Chikungunya virus (CHIKV), dengue virus (DENV), and Zika virus (ZIKV) are 3 of the most common arthropod-borne viruses (arboviruses) that infect humans. All are transmitted by the anthropophilic and urban-adapted Aedes aegypti and Ae. albopictus mosquito vectors (1). Driven by human movement and climate trends, the distribution of these mosquitoes is expanding along with the arboviruses they transmit (2). In Latin America, CHIKV and ZIKV have emerged since the mid-2000s, joining DENV, which is already endemic there (3). In this region, only Uruguay and Chile did not report autochthonous transmissions of one of these arboviruses during 2014–2019, highlighting the current state of endemicity (4).

Paraguay has been severely affected by emergent Zika and chikungunya viruses, and dengue virus is endemic. To learn more about the origins of genetic diversity and epidemiologic history of these viruses in Paraguay, we deployed portable sequencing technologies to strengthen genomic surveillance and determine the evolutionary and epidemiologic history of arthropod-borne viruses (arboviruses). Samples stored at the Paraguay National Central Laboratory were sequenced and subjected to phylogenetic analysis. Among 33 virus genomes generated, we identified 2 genotypes of chikungunya and 2 serotypes of dengue virus that circulated in Paraguay during 2014–2018; the main source of these virus lineages was estimated to be Brazil. The evolutionary history inferred by our analyses precisely matched the available travel history of the patients. The genomic surveillance approach used was valuable for describing the epidemiologic history of arboviruses and can be used to determine the origins and evolution of future arbovirus outbreaks.

Chikungunya virus (CHIKV), dengue virus (DENV), and Zika virus (ZIKV) are 3 of the most common arthropod-borne viruses (arboviruses)
of chikungunya fever in Paraguay were first reported in June 2014; autochthonous cases were first detected in 2015, and CHIKV caused seasonal outbreaks every year until 2018. Zika was first detected in November 2015, and autochthonous infections were confirmed soon after (9). To date, however, little is known about the genetic diversity of CHIKV and ZIKV that circulate in Paraguay.

The potential triple epidemic scenario (i.e., CHIKV, DENV, ZIKV) in Paraguay could pose serious public health and economic burdens. Arbovirus surveillance is critical for assisting health services with preparedness, providing key information about the seasonality of infections and diversity of circulating viral lineages. When resources allow, such surveillance can now involve genomic surveillance via portable sequencing technologies. For example, this approach was successfully used to study the ZIKV epidemic in the Americas (10,11), the reemergence of yellow fever virus in Brazil (12,13), and recurrent outbreaks of CHIKV in several regions of Brazil (14–16).

To help reinforce arbovirus surveillance in Paraguay, we performed portable genome sequencing under the scope of the ZIBRA project (http://zibra-project.org) at the Laboratorio Central de Salud Pública in Asunción, Paraguay. During July 16–20, 2018, a team of molecular biologists from Brazil and Paraguay worked on a group of samples selected to determine the recent history of arboviruses in the country, generating 33 viral genomes and building capacity skills among the local laboratory staff. We report the analysis of the origins and spread of CHIKV in Paraguay as well as the current dynamics of DENV. The project was reviewed and approved by the Comisión Nacional de Ética en Investigación (CONEP) from the Brazilian Ministry of Health as part of the arbovirus genomic surveillance efforts within the terms of CONEP Resolution 510/2016 by the Pan American Health Organization Ethics Review Committee (PAHO-2016-08-0029) and by the Paraguayan Ministry of Public Health and Social Welfare (MSPyBS/S.G. no. 0944/18).

