184, Brazil). During April 2015–January 2016, acute Zika virus infection was diagnosed for 15 patients whose serum samples tested positive by a qualitative reverse transcription PCR (RT-PCR) by using primers targeting the nongenomic 5 gene (8). Clinical samples were retested for Zika virus positivity by using a separate quantitative RT-PCR (QuantiTect SYBR Green PCR kit; QIAGEN, Valencia, CA, USA) and primers targeting the envelope gene (9). Metagenomic next-generation sequencing libraries were constructed from serum RNA extracts, as described (10,11; online Technical Appendix, http://wwwnc.cdc.gov/EID/article/22/10/16-0663-Techapp1.pdf). Pathogen identification from metagenomic next-generation sequencing data was performed by using the Sequence-based Ultra-Rapid Pathogen Identification bioinformatics pipeline (12; http://chiulab.ucsf.edu/surpi/). Results of the metagenomic analyses and identification of co-infections with chikungunya virus are reported elsewhere (13).

For Zika virus genome sequencing, 2 isolates (Bahia07 and Bahia09; Table) with Zika virus titers >10¹⁰ copies/mL generated sufficient viral metagenomic data for complete genome assembly. For the remaining samples with lower titers, metagenomic next-generation sequencing libraries were enriched for Zika virus sequencing by using xGen biotinylated lockdown capture probes (Integrated DNA Technologies, San Carlos, CA, USA) and primers targeting the envelope gene (9). Metagenomic next-generation sequencing libraries were constructed from serum RNA extracts, as described (10,11; online Technical Appendix, http://wwwnc.cdc.gov/EID/article/22/10/16-0663-Techapp1.pdf). Pathogen identification from metagenomic next-generation sequencing data was performed by using the Sequence-based Ultra-Rapid Pathogen Identification bioinformatics pipeline (12; http://chiulab.ucsf.edu/surpi/). Results of the metagenomic analyses and identification of co-infections with chikungunya virus are reported elsewhere (13).

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Genbank accession no. | Zika virus RT-PCR C<sub>r</sub> | Viral load, copies/mL | 160-nt single-end metagenomic reads genome recovery, %‡ | Mean fold coverage | 250-nt paired-end Zika virus-specific enrichment genome recovery, %‡ | Mean fold coverage
---|---|---|---|---|---|---
Bahia01 | KX101066 | 34.6 | 1,042 | 23.1 | 0.4 | 65.3 | 16,288.2
Bahia02 | KX101060 | 32.5 | 4,086 | 26.0 | 0.4 | 73.4 | 20,045.8
Bahia03 | KX101061 | 32.8 | 3,272 | 1.1 | 0.0 | 77.7 | 220.0
Bahia04 | KX101062 | 34.1 | 1,461 | 5.1 | 0.1 | 42.0 | 4,659.5
Bahia05 | KX101063 | 33.7 | 1,901 | 5.0 | 0.1 | 42.8 | 8,547.5
Bahia07 | KU940228 | 13.7 | 9.1 × 10<sup>8</sup> | 100 | 3,603.5 | ND | ND
Bahia08 | KU940227 | 33.3 | 2,470 | 75.1 | 9.2 | 84.9 | 23,805.1
Bahia09 | KU940224 | 29.9 | 23,121 | 99.98 | 41.5 | ND | ND
Bahia11 | KX101064 | Neg (no C<sub>r</sub>) | NA | 27.8 | 0.9 | 64.0 | 28,704.1
Bahia12 | KX101067 | 34.2 | 1,327 | 11.2 | 0.2 | 50.4 | 10,461.8
Bahia15 | KX101065 | Neg (no C<sub>r</sub>) | NA | 4.6 | 0.2 | 45.4 | 3,706.8

*C<sub>r</sub>, cycle threshold; NA, not applicable; ND, not done; Neg, negative; Pos, positive; qRT-PCR, quantitative reverse transcription PCR; RT-PCR, reverse transcription PCR; U, unknown.
†Samples were collected from Salvador in Bahia, Brazil, except for Bahia05, which was collected in Camaçari, Bahia, Brazil.
‡Assumes a genome size of 10,676 nt, the size of the prototype Brazilian Zika virus strain SPH2015 (KU321639).

Technologies, Redwood, CA, USA) designed to tile across all sequenced Zika virus genomes >10,000 nt in GenBank (http://www.ncbi.nlm.nih.gov/genbank) as of March 1, 2016. Capture probes were curated for redundancy at a 99% nt similarity cutoff. Enrichment was performed on the metagenomic libraries in pools of 8 libraries (including Zika virus–negative serum sample controls) by using the xGen lockdown probe protocol and the SeqCap EZ Hybridization and Wash Kit (Roche, Indianapolis, IN, USA). Eleven Zika virus genomes with >40% genome recovery (mean 69.4% ± 2.0%) were assembled (Table). Distribution of single nucleotide variants across the 11 recovered genomes exhibited distinct patterns (online Technical Appendix Figure 1), indicating that the assembled genomes were unlikely to result from cross-contamination by a single high-titer Zika virus sample.

