Occupational Exposure to Zoonotic Tuberculosis Caused by *Mycobacterium caprae*, Northern Greece, 2019

Dimitrios Papaventsis, George Dougas, Ourania Kalkouni, Simona Karabela, Katerina Manika

Author affiliations: Sotiria Chest Diseases Hospital, Athens, Greece (D. Papaventsis, S. Karabela); National Public Health Organization, Athens (G. Dougas, O. Kalkouni); G. Papanikolaou Hospital, Thessaloniki, Greece (K. Manika)

DOI: https://doi.org/10.3201/eid2707.204399

Pulmonary tuberculosis caused by *Mycobacterium caprae* was diagnosed in a 65-year-old goat breeder from northern Greece. This case represents a documented occupational transmission of *M. caprae* and highlights the importance of enhanced laboratory screening and increased surveillance for zoonotic tuberculosis control.

*Mycobacterium caprae*, a member of *Mycobacterium tuberculosis* complex (MTBC), causes caprine tuberculosis (TB) (1). *M. caprae* has been isolated mainly in continental Europe and infects animals and humans (2–4). Although clinically indistinguishable from *M. tuberculosis* or *M. bovis*, *M. caprae* causes nearly one third of *M. bovis*–associated TB cases (5). Bovine TB control and pasteurization have made zoonotic TB rare; however, occupational risk exists for livestock farmers, veterinarians, slaughterhouse workers, butchers, and other persons working in close contact with livestock.

In August 2019, a 65-year-old male goat breeder from northern Greece was admitted to the Aristotle University Pulmonology Department at G. Papanikolaou Hospital (Thessaloniki, Greece) for weight loss, productive cough, and fever <38.5°C that was not responsive to antibiotics. He reported chronic alcohol abuse, smoking, and had no other medical history. He reported living in the goat shelter, occasionally consuming unpasteurized milk, and having no contact with or hunting practices involving other animal species. Laboratory tests showed normocytic anemia (hemoglobin 10.4 g/dL) and an erythrocyte sedimentation rate of 112 mm/h. Chest radiograph and computed tomography scans revealed upper lobe cavities and infiltrations, predominantly on the left side (Figure, panels A–C). A tuberculin skin test result was 12 mm in size. TB was suspected; sputum samples were sent for molecular and bacteriologic testing to the National Reference Laboratory for Mycobacteria at Sotiria Chest Diseases Hospital (Athens, Greece). Acid-fast stain results were positive and MTBC detected by using TRCReady-80 (Tosoh Corp., https://www.tosoh.com). We determined drug sensitivity on a strain grown on sodium pyruvate-LJ medium by Genotype MTBDRplus (Hain LifeScience, https://www.hain-lifescience.de) and on solid media. We documented sensitivity to first-line anti-TB drugs and pyrazinamide. Because of the patient’s occupation, laboratory investigation included zoonotic species. GenoType MTBC differentiated the isolate as *M. bovis* subspecies *caprae*.

We used the optimized 24-loci Genoscreen MIRU-VNTR (mycobacterial interspersed repetitive unit–variable-number tandem-repeat) Typing Kit (Genoscreen, https://www.genoscreen.fr) on crude DNA, using 6 quadruplex PCR and fluorescent primers with capillary electrophoresis. Identification was performed with the MIRU-VNTRplus database (6). Because no match was detected after initial best-match analysis, we used a tree-based identification scheme by applying the UPGMA method and using the database reference strains. Identification was confirmed with a unique MIRU-VNTR pattern (*M. caprae_Kilkis:255326322553434243231432*) showing close phylogenetic match with *M. caprae* reference strains from central and eastern Europe and clear genetic distance from the Iberian cluster, mainly characterized by absence of spoligotype spacers 30–33 (3) (Appendix Figure, https://wwwnc.cdc.gov/EID/article/27/7/20-4399-App1.pdf).

The patient received a standard anti-TB regimen for 9 months and responded favorably. Cultures and acid-fast bacilli turned negative 10 days after treatment and were negative 2.5 months after treatment. Erythrocyte sedimentation rate dropped to 25 mm/h, and the patient gained 17 kg. Chest radiograph and computed tomography scans showed remarkable improvement (Figure, panels D–F). Contact tracing did not reveal further human cases. Veterinary investigation identified no tuberculin skin test reactors among >300 goats of the epidemiologically linked herd.

Zoonotic TB causes an estimated 147,000 new human cases and 12,500 deaths annually worldwide (7). Zoonotic TB is a reemerging and underrecognized infection in Europe; 170 confirmed human cases were reported in 2018 (0.05 cases/100,000 population) (4). Among countries not officially TB-free, Greece, Ireland, and Spain reported a plateaued prevalence of 2%–5% in cattle herds over the past decade (4). The actual global burden of *M. caprae* disease is further underestimated because of differences in laboratory...
capacity and lack of routine surveillance data. Zoonotic TB is a public health hazard, resulting in serious economic losses and having a substantial effect on poor and marginalized communities (7).