Methods

Sample and Data Collection

This study was necessarily based on convenience sampling; de-identified samples were obtained from material exceeding the routine number of arbovirus diagnoses and stored at the Laboratorio Central de Salud Pública de Asunción, Paraguay, which concentrates biological samples collected throughout the country. On the basis of resources and time availability, we selected 50 acute-phase serum samples that were positive for DENV or CHIKV with PCR cycle threshold (Ct) values ≤28 at the time of diagnosis. Using epidemiologic data, we chose samples to represent geographic departments in Paraguay with the highest number of cases. ZIKV-positive samples with low Ct and historical samples for DENV were unavailable; hence, for DENV, we studied only the 2018 epidemic. The Dirección General de Vigilancia de la Salud de Paraguay provided temporal data on the incidence of chikungunya and dengue cases by department within Paraguay.

Virus Amplification and Whole-Genome Sequencing

We extracted viral RNA from the selected samples by using the QIAamp Viral RNA Mini Kit (QIAGEN, https://www.qiagen.com) and subjected the RNA to real-time reverse transcription quantitative PCR to detect CHIKV and DENV serotypes 1–4 as described previously (17–19). To increase the genome coverage, we selected only samples with Ct ≤35 for sequencing. Extracted RNA was converted to cDNA by using the Protoscript II First Strand cDNA Synthesis Kit (New England Biolabs, Inc., https://www.neb.uk.com) and random hexamer priming. We attempted whole-genome amplification by multiplex PCR as previously described (20).

We purified amplicons by using 1x AMPure XP Beads (Beckman Coulter, https://www.beckman.com) and quantified them on a Qubit 3.0 fluorimeter by using a Qubit dsDNA HS Assay Kit (ThermoFisher Scientific, https://www.thermofisher.com). We performed DNA library preparation by using a Ligation Sequencing Kit and Native Barcoding Kit (NBD103; Oxford Nanopore Technologies, https://nanoporetech.com). We generated sequencing libraries from the barcoded products by using the Genomic DNA Sequencing Kit SQK-MAP007/SQK-LSK208 and loaded them into an R9.4 flow cell (Oxford Nanopore Technologies).

Generation of Consensus Sequences

We base-called raw files by using Albacore software, demultiplexed and trimmed by using Porechop software (https://github.com) and then mapped with Burrows-Wheeler Aligner software to a reference genome. On the basis of PCR analyses, we used GenBank accession nos. KP164568 for CHIKV, KF672760 for DENV-1, and JN559741 for DENV-4 as reference sequences. To detect single-nucleotide variants to the reference genome, we applied Nanopolish software variant calling (https://nanoporetech.com) to the assembly. Nonoverlapped primer binding sites and
sites for which coverage was <20× were replaced with ambiguity code N.

**Phylogenetic Analyses**

We first investigated sequence genotypes by using the arbovirus genotyping tool (21). To investigate the origins and spatial dynamics of arboviruses in Paraguay, we downloaded all sequences assigned as CHIKV, DENV-1, and DENV-4 from GenBank. We excluded sequences without sampling date and location and sequences covering <50% of the virus genome. Sequence alignment was performed by using MAFFT (22) (FFT-NS-2 algorithm) and visually inspected in Aliview (23). We estimated maximum-likelihood phylogenies in IQ-TREE (24) by using the best-fit model of nucleotide substitution as indicated by the ModelFinder application (implemented in IQ-TREE). Branch support was assessed by the SH-like approximate-likelihood ratio test, and we submitted highly supported (>0.9) clades containing the DENV genotypes from Paraguay (Appendix 1 Figure 1, https://wwwnc.cdc.gov/EID/article/27/5/20-4224-App1.pdf) and the clades of CHIKV from the Americas to TempEst (25) to assess the strength of temporal signal in these data.

Time-scaled phylogenetic trees were inferred by using the BEAST package (26). We chose the uncorrelated relaxed molecular clock model as indicated by the marginal likelihood estimation model test procedure. We also used the codon-based SRD06 model of nucleotide substitution and the nonparametric Bayesian Skygrid coalescent model. A discrete phylogeographical model (27) was used to reconstruct the spatial diffusion of the virus across the compiled dataset sampling locations (Appendix 2, https://wwwnc.cdc.gov/EID/article/27/5/20-4224-App2.xlsx). Phylogeographic analyses were performed by applying an asymmetric model of location transitioning coupled with the Bayesian stochastic search variable selection procedure. We complemented this analysis with Markov jump estimation that counts location transitions per unit time along the tree. We ran Monte Carlo Markov chains long enough to ensure stationarity and an adequate effective sample size of >200.

