Two cases of newly characterized Neisseria species, Brazil

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Abstract: We describe 2 human cases of infection with a new Neisseria species (putatively N. brasiliensis), 1 of which involved bacteremia. Genomic analyses found that both isolates were distinct strains of the same species, were closely related to N. iguanae, and contained a capsule synthesis operon similar to N. meningitidis.

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closest reported autochthonous human cases were in southwestern Hungary (5).

E. multilocularis tapeworms have been reported in foxes in western and central Croatia (6) and likely is in eastern areas, such as Vukovar, because it was found in 17.9% of foxes and 14.3% of golden jackals in the region of Serbia directly across the Danube River from Vukovar (7). Since 2013, rabies vaccination has increased in Croatia, which might give the fox population an opportunity to expand and increase transmission of E. multilocularis tapeworms to humans, as noted in Switzerland (8).

Correct diagnosis for this patient took 2.5 years because radiologic findings were inconsistent with cystic echinococcosis and clinicians assumed that was the only type of human echinococcosis in Croatia (9). This case highlights the need for clinicians to include alveolar echinococcosis in differential diagnosis of liver lesions. Imaging provides the first-line approach to such a diagnosis and serology provides strong complementary support. Our case also highlights the usefulness of considering pleural effusion and analyzing archival biopsies to retrospectively diagnose alveolar echinococcosis.

About the Author

Dr. Dušek is an attending physician in the University Hospital for Infectious Diseases in Zagreb, Croatia. Her research interests include viral hepatitis and infectious diseases in immunocompromised patients, especially patients with solid organ or stem cell transplants.

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Two Cases of Newly Characterized Neisseria Species, Brazil

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We describe 2 human cases of infection with a new Neisseria species (putatively N. brasiliensis), 1 of which involved bacteremia. Genomic analyses found that both isolates were distinct strains of the same species, were closely related to N. iguanae, and contained a capsule synthesis operon similar to N. meningitidis.
Neisseria is a genus containing diverse organisms; most are rarely pathogenic. N. meningitidis and N. gonorrhoeae are the most clinically relevant species. The polysaccharide capsule is the most critical meningococcal virulence factor, a vaccine target, and the basis for classifying meningococci into serogroups (1). During routine laboratory-based public health surveillance in Brazil, we identified 2 cases of infection caused by a previously uncharacterized species of the Neisseria genus.

Clinicians reported 2 cases to the National Reference Laboratory, Adolfo Lutz Institute (IAL), São Paulo, Brazil. Case-patient 1 was a 64-year-old man from Rio Grande do Sul state, Brazil, who, in June 2016, had congestive heart failure with bilateral pulmonary infiltrates and pleural effusion on chest radiograph. Case-patient 2 was a 74-year-old woman with leprosy from Paraná state, Brazil, who, in February 2016, developed a polymicrobially infected ulcer of the left lower extremity. The 2 cases were separated in time and by >400 km and had no known epidemiologic link.

Overnight cultures of blood from case-patient 1 and ulcer exudate from case-patient 2 on brain–heart infusion agar containing 10% chocolate and horse blood at 37°C in the presence of 5% CO2, both revealed brownish colonies uncharacteristic of N. meningitidis. We identified both isolates (N.95-16, from case-patient 1, and N.177-16, from case-patient 2) as gram-negative glucose-fermenting diplococci with positive catalase and oxidase tests. The isolates fermented maltose, lactose, sucrose, and fructose but not mannose; they reduced nitrate and produced a starch-like polysaccharide detected with Gram’s iodine but did not produce DNase. Assessment by matrix-assisted laser desorption/ionization time-of-flight mass spectrometry found no species match; the closest matches were N. iguanae and the proposed N. weixii, isolated from the intestinal contents of a Tibetan Plateau pika (Figure, panel A) (PubMed identification 56407–56409; GenBank accession no. CP023429). Both genomes shared identical rMLST profiles (rST 61343); the 4 proposed N. weixii genomes shared only 1–2 alleles of 53 rMLST loci with these isolate genomes, and N. iguanae shared no rMLST alleles. Both genomes contained an intact capsule gene cluster (cps) that was similar in gene organization and sequence identity to N. meningitidis (Figure, panel B). The ctrA-csaA/cssA promoter region was conserved in both isolates. However, both genomes contained only 1 copy of galE-rfbCAB (Region D), compared with 2 copies found in meningococcal reference genomes; the tex gene was located >10 kb outside cps, upstream from ctrD (Figure, panel B). The 2 isolates differed in their sequence of sialic acid biosynthesis genes within region A; isolate N.95-16 contained cssABC genes that shared 98% amino acid identity with the meningococcal serogroup X reference strain α388 (1), and isolate N.177-16 contained cssABC-csb genes that shared 99% amino acid identity with serogroup B reference strain H44/76 (Figure, panel B). The cps differences observed between the isolates were similar to the mosaic recombination pattern associated with meningococcal capsular switching (9). Taken together, the presence of cps genes sharing substantial similarity to meningococcal homologs suggests that both isolates have the potential to synthesize meningococcal-like capsules.

