In conclusion, we report a rare case of *Spiroplasma* sp. blood infection in a patient after surgery for type A aortic dissection. *Spiroplasma* is an arthropod-infecting bacterium that may be part of the commensal microbiome of the human gut; there are 13 pieces of relevant information deposited into the gutMEGA database (http://gutmega.omicsbio.info) (10). *Spiroplasma* detection is challenging, and the discovery and diagnosis of emerging pathogens, such as the one we have described, can be aided by new technologies such as 16S rRNA gene sequencing and metagenomic next-generation sequencing.

This research was supported by the National Science and Technology Fundamental Resources Investigation Program of China (grant no. 2021FY100900).

**About the Author**

Dr. Ningning Xiu works in the Laboratory Department at Dongguan Kanghua Hospital, Dongguan, China. Her primary research interests are clinical microbiological laboratory diagnosis.

**References**


**Emergence of Dengue Virus Serotype 2 Cosmopolitan Genotype, Colombia**

David Martínez, Marcela Gómez, Carolina Hernández, Marina Muñoz, Sandra Campo-Palacio, Marina González-Robayo, Marcela Montilla, Norma Pavad-Escobar, Juan David Ramírez

Author affiliations: Universidad del Rosario, Bogotá, Colombia (D. Martínez, M. Gómez, C. Hernández, M. Muñoz, J.D. Ramírez); Universidad de Boyacá, Tunja, Colombia (M. Gómez); Centro de Tecnología en Salud (CETESA), Innovaseq SAS, Bogotá (C. Hernández); Laboratorio de Salud Pública, Secretaría de Salud Departamental Meta, Villavicencio, Colombia (S. Campo-Palacio, M. González-Robayo, M. Montilla, N. Pavad-Escobar); Universidad Cooperativa de Colombia, Villavicencio, Colombia (M. Montilla, N. Pavad-Escobar); Icahn School of Medicine at Mount Sinai, New York, New York, USA (J.D. Ramírez)

Using Oxford Nanopore technologies and phylogenetic analyses, we sequenced and identified the cosmopolitan genotype of dengue virus serotype 2 isolated from 2 patients in the city of Villavicencio, Meta department, Colombia. This identification suggests the emergence of this genotype in the country, which warrants further surveillance to identify its epidemic potential.
Dengue fever is a viral disease transmitted by *Aedes* spp. mosquitoes; the Americas are one of the most severely affected regions (1). The causative agent of dengue fever is the dengue virus (DENV), a positive-sense single-stranded RNA virus with a genome size of ≈10.7 kilobase. This virus is categorized into 4 distinct serotypes (DENV-1–4), classified on the basis of their surface antigens, and each serotype further consists of different genotypes that are phylogenetically distinct (2,3).

Recent epidemics in South America have been primarily attributed to the DENV-2 serotype, according to epidemiologic reports from the region (4). In Colombia, 70,418 cases of dengue fever have been reported as of August 2023; DENV-2 has been identified in most cases (5). Currently, this serotype consists of 5 genotypes named according to the region in which they circulate. Asian I and II genotypes are predominantly found in Asia, whereas the American genotype, which is no longer in circulation, was once prevalent in Central and South America. In the 1980s, the American genotype was replaced by the Asian-American genotype, which now circulates in Southeast Asia and the Americas. Last, the cosmopolitan genotype is noteworthy for its extensive global distribution, spanning 5 continents (6).

The cosmopolitan genotype has recently expanded in Africa and the Americas (7). This widespread dispersal has led to substantial intragenotype heterogeneity, reflecting the evolutionary forces acting within this genotype that are associated with its transmission. An outbreak attributed to the cosmopolitan genotype was reported in Madre de Dios Province, Peru, in 2019, coinciding with its recent expansion in Africa (8,9). In 2021, an additional 2 reports were documented in the states of Acre and Goiás in Brazil (4). Those reports shed light on a potential introduction route of the genotype into Brazil, specifically from the border with Peru (4). In 2023, the World Health Organization reported an outbreak in Latin America, generating a state of alert because of the increase in DENV cases (10). The genetic characteristics acquired during the extensive dissemination of the cosmopolitan genotype emphasize the need for further research into its diversity, evolution, and transmission dynamics within DENV-endemic areas.