**Results**

Of the 50 samples tested, 25 were positive for CHIKV, 14 for DENV-1, and 11 for DENV-4. For positive samples, the average PCR C<sub>t</sub> value was 26.36 (range 16–37). From the 50 samples, we were able to generate 33 complete or near-complete genome sequences (17 CHIKV and 16 DENV genomes) (Table). The GenBank accession numbers of newly generated sequences are MT038393–409 (CHIKV) and MT040672–87 (DENV). The collection dates of the samples sequenced were November 3, 2014, through July 10, 2018, and locations covered 15 municipalities and 8 departments (the first-level administrative subdivisions) of Paraguay (Figure 1). Women accounted for 58% of the samples, and the median patient age was 34 years. A TempEst analysis of all arbovirus lineages found here revealed a strong correlation between the sampling time and the root-to-tip divergence (Appendix 1 Figure 2).

**The CHIKV Epidemic in Paraguay during 2014–2018**

Of the 17 CHIKV genomes, 12 were classified as Asian genotype (sampled during 2014–2016) and 5 as East/Central/South African (ECSA) genotype (sampled during 2018). The oldest CHIKV sample analyzed (patient PY45) was obtained in November 2014 from the department of Amambay and was identified as an Asian genotype (Figure 1). However, autochthonous transmission of CHIKV was not confirmed until February 2015, followed by an increased number of reported infections (9) (Figure 2, panel A). Phylogeographic analysis revealed that the most likely origin of PY45 was Central or South America (Figure 3, panel A). The travel history for PY45 is in agreement with these results because the patient reported having visited Panama and San Andrés, a Colombian cluster of islands in the Caribbean region. In November 2014, another introduction of CHIKV in Paraguay was detected in the Central Department. That sequence (patient PY44) clustered with high support (posterior probability [PP] = 1) among sequences from Colombia and 1 sequence from Nicaragua, and the ancestral state of the most recent common ancestor (MRCA) of this clade was South America (PP = 0.8). Patient PY44 reported traveling to Cartagena, Colombia, supporting the origin estimated by the phylogeographic analysis.

In 2015, a large CHIKV epidemic occurred in Paraguay, resulting in ≈10,000 cases (suspected and confirmed); the main affected departments were Central, Asunción, and Paraguari (Figure 2, panel A). All genomes generated from the 4 samples from 2015 were classified as members of the Asian genotype and grouped together with high support (PP = 1) (Figure 3, panel A) in a clade for which time to MRCA (tMRCA) was October 2014 (95% highest posterior density [HPD] May 2014 to November 2014) (Figure 2, panel B). From our analysis, we estimated that the geographic origin of the variant circulating in Paraguay in 2015 was Puerto Rico (PP = 0.98) (Figure 3, panel A). The first patient with autochthonous
CHIKV infection in Paraguay is believed to be the housemaid of a family returning from Puerto Rico in October 2014, who sought healthcare services for symptoms of chikungunya fever. CHIKV infections were confirmed for the housemaid and the family, and our phylogenetic analysis confirmed this epidemiologic history.