Multiple sequence alignment was performed by using MAFFT version 7 (http://mafft.cbrc.jp/alignment/software/); maximum-likelihood (ML) and Bayesian phylogenetic inferences were determined by using PhyML version 3.0 (http://www.atgc-montpellier.fr/phyml/) and BEAST version 1.8.2 (http://beast.bio.ed.ac.uk/), respectively. The best-fit model was calculated by using jModelTest2 (https://github.com/ddarriba/jmodeltest2; details in online Technical Appendix). Coding regions corresponding to the 11 complete or partial genomes from Bahia were aligned with all published and available near-complete Zika virus genomes and longer subgenomic regions (>1,500 nt) of the Asian genotype as of April 2016 (mean sequence size 8,402 nt with 1,652 distinct nucleotide site patterns).

The isolates from patients in Salvador clustered together within 1 strongly supported clade (posterior probability 1.00, bootstrap support 100%, Bahia clade C) (Figure; online Technical Appendix Figure 2). This support is notable; most Zika virus genomes in this clade are incomplete, and uncertainty is accounted for in phylogenetic inference. The tree topology accords with previous findings (2,4,5), and time to most recent common ancestor (TMRCA) of the epidemic in the Americas is similar to that previously estimated (2) (American epidemic clade A; Figure). The overall ML and molecular clock phylogenies exhibited many well-supported internal nodes with bootstrap support >60% and posterior probability >0.80 (Figure; online Technical Appendix Figure 2), although several nodes near the ancestor of clade A were less well supported.
**Figure.** Timeframe of Zika virus outbreaks in the Americas. A molecular clock phylogeny is shown with the Zika virus outbreak lineage estimated from complete and partial (>1,500 nt) coding region sequences. For visual clarity, 5 basal Southeast Asia sequences (GenBank accession nos. HQ23499 [Malaysia, 1966]; EU545988 [Micronesia, 2007]; KU681082 [Philippines, 2012]; JN860885 [Cambodia, 2010]; and KU681081 [Thailand, 2013]) are not displayed. Blue horizontal bars represent 95% Bayesian credible intervals for divergence dates. A, B, and C denote the current American epidemic, the northeastern Brazil (Maranhão sequence and Bahia), and the Bahia clades, respectively; numbers next to the clade denote posterior probabilities and bootstrap scores in percentages. Circle sizes at each node represent the posterior probability support of that node. Taxa are labeled with the GenBank accession numbers, sampling location, and sampling date. Names of sequences generated in this study are in bold. The inset graph on the left shows the posterior probability distributions of the estimated ages (time to most recent common ancestor) for clades A, B, and C. The posterior probability density is plotted on the vertical axis as a function of time on the horizontal axis (tick marks designate 3-month intervals). Estimated ages were determined with BEAST version 1.8.2 (http://beast.bio.ed.ac.uk) by using the best-fitting evolutionary model. The posterior probability distributions were visualized by using Tracer version 1.6 (http://tree.bio.ed.ac.uk/software/tracer/). Brazil states: BA, Bahia; CE, Ceará; MA, Maranhão; PA, Pará; PB, Paraíba; RN, Rio Grande do Norte; RJ, Rio de Janeiro; SP, São Paulo.
The updated phylogenetic analyses, including the newly identified clade C, suggest that Zika virus was introduced in Bahia during March–September 2014. An isolate from Maranhão in northeastern Brazil (≈1,000 km from Bahia) is ancestral to the Bahia clade (posterior probability 1.00, bootstrap support 74%, northeastern Brazil clade B) (Figure; online Technical Appendix Figure 2). The TMRCA of clade B (comprising the Bahia clade and the Maranhão sequence) is estimated to be September 2013–April 2014, an early stage of the epidemic. This TMRCA is consistent with the hypothesis that Zika virus in the Americas originated in Brazil (2). A previously reported sequence from Bahia (6) clustered with an isolate from Belém in the state of Pará in northern Brazil, ≈3,000 km from Bahia (posterior probability 0.99, bootstrap support 81%) (Figure; online Technical Appendix Figure 2). The patient denied history of travel, suggesting that multiple Zika virus lineages may circulate in Bahia.

Conclusions
Our results suggest an early introduction and presence (mid-2014) of Zika virus in the Salvador region in Bahia, Brazil. Given the size of the cluster and statistical support for it, this lineage likely represents a large and sustained chain of transmission within Bahia state. Most cases of this Zika virus lineage clustered closely to a sequence from Maranhão, and we found evidence for an additional potential introduction to Bahia from Pará state. Consequently, Zika virus in Salvador during mid-2014 was likely introduced from other regions in Brazil rather than from outside the country. Current findings of Zika virus emergence in Bahia state during mid-2014 are consistent with first-trimester viral infection in pregnant women corresponding to the initial reported cases of fetal microcephaly, which began in January 2015 (5) and peaked in November 2015.

Broader sampling across Bahia is needed to determine whether the Salvador lineage (clade C) identified in this article comprises most Zika virus cases in the state. Brazil currently faces a major public health challenge from co-circulation of Zika, dengue, and chikungunya viruses (2–4,14,15). Additional molecular surveillance in the Americas and beyond is urgently needed to trace and predict transmission of Zika virus.