In this report, we discuss 2 cases of the cosmopolitan genotype DENV-2 identified in Villavicencio, a city in the Meta department of Colombia. Of note, this department had the highest number of DENV cases in Colombia in 2023, accounting for 15.4% (10,859 cases) of total cases reported nationwide as of August (5). The 2 cases involved 2 young men with no travel history residing in suburban neighborhoods in southern Villavicencio (Figure, panel A). Both patients exhibited symptoms of fever, headache, myalgia, intense and continuous abdominal pain, and a platelet count of <100,000. Those symptoms align with the classification of DENV infection with warning signs, and dates of symptom onset were April 26, 2023, and

---

**Figure.** Phylogenetic analysis of dengue virus 2 cosmopolitan genotype, Colombia. A) Geographic location of the neighborhoods where the patients’ residences are situated. B) Maximum-likelihood tree rooted at the midpoint depicts the evolutionary relationships of the complete genome sequence of the dengue virus 2 cosmopolitan genotype identified in 2 patients from the city of Villavicencio in Meta department, Colombia (red circles), along with 1,001 publicly available sequences from GenBank. The highlighted blue area is shown in a time-resolved maximum-likelihood tree in expanded panel; colors represent different sampling locations. Scale bar indicates number of substitutions per site.
April 29, 2023.

Serum samples were collected and sent to the microbiology laboratory at Universidad del Rosario in Bogotá, Colombia for processing. We extracted viral RNA using the Quick-RNA Viral Kit (Zymo Research, https://zymoresearch.eu). The infection was confirmed to be caused by the DENV-2 serotype using the previously described protocol (Appendix, https://wwwnc.cdc.gov/EID/article/30/1/23-0972-App1.pdf). We performed whole-genome sequencing using MinION (Oxford Nanopore Technology, https://nanoporetech.com) to determine the corresponding genotype classification and to conduct subsequent analysis of the local distribution of DENV (Appendix). The Technical Research Committee and Ethics Research Board from Universidad del Rosario in Bogotá, Colombia approved the protocol implemented in this study (approval no. DVO005 1585-CV142).

We conducted an initial maximum-likelihood phylogenetic analysis to identify the genotype. The analysis revealed that the sequences obtained from the patients were closely related, belonged to the DENV-2 cosmopolitan genotype, and were placed within the South America sequences found in Tefé and Tabatinga, Brazil, and Madre de Dios in Peru (Figure, panel B).

Further examination using a time-resolved maximum-likelihood tree demonstrated that those sequences were closely related to sequences reported in the Tabatinga province in Brazil. The bootstrap support for this relationship was 95% (Figure, panel B). This finding suggests potential cross-border transmission in the Tabatinga province, highlighting the possibility of viral spread across borders.

In conclusion, although genetic data alone cannot provide conclusive evidence about the directionality of the introduction of the DENV-2 cosmopolitan genotype, insights gained from phylogenetic reconstruction and temporal information suggest a potential introduction from Tabatinga, Brazil, with subsequent spread northwards in Colombia. Tabatinga is located in the tripartite border region between Brazil, Colombia, and Peru adjacent to the Amazonas department in southern Colombia. Because of the limited research available on the cosmopolitan genotype, our understanding of its effects on dengue disease dynamics in Colombia remains incomplete. Further investigations are required to gain a more comprehensive insight into its potential for local, regional, and global epidemics. Our findings highlight the importance of implementing robust genomic surveillance in the region, especially considering the ongoing outbreak in Latin America.

This work was supported by the Colombian Ministerio de Ciencia, Tecnología e Innovación Minciencias (grant no. 143889685192-2021).