The time distributions of CHIKV infection cases from 2016 were very similar to those in 2015 (Figure 2, panel A), although they did reveal that at least 2 lineages were circulating in the country (Figure 3, panel A). A new introduction is likely to have occurred in the Paraguari Department at the mean time point of November 2015 (95% HPD June 2015 to January 2016); the most likely place of origin was Central America (PP = 0.76) (Figure 3, panel A). However, the sequence from Amambay Department, isolated in 2014, is positioned basally to the Paraguari cluster, suggesting that the same variant persisted in the country up to 2016. Year-round persistence of a CHIKV strain is clearly observed in the Central/Asunción Department cluster in which the genome

| Sample | Virus              | Department, municipality | Collection date | Patient age, c/s | Cc | Reads | Genome coverage, % |
|--------|--------------------|--------------------------|-----------------|------------------|----|-------|-------------------|
| PY02   | CHIKV-Asian genotype | Paraguari, Yaguarón      | 2016 Jan 20     | 40/M             | 23.6 | 204,763 | 88.1              |
| PY03   | CHIKV-Asian genotype | Paraguari, Yaguarón      | 2016 Jan 21     | 67/M             | 16.9 | 215,137 | 87.0              |
| PY06   | CHIKV-Asian genotype | Paraguari, Yaguarón      | 2016 Feb 16     | 9/M              | 26.9 | 282,182 | 88.3              |
| PY07   | CHIKV-Asian genotype | Paraguari, Yaguarón      | 2016 Feb 19     | 34/F             | 29.7 | 267,784 | 87.2              |
| PY08   | CHIKV-Asian genotype | Paraguari, Yaguarón      | 2016 Feb 22     | 39/F             | 30.2 | 142,555 | 87.1              |
| PY09   | CHIKV-Asian genotype | Asunción, Asunción       | 2016 Mar 21     | 27/F             | 28  | 265,596 | 87.9              |
| PY12   | CHIKV-ECSA/BR      | Amambay, Pedro Juan Caballero | 2018     | 21/M | 29 | 236,285 | 84.8 |
| PY13   | CHIKV-ECSA/BR      | Amambay, Pedro Juan Caballero | 2018 Jun 25 | 35/F | 28 | 175,112 | 85.3 |
| PY15   | CHIKV-ECSA/BR      | Amambay, Bella Vista Norte | 2018 Jun 29 | 40/F | 34 | 17,030  | 70.9 |
| PY17   | CHIKV-ECSA/BR      | Amambay, Bella Vista Norte | 2018 Jul 3     | 22/M | 34 | 320,142 | 88.6 |
| PY18   | CHIKV-ECSA/BR      | Amambay, Bella Vista Norte | 2018 Jul 10    | 57/M | 29 | 315,588 | 86.0 |
| PY19   | DENV-4             | Guairá, Villarrica        | 2018 Apr 23     | 38/F | 23 | 22,041  | 82.3 |
| PY21   | DENV-4             | Guairá, Villarrica        | 2018 Apr 26     | 68/F | 22 | 21,042  | 96.0 |
| PY22   | DENV-4             | Guairá, Villarrica        | 2018 Apr 27     | 52/M | 22 | 13,213  | 96.0 |
| PY23   | DENV-4             | Central, San Lorenzo     | 2018 May 6      | 19/M | 27 | 11,548  | 74.6 |
| PY24   | DENV-1             | San Pedro, San Pedro De Ycuamandju | 2018 May 4 | 29/F | 20 | 7,265  | 89.1 |
| PY25   | DENV-4             | Central, San Lorenzo     | 2018 May 5      | 38/F | 21 | 17,299  | 96.0 |
| PY27   | DENV-4             | Alto Paraná, Domingo Martinez De Irala | 2018 May 9 | 27/F | 19 | 21,188  | 96.0 |
| PY28   | DENV-4             | Alto Paraná, Hernandarias | 2018 May     | 30/M | 21 | 22,800  | 86.6 |
| PY31   | DENV-4             | Alto Paraná, Hernandarias | 2018 May 22    | 14/M | 32 | 8,770   | 95.9 |
| PY32   | DENV-4             | Central, San Lorenzo     | 2018 May 31     | 28/M | 28 | 6,907   | 96.0 |
| PY33   | DENV-1             | Asunción, Asunción        | 2018 May 31     | 3/F  | 22 | 9,846   | 76.8 |
| PY34   | DENV-4             | Alto Paraná, Juan Leon Mallorquin | 2018 May 28 | 47/F | 26 | 7,945   | 96.0 |
| PY35   | DENV-1             | Itapúa, Encarnacion       | 2018 Jun 7      | 62/F | 23 | 119,293 | 89.1 |
| PY36   | DENV-1             | Itapúa, Encarnacion       | 2018 Jun 8      | 61/M | 23 | 6,448   | 76.8 |
| PY38   | DENV-1             | Itapúa, Cambyreta         | 2018 Jun 8      | 6/F  | 25 | 111,057 | 89.1 |
| PY43   | DENV-1             | Guairá, Villarrica        | 2018 Jun 4      | 53/F | 25 | 8,779   | 89.1 |
| PY44   | CHIKV-Asian genotype | Central, Luque           | 2014 Nov 30     | 33/F | 24 | 13,687  | 85.1 |
| PY45   | CHIKV-Asian genotype | Amambay, Pedro Juan Caballero | 2014 Nov 3 | 54/M | 24 | 12,214  | 86.6 |
| PY47   | CHIKV-Asian genotype | Central, Guarambaré       | 2015 Apr 26     | 25/M | 17 | 9,536   | 85.2 |
| PY48   | CHIKV-Asian genotype | Central, Fernando De La Mora | 2015 Apr 28    | 50/F | 18 | 6,002   | 86.6 |
| PY49   | CHIKV-Asian genotype | Central, Fernando De La Mora | 2015 May 3    | 12/F | 27 | 53,928  | 86.4 |
| PY50   | CHIKV-Asian genotype | Central, Fernando De La Mora | 2015 May 6    | 25/F | 16 | 49,813  | 86.8 |