Contaminated food and airborne transmission pose considerable risks to persons in contact with infected animals or animal products (7). In this study, caprine TB was occupationally acquired by a goat farmer with a history of routine close contact with livestock. Pyrazinamide sensitivity and lung localization could have caused an initial misdiagnosis of *M. tuberculosis* infection. Since 2019, enhanced laboratory screening and surveillance for high-risk patients related to animal breeding have been used in Greece. Timely application of bacteriologic and genotyping techniques, less frequently used in routine laboratory investigation until recently, highlighted the role of *M. caprae* as a human pathogen. MIRU-VNTR revealed a unique isolate similar to patterns reported in animals in Eastern Europe and the Balkans (3,8). Data on *M. caprae* zoonotic TB in Greece are limited. A dairy goat farm outbreak in northern Greece was the first documented in 2005; *M. caprae* caused pneumonia among animals, and the mortality rate reached 92% (9). Further, Neonakis et al. (10) reported a human *M. bovis* isolate identified as *M. bovis* subsp. *caprae* in a Geno-Type MTBC assay evaluation study.

*M. caprae*, a recently identified separate MTBC species, should be carefully addressed nationally and internationally. Key risk populations in countries not officially TB-free and with sizeable goat populations must be identified, occupational history sought, testing capacity extended, and monitoring and reporting improved. A One Health approach that highlights interdependence of human and animal health sectors is needed to curb the spread of this pathogen.

**Acknowledgments**

We thank all members of the staff involved in the investigation of this study at the National Reference Laboratory for Mycobacteria, Sotiria Chest Diseases Hospital, Athens; the National Public Health Organization, Athens; the Aristotle University Pulmonary Department, G. Papanikolaou Hospital, Thessaloniki; the General Directorate of Veterinary Medicine, Athens; and the Directorate of Public Health, Kilkis Prefecture, Kilkis, Greece.

**About the Author**

Dr. Papaventsis is a medical director in Clinical Microbiology–Laboratory Medicine at the National Reference Laboratory for Mycobacteria, Sotiria Chest Diseases Hospital, Athens, Greece.
Outbreak of Rabbit Hemorrhagic Disease Virus 2 Infection, Ghana

Aruna Ambagala, Patrick Ababio, Lindsey Lamboo, Melissa Goolia, Oliver Lung, Yohannes Berhane, Theophilus Odoom

Author affiliations: National Centre for Foreign Animal Disease, Canadian Food Inspection Agency, Winnipeg, Manitoba, Canada (A. Ambagala, L. Lamboo, M. Goolia, O. Lung, Y. Berhane); Veterinary Services Directorate, Accra Laboratory, Accra, Ghana (P. Ababio, T. Odoom)

DOI: https://doi.org/10.3201/eid2707.210005

In September 2019, high mortality in commercial rabbits was reported in the Greater Accra Region of Ghana. Rabbit hemorrhagic disease virus 2 phylogenetically related to isolates from 2015–2017 outbreaks in the Netherlands was confirmed as the causative agent. The virus has not yet been detected in native rabbits in Ghana.

Rabbit hemorrhagic disease (RHD) is an acute, fatal, highly contagious viral hepatitis in European rabbits (Oryctolagus cuniculus) (1). It causes severe economic losses in the rabbit meat and fur industries and can have a substantial negative ecological impact on wild rabbit populations and their predators. The causative agent, RHD virus (RHDV; family Caliciviridae, genus Lagovirus) has a single-stranded positive-sense RNA genome ≈7.4 kb in length. Pathogenic RHDV strains exist in 2 main genotypes, GI.1 and GI.2 (RHDV2). RHDV GI.1 has several variants: GI.1a (proposed as G6/RHDVa), GI.1b (G1), GI.1c (G2), and GI.1d (G3–G5) (2). It is considered enzootic in domestic and wild European rabbits in Asia and Europe; sporadic outbreaks occur in the Americas, Middle East, and Africa (1). RHDV2 GI.2, a variant first reported in France in 2010 (3), differs antigenically from RHDV GI.1 and is therefore considered a distinct serotype. RHDV2, which has replaced RHDV GI.1 in many countries in Europe, infects rabbits of all ages and crosses the species barrier to affect non-European rabbit species (4).

Rabbit production, because of its low costs, has been promoted to reduce poverty in Africa but has been threatened by RHD outbreaks since the late 1980s. In 2015, RHDV2 was detected on Tenerife in the Canary Islands (5). In April 2017, an outbreak was reported in northern Morocco (6) caused by a recombinant GI.1b/GI.1b/GI.2 RHDV2 strain closely related to isolates identified in Portugal in 2014.
Appendix Figure. Phylogenetic tree of a *Mycobacterium caprae* isolate from a 65-year-old goat breeder from Kilkis, northern Greece (M.caprae_Kilkis: MIRU-VNTR 24-loci pattern 255326322553434243231432). The tree was calculated and drawn by using the UPGMA method and utilizing the MIRU-VNTRplus database reference strains (1). VNTR patterns were first searched against the MIRU-VNTRplus database using a default stringent distance cut-off of 0.17, which corresponds to a tolerance of ≤4 locus differences. Since no match was detected after the initial best-match analysis, a tree-based identification was used. Species, Lineage, MLVA Mtbc15-9, SpolDB4 Type, and Spoligo pattern information has been included. The patient’s isolate shows tight genetic clustering with *M. caprae* reference strains that dominate in central and eastern Europe (spoligotypes lacking spacers 1, 3 to 16, 28, and 39 to 43, BOV_4-CAPRAE, SIT647, SB0418). A clear genetic separation from the Iberian isolates, mainly characterized by the additional absence of spacers 30-34 (BOV-CAPRAE, SIT644/SB0157) has been demonstrated. The *M. bovis* BCG control is also depicted. Scale bar represents 10% sequence divergence. BCG, Bacillus Calmette–Guérin; MLVA, Multiple-locus variable-number tandem repeat analysis; VNTR, variable number of tandem DNA repeats.
Reference