D.M. and J.D.R. conceived the study; M.G., C.H., and M.M. analyzed the data; and S.C.P., M.G.R., M.M., and N.P.E. collected the samples. All authors have read and agreed to the published version of the manuscript.

About the Author

Mr. Martínez is a biologist and master’s student in natural sciences at Universidad del Rosario, Bogotá, Colombia. His primary research interest is the genomic surveillance of dengue virus. Dr. Ramírez is an associate professor at Universidad del Rosario and the Icahn School of Medicine at Mount Sinai. His primary research interests are the genomic surveillance and evolution of viruses and parasites.

References

Mycobacterium senegalense Infection in Kidney Transplant Patient with Diabetes, Memphis, Tennessee, USA

Nupur Singh, Reeti Khare, Shirin Mazumder

Author affiliations: University of Tennessee Health Science Center, Memphis, Tennessee, USA (N. Singh, S. Mazumder); National Jewish Health, Denver, Colorado, USA (R. Khare); Methodist University Hospital, Memphis (S. Mazumder)

DOI: https://doi.org/10.3201/eid3001.231013

Mycobacterium senegalense, also referred to as M. conceptionense, is a nonpigmented rapid-growing mycobacterium belonging to the M. fortuitum group, which was first isolated in 2006 from a post-traumatic osteitis inflammation in France (1,2). Only a handful of M. conceptionense cases are readily identifiable in existing literature. Infections can manifest with pulmonary involvement but more commonly manifest as skin or subcutaneous infection, such as after face rejuvenation surgery, breast augmentation surgery, gastric carcinoma resection, or subcutaneous ankle infection (3–6). Mycobacterial species like M. senegalense have been found in irrigation systems, soil, domestic and wild animals, and dairy products (7,8). Cases of M. senegalense infection have been observed in France, Iran, Taiwan, South Korea, Japan, and the United States, demonstrating an unidentifiable pattern of regional bacterial prevalence.

Establishing an accurate diagnosis of M. senegalense infection is incredibly difficult, requiring histological examination and extensive mycobacterial cultures (9). The limited susceptibility data also mean an optimal therapy has not been completely established, which leaves certain patient populations, particularly the elderly and immunocompromised, susceptible to increased illness and death from M. senegalense infection (10). We report a complicated case of M. senegalense infection in a patient with a previous kidney transplant and insulin-dependent diabetes mellitus in Memphis, Tennessee, USA.

A 70-year-old Black woman with end-stage kidney disease sought care for a painful, swollen, abdominal wall abscess. She had first noticed the lesion ≈3 weeks before in the left mid-abdomen, where she frequently injected insulin. The patient denied any recent travel, drainage at the site, or fever. The 7-×4-cm abscess was drained the next day without complications, and we sent the custard-like purulent material for laboratory testing.

The patient had undergone a right-sided cadaveric renal transplant 8 years before; her immunosuppressive regimen consisted of tacrolimus, mycophenolate, and prednisone. The patient’s past diagnoses at the time of infection included type 2 diabetes mellitus, lupus, hypertension, hyperlipidemia, sleep apnea, and coronary artery disease. Insulin injections create small open wounds where pathogens can enter and cause infection. An environmental source of the infection was not sought. The hospital microbiology laboratory detected acid-fast bacilli on direct AFB smear. The patient immediately began empiric antimicrobial drugs, including doxycycline (100 mg 2×/d) and levofloxacin (250 mg 1×/d), adjusted for her creatinine clearance.

The isolate was sent to the National Jewish Mycobacteriology Reference Laboratory (Denver, Colorado, USA) for confirmation and susceptibility testing. Sanger sequencing analysis was performed; BLAST (https://blast.ncbi.nlm.nih.gov/Blast.cgi) testing of the rpoB sequence results against the public GenBank database identified M. senegalense (>99% homology to existing sequences). A line probe assay for common nontuberculous mycobacteria, GenoType NTM-DR (Hain Lifescience, https://www.hain-lifescience.de),