*CHIKV, chikungunya virus; Cc, cycle threshold; DENV, dengue virus; ECSA/BR, East/Central/South African genotype from Brazil.
isolated in 2016 clustered with the genomes isolated in 2015 (Figure 3, panel A).

Besides the Asian genotype of CHIKV, an outbreak of the ECSA genotype occurred in Paraguay in 2018. After a year of very few CHIKV infections in 2017 (Figure 2, panel A), a new outbreak was observed in 2018 (although milder than that of 2015 and 2016), and the main affected department was Amambay. Five genomes from this department revealed circulation of the CHIKV ECSA genotype in 2018, and the mean tMRCA of this cluster was December 2017 (95% HPD July 2017 to April 2018) (Figure 2, panel B). The source of introduction of this new CHIKV lineage in Paraguay was estimated to be Brazil, most likely the Northeast Region (PP = 0.64), or perhaps the North Region (PP = 0.32) (Figure 3, panel B).

We also summarized all geographic locations that had significantly (Bayes factor >3) seeded new CHIKV lineages to Paraguay and superimposed it onto the tMRCA of the 3 main CHIKV clusters detected there (Figure 2, panel B). Most transitions in the Asian genotype occurred in 2014, when CHIKV was spreading rapidly through the Americas. Far fewer transitions were estimated to have occurred in 2015, which accords with the hypothesis that the 2016 outbreak in Paraguari was caused by a lineage already circulating in the country. For the ECSA genotype of CHIKV, the importations from Brazil were widespread between the middle of 2015 and the beginning of 2019. These widespread importations result from the long branch connecting the Paraguay ECSA cluster to the Brazilian sequences (Figure 3, panel B), increasing uncertainty in the relevant parameter estimates.

**Genetic Diversity of DENV in 2018**

The number of DENV cases in Paraguay during 2015–2018 shows a very similar pattern to that for CHIKV (Figure 2, panel A; Figure 4, panel A). Case numbers
were higher at the beginning of each year, except for 2017 when the DENV season was atypically mild. Our sampling from 2018 captured the 2 DENV serotypes (DENV-1 and DENV-4) circulating in the country (9), and molecular clock analyses estimated that DENV-4 was introduced in Paraguay just before the beginning of the 2018 outbreak, whereas DENV-1 was already circulating in 2017 (Figure 4, panel B).