In summary, we describe 2 sporadic cases of a new Neisseria species (which we propose to name
Neisseria brasiliensis, 1 of which also involved bacteremia. Both genomes contain an intact repertoire of genes for capsule synthesis, a key meningococcal virulence factor. The significance of capsule genes and potential capsule synthesis in nonmeningococcal Neisseria is unknown (10). Continued surveillance is required to establish the pathogenic potential and host range for this apparent new species.

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Hepatitis A Virus Genotype IB Outbreak among Internally Displaced Persons, Syria

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In 2018, a hepatitis A virus outbreak was identified among internally displaced persons in Syria. Sequence analysis based on the viral protein 1/2A junction revealed that the causative virus belonged to genotype IB. A high displacement rate, deteriorated sanitary and health conditions, and poor water quality likely contributed to this outbreak.

Hepatitis A virus (HAV) is the leading cause of acute hepatitis infections worldwide, infecting ≈1.5 million persons annually (1). Symptoms, which are usually mild, include nausea, vomiting, abdominal pain, restlessness, body weakness, myalgia, loss of appetite, and fever. However, HAV may progress into fulminant liver failure, necessitating liver transplant. Generally, HAV is self-limiting (2). HAV (genus Hepatovirus, family Picornaviridae) is a nonenveloped virus with a single-stranded, positive-sense RNA linear genome (7.5 kb). The viral protein (VP) 1/2A junction (168 nt) is used to classify HAV into 6 genotypes: I–III (subgenotypes A and B) of human origin and IV–VI of simian origin (3). Genotype IA is the most commonly reported worldwide, whereas genotype IB is predominant in the Middle East (4–6).

On September 9, 2018, the governorate of Aleppo, Syria, informed the World Health Organization office in Syria that internally displaced persons (IDPs; displaced since early 2018) and local host community members in Tal Refaat, Fafin, and surrounding areas in the northwestern and western parts of Aleppo were experiencing a suspected hepatitis outbreak. The affected area included 17 locations in Azaz and Jabal Sem’an districts in western Aleppo (Appendix, https://wwwnc.cdc.gov/EID/article/26/2/19-0652-App1.pdf). Outbreak field investigation found sporadic cases of the disease among IDPs starting July 21, 2018; as of November 8, a total of 638 cases of suspected acute hepatitis infection had been reported. Most patients (98.59%) were <15 years of age and the rest 16–54 years of age. A total of 105 patients (16.5%) were admitted into the Fafin hospital; no fatalities were reported. No field investigations were performed in the first half of 2018 because of the crisis that led to weakness in the routine surveillance system.

A total of 48 unidentified serum and plasma samples were collected from 24 IDP children with suspected hepatitis and sent to the laboratory on October 29. The specimens originated from 3 locations in Syria: 13 from Fafin camp in Aleppo, 6 from eastern rural Daraa, and 5 samples from rural Quneitra. Even though the main outbreak was in the Aleppo governorate, Daraa and Quneitra were also experiencing a notable upsurge in reported cases of suspected acute hepatitis infection. For this reason, additional samples were collected from these governorates.

We analyzed the serum specimens by serology (total HAV antibodies and HAV IgM) using the enzyme-linked fluorescent assay VIDAS (bioMérieux Diagnostics, https://www.biomerieux-diagnostics.com) and the plasma specimens by real-time reverse transcription PCR (RT-PCR) for the detection of HAV (using the HAVNET protocol) and hepatitis E virus (HEV) (7). Seven samples had insufficient volume to perform both total HAV antibody and HAV IgM tests; thus, only the IgM test was performed. Overall, 19 plasma specimens were positive for HAV and none for HEV by PCR (Table). Eighteen serum specimens had detectable HAV IgM. All the specimens with sufficient volume (n = 17) were positive for HAV and none for HEV by PCR (Table). Eighteen serum specimens had detectable HAV IgM. All the specimens with sufficient volume (n = 17) were positive for HAV and none for HEV by PCR (Table).