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**EMERGING INFECTIOUS DISEASES**

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Distinct Zika Virus Lineage in Salvador, Bahia, Brazil

Technical Appendix

Next-Generation Sequencing Library Construction

RNA metagenomic libraries were constructed from the patients’ serum sample as previously described (1,2). Total nucleic acid was extracted by using the QIAamp Viral RNA mini kit (Qiagen, Valencia, CA, USA). Extract (20 μL) was treated with Turbo DNase (Ambion, Waltham MA) and Baseline-ZERO DNase (Epicentre, Madison, WI, USA), followed by cleanup with the RNA Clean & Concentrator kit (Zymo Research, Irvine, CA, USA). Eluted RNA was reverse transcribed to cDNA by using Superscript III Reverse Transcriptase (Invitrogen, Carlsbad, CA, USA) mediated by random hexamers. Second strand synthesis was conducted by using Sequenase version 2.0 DNA Polymerase (Affymetrix, Santa Clara, CA, USA) for 10 minutes at 37°C following denaturation. Double-stranded cDNA was cleaned up by using the DNA Clean & Concentrator kit (Zymo Research, Irvine, CA, USA) and next-generation sequencing (NGS) libraries were generated from the entirety of the eluate by using the Nextera XT DNA Library Prep Kit (Illumina, San Diego, CA, USA). The Nextera XT library was cleaned by using Agencort AMPure XP beads (Beckman-Coulter, Brea, CA, USA). Dual-indexed, barcoded NGS libraries were quantitated on the Qubit 3.0 fluorometer (Fisher Scientific, Waltham, MA, USA), mixed equally by concentration, and quantified for size and concentration on the Bioanalyzer (Agilent, Santa Clara CA, USA) using the Agilent High Sensitivity dsDNA kit. Libraries were run on the Illumina HiSeq sequencing system (1 × 160 nt single-end runs and 2 × 250 nt paired-end runs). Metagenomic NGS data was analyzed for pathogens by using the automated Sequence-Based Ultra-Rapid Pathogen Identification (SURPI) computational pipeline (http://chiulab.ucsf.edu/surpi/) and the March 2015 National Center for Biotechnology Information nucleotide database (Bethesda, MD, USA) (3).
**Phylogenetic Analyses**

By using the MAFFT program (4), multiple sequence alignment of the coding regions corresponding to the 11 complete or partial genomes from Bahia, Brazil, which were recovered in this study were aligned together with all published and available near-complex Zika virus genomes and longer subgenomic regions (>1,500 nt) of the Asian genotype submitted to GenBank as of April 2016. Maximum likelihood and Bayesian phylogenetic inferences were determined by using PhyML version 3.0 (5; http://www.atgc-montpellier.fr/phylml/) and BEAST version 1.8.2 (6; http://beast.bio.ed.ac.uk/), respectively. We used the program jModelTest2 (7) to determine the best-fitted general time reversible nucleotide substitution model with a proportion of invariant sites (GTR+I).

**Data Availability**

The 11 assembled complete and partial Zika virus genomes recovered in this study have been deposited in GenBank (accession nos. KU940224, KU940227–KU940228, and KX101060–KX101067). Raw FASTQ sequence files with human reads removed by using Bowtie2 (8; http://bowtie-bio.sourceforge.net/bowtie2/index.shtml) in local alignment mode at default parameters have been deposited in the Sequence Read Archive (National Center for Biotechnology Information; accession no. SRP072069).

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Technical Appendix Figure 1. Multiple sequence alignment of 11 whole and partial Zika virus genomes from the Salvador region of Bahia, Brazil. Viral genomes were aligned by using the MUSCLE program (9; http://www.drive5.com/muscle/) at default parameters. Aligned genomic regions are shown in light tan, with missing genomic regions from partially recovered genomes shaded in dark tan. Single nucleotide variants differing from the fully sequenced Bahia09 strain (in red) are plotted as vertical black lines according to nucleotide position along the viral genome (horizontal axis). The single nucleotide variant patterns differ for each strain; none are identical to Bahia07 or Bahia09, the Zika virus isolates with the highest viral titers. NS, nonstructural gene; UTR, untranslated region.
Technical Appendix Figure 2. Maximum likelihood phylogeny of Zika virus Asian genotype. The tree was estimated from complete and partial (>1,500 nt) coding region sequences. Taxa are labeled with GenBank accession numbers (http://www.ncbi.nlm.nih.gov/genbank/), sampling locations, and sampling dates. Names of sequences generated in this study are shown in bold. Branch lengths are drawn proportionally to the number of nucleotide substitutions per position (indicated by the scale bar). Numbers next to phylogenetic nodes denote bootstrap percentages (1,000 replicates). Brazil states: BA, Bahia; CE, Ceará; MA, Maranhão; PA, Pará; PB, Paraíba; SP, São Paulo; RN, Rio Grande do Norte; RJ, Rio de Janeiro.