DENV-1 and DENV-4 sequences from Paraguay clustered together with high support (PP>0.9) and belonged to genotypes V and II, respectively, which predominate in Latin America (Figure 5). The most likely origin of the DENV-1 strain circulating in Paraguay in 2018 was estimated to be Brazil (PP = 0.75) (Figure 5, panel A), and the mean tMRCA was estimated to be October 2016 (95% HPD February 2016 to May 2017) (Figure 4, panel B). DENV-4 was also estimated to have an origin in Brazil; the best-supported regions of origin were Central-West (PP = 0.64) and North (PP = 0.34) (Figure 5, panel B). The mean tMRCA of DENV-4 was September 2017 (95% HPD April 2017 to February 2018), ≈1 year later than DENV-1. Examining the DENV-4 cluster in Paraguay in more detail revealed that sequences from the Alto Paraná Department are basal and that sequences from the Central and Guairá Departments group together in a highly supported (PP>0.9) and distal cluster (Figure 5, panel B). Alto Paraná borders Brazil, and although not formally tested because of the small number of sequences, Alto Paraná could be the point of introduction of the current DENV-4 lineage into Paraguay.

**Discussion**

The first CHIKV outbreaks in the Americas (the Asian genotype) were reported for the French Caribbean islands of Saint Martin and Martinique in December 2013 (28). The virus rapidly spread throughout the Caribbean and Central America in 2014, and autochthonous transmissions were reported in almost all countries/territories of these regions. In 2014, Paraguay reported imported cases only, mostly in persons with a history of travel to Central America or the Caribbean. Our analysis of virus sequences from 2 of these persons with imported cases, and the virus phylogenetic relatedness to foreign viruses, matched the travel history with precision. In addition, the estimated origin of the first outbreak of CHIKV in Paraguay in 2015 agreed both in time (October 2015) and location (Puerto Rico) with the travel data collected by the Paraguay surveillance system.

The CHIKV epidemic in Paraguay in 2016 was very similar to that in 2015, when the most affected departments were Central, Asunción, and Paraguari. These neighboring departments are located in the most densely populated region of Paraguay, which might lead to consecutive outbreaks. In our sampling, most sequences from 2016 were from the Paraguari Department and formed a separate clade

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**Figure 2.** Chikungunya virus (CHIKV) outbreaks in Paraguay during 2014–2018 and the spatial–temporal history of virus diffusion. A) Total number of cases of CHIKV infection reported by epidemiologic week in the departments from which genome sequences were available. B) Location transitions to Paraguay inferred from the posterior distribution of phylogenetic trees by the Markov jump approach, and the time to most recent common ancestor for the CHIKV clusters detected in the country. Lines are colored according to the origin of the estimated transition toward Paraguay. Violin plots show 95% CIs with internal boxplots showing median and interquartile ranges. Brazil-N, Brazil North Region; Brazil-NE, Brazil Northeast Region; ECSA, East/Central/South African genotype.
Figure 3. Time-scaled phylogenetic trees of chikungunya virus (CHIKV) genomes isolated in the Americas. A) CHIKV Asian genotype; B) CHIKV East/Central/South African genotype from Brazil. Tips and internal branches are colored according to the most likely geographic location, and ancestral states were estimated by phylogeographic methods. Clusters relevant to the epidemic in Paraguay are shown in detail where the most likely ancestral state estimation is annotated. Asterisks indicate highly supported clusters (posterior probability >0.9).
from the 2015 epidemic, suggesting a new introduction event. However, because of the co-clustering of sequences from the Asunción and Central Departments isolated in 2015 and 2016, we cannot exclude year-round persistence of the virus. Models of *Aedes* spp. mosquito competence for DENV transmission have shown that the Paraguay climate is conducive to year-round persistence (29).

In Brazil, 2 distinct lineages of CHIKV were detected at the end of 2014, the ECSA genotype in the Northeast Region and the Asian genotype in the North Region (30). Whereas the CHIKV outbreak of the Asian genotype remained restricted to a small number of cases, the ECSA lineage spread throughout Brazil. In this study, we determined that the ECSA genotype was the causative agent of a CHIKV outbreak in Amambay (a Paraguay department on the Brazil border) in 2018. To our knowledge, these are the only 2 countries in the Americas to report outbreaks of both the Asian and ECSA genotypes to date. Our analysis revealed a mean time of entry into Paraguay of around December 2017, most likely from the Northeast Region in Brazil. However, because of the small number of samples from states in Brazil that border Paraguay, all such inferences of geographic origins should be interpreted with caution. It is possible that previous exposure to the CHIKV Asian genotype may have created some population immunity that restricted ECSA circulation in other parts of Paraguay (e.g., Central and Asunción Departments) in 2018. Unfortunately, no data on CHIKV seroprevalence in Paraguay are available to test this hypothesis. Amambay, on the other hand, reported few CHIKV cases during the epidemics of 2015 and 2016 (Figure 2, panel A), potentially enabling the emergence of the ECSA genotype in 2018.

The 2018 dengue season in Paraguay was dominated by DENV-1, reported in all departments (9). Three departments (Central, Alto Paraná, and Guaira) also reported the circulation of DENV-4. Our analysis suggests that the origin of both serotypes in Paraguay is Brazil, supporting findings of previous studies (31). Although our sampling was restricted to 2018, we observed that the tMRCA of DENV-1 was much earlier (October 2016), suggesting that this lineage may have persisted during the 2017 and 2018 seasons, when it may have been responsible for most cases (9). DENV-4, on the other hand, was introduced in September 2017, just before the start of the dengue season, and was responsible for few infections until the 2019–20 season, when it dominated the epidemic (32, 33).

Despite screening all publicly available (GenBank) sequences of DENV from the Americas, we found that the DENV datasets were sparsely distributed in time (DENV-4) or contained large temporal gaps (DENV-1) (Appendix 1 Figure 2), potentially biasing our results. The paucity of available DENV complete genomes in South America constrains the applicability of phylogenetic tools for studying virus population dynamics. It also highlights the value of intensifying sequencing efforts in line with the genomic surveillance approach and for real-time generating and sharing of data. The CHIKV datasets were much more comprehensive; for instance, the Asian lineage dataset analyzed included 291 genomes sampled during 2014–2018, representing 38 countries/territories (Appendix 2). The more comprehensive CHIKV datasets reflect the fact that CHIKV emerged in the Americas in the era of next-generation sequencing, when the development of numerous platforms reduced the cost and shortened the time from sample preparation to...
Figure 5. Time-scaled phylogenetic trees of dengue virus (DENV) serotypes 1 and 4 genomes isolated in Paraguay and related sequences. A) DENV-1; B) DENV-4. Tips and internal branches are colored according to geographic location, and ancestral states were estimated by phylogeographic methods. Clusters relevant to the epidemic in Paraguay are shown in detail where the most likely ancestral state estimation is annotated. Asterisks indicate highly supported clusters (posterior probability >0.9).
data generation. Hence, this increased availability of CHIKV virus genomes allowed a more detailed analysis of the virus’s spatiotemporal history in Paraguay. However, another study limitation is the convenience sampling used, with a narrow breadth, potentially biasing molecular clock dating and location ancestral reconstruction. The limited availability of stored samples from years before 2018, and their possibly limited RNA integrity, impaired genome sequencing from previous outbreaks. Nevertheless, we suggest that the 33 genomes generated here are representative of the current DENV diversity and the recent CHIKV evolutionary history in Paraguay.

Of note, the mean tMRCAs for the 3 clusters of CHIKV and the 2 clusters of DENV were estimated in the last trimester of the year (September–December), the start of arbovirus season in many tropical and subtropical regions in the Southern Hemisphere (e.g., Paraguay). A previous study (34) has modeled the timing and scale of arbovirus transmission potential and found that in many cities in Brazil with climates similar to that of Paraguay, transmission starts to increase around September. Thus, the tMRCAs estimated here probably reflect the onset of the arbovirus season, which peaks during January–March. Oddly, the 2017 season was marked by a noticeable reduction in cases of both DENV and CHIKV in Paraguay (Figure 2, panel A; Figure 5, panel A). This pattern was observed for dengue throughout the Americas, where cases decreased by 73% in 2017 compared with 2016 (35). It is possible that the mild season in 2017 might be explained by a transient strengthening of vector control interventions, implemented after the arrival of CHIKV and ZIKV in the Americas and the consequent public health emergency triggered by these pathogens. In addition, cross-immunity between ZIKV and DENV has been observed in the laboratory (36). Hence, population immunity to ZIKV after the 2015–2016 epidemic may have provided some transitory protection against DENV, resulting in lower incidence in 2017 (37,38), although this protection alone would not explain the decreased CHIKV cases in 2017 in Paraguay.

In conclusion, our study reveals a complex pattern of arbovirus circulation in Paraguay. We identify Brazil as a source of CHIKV and DENV lineages and show that other countries from South America and the Caribbean, mainly tourist destinations, were also hubs of virus spread toward Paraguay. Our sequencing and phylogenetic analyses proved to be powerful tools for revealing the transmission dynamics between the sampled locations and matched, with striking precision, available patient travel history. With support from the Pan American Health Organization, this project developed capacity-building skills in Paraguay, which can be applied in future arbovirus outbreaks.

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- Characteristics of Patients Co-infected with Severe Acute Respiratory Syndrome Coronavirus 2 and Dengue Virus, Buenos Aires, Argentina, March–June 2020
- Characteristics and Timing of Initial Virus Shedding in Severe Acute Respiratory Syndrome Coronavirus 2, Utah, USA
- Zika Virus–Associated Birth Defects, Costa Rica, 2016–2018
- *Plasmodium ovale wallikeri* and *P. ovale curtisi* Infections and Diagnostic Approaches to Imported Malaria, France, 2013–2018
- Symptom Profiles and Progression in Hospitalized and Nonhospitalized Patients with Coronavirus Disease, Colorado, USA, 2020
- Addressing COVID-19 Misinformation on Social Media Preemptively and Responsively
- Universal Admission Screening for SARS-CoV-2 Infections among Hospitalized Patients, Switzerland, 2020
- Excess Deaths during Influenza and Coronavirus Disease and Infection–Fatality Rate for Severe Acute Respiratory Syndrome Coronavirus 2, the Netherlands
- Murine Typhus in Canary Islands, Spain, 1999–2015
- *Plasmodium falciparum* Histidine-Rich Protein 2 and 3 Gene Deletions in Strains from Nigeria, Sudan, and South Sudan
- Hepatitis C Virus Transmission Clusters in Public Health and Correctional Settings, Wisconsin, USA, 2016–2017
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- Use of Commercial Claims Data for Evaluating Trends in Lyme Disease Diagnoses, United States, 2010–2018
- Highly Pathogenic Avian Influenza A(H5N8) Virus Spread by Short- and Long-Range Transmission, France, 2016–17
- Emergence of Lyme Disease on Treeless Islands, Scotland, United Kingdom
- SARS-CoV-2 Transmission between Mink (Neovison vison) and Humans, Denmark
- Shuni Virus in Cases of Neurological Disease in Humans, South Africa
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Appendix 1. Maximum likelihood phylogenetic tree with DENV1 (left) and DENV4 (right) complete genomes. Tips’ circles are colored according to the sampling continent and Paraguayan genomes generated in this study are shown. Highlighted clades were submitted to Bayesian phylogenetic analysis.
Appendix Figure 2. Regression analysis of the root-to-tip divergence against sampling date for the four viral lineages found in Paraguay in this study.