facilities to treat EVD (9,10). A tiered nationwide network of healthcare facilities that can rapidly identify, isolate, and treat patients with EVD has been established to improve the nation’s preparedness for EVD and can serve as a valuable resource for future outbreaks of other highly infectious diseases. Ongoing resources will be needed to sustain the readiness of such a network.

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Address for correspondence: John J. Lowe, College of Public Health, University of Nebraska Medical Center, 984388 Nebraska Medical Center, Omaha, NE, USA 68198; email: jjlowe@unmc.edu

Detection of Influenza D Virus among Swine and Cattle, Italy

Chiara Chiaipponi,1 Silvia Faccini,1 Aurora De Mattia, Laura Baioni, Ilaria Barbieri, Carlo Rosignoli, Arrigo Nigrelli, Emanuela Foni

Author affiliations: Istituto Zooprofilattico Sperimentale della Lombardia ed Emilia Romagna, Brescia, Italy (C. Chiaipponi, S. Faccini, A. De Mattia, L. Baioni, I. Barbieri, C. Rosignoli, A. Nigrelli, E. Foni); World Organisation for Animal Health Reference Laboratory for Swine Influenza, Parma, Italy (C. Chiaipponi, L. Baioni, E. Foni)

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To the Editor: Recent studies have identified a new genus of the Orthomyxoviridae family (1–5). The virus, distantly related to human influenza C virus, has been provisionally designated as influenza D virus. This novel virus was identified for the first time in pigs with influenza-like illness (I), but subsequent serologic and virologic surveys have suggested cattle as a possible reservoir (2–4). Moreover, the virus was shown to infect ferrets used in laboratories as surrogates for humans when investigating influenza infection (I). In a serologic study conducted on 316 human samples, low antibody titers and a low level of positive samples (1.3%) were detected (I), suggesting that humans are a possible host to be studied in depth. To investigate the circulation of influenza D viruses among pigs and cattle in Italy, we performed biologic and virologic tests on clinical samples collected from respiratory outbreaks in Po Valley, the area in Italy with the highest density of swine and cattle farms.

We screened clinical specimens from swine (n = 150) and cattle (n = 150) for influenza D virus by reverse transcription quantitative PCR (I). Three nasal swab samples were found positive: 1 from a sow and 2 from cattle, collected from 3 farms located at linear distances ranging from 47 to 80 km. All positive samples were confirmed by partial polymerase basic 1 gene sequencing and submitted to viral isolation in cell cultures as previously described (5,6). The virus was isolated on CACO-2 and HRT18 cell cultures only from the sow sample (D/swine/Italy/199723-3/2015). Cell cultures were tested by using reverse transcription quantitative PCR. Viral RNA was isolated from clinical samples or cell culture by using One-For-All Vet Kit (QIAGEN, Milan, Italy). Full-genome amplification from influenza D virus–positive samples was achieved as previously described (3). A sequencing library of the purified amplicons was prepared by using NEXTERA-XT kit and

1These authors contributed equally to this article.
sequenced by using a MiSeq Reagent Kit v2 in a 250-cycle paired-end run (both from Illumina Inc., San Diego, CA, USA). Sequencing reads were assembled de novo or by using D/swine/Oklahoma/1334/2011 (GenBank accession nos. JQ922305–JQ922311) as a template by Seqman NGen DNASTAR version 11.2.1 (DNASTAR, Madison, WI, USA). Gene sequences from the 3 influenza D viruses isolated in Italy and all the available influenza D virus sequences retrieved from GenBank were aligned with ClustalW by using MEGA5 (7). We analyzed the predicted amino acid sequences for each gene.

Phylogenetic trees of the individual segments were inferred by using the maximum-likelihood method implemented in the IQ-TREE package 0.9.6 (8). The robustness of the maximum-likelihood trees was evaluated by bootstrap analysis by comparison to 1,000 bootstrap samples. The swine isolate D/swine/Italy/199723-3/2015 was fully sequenced (GenBank accession nos. KT592530–KT592536). Full genome sequences of D/bovine/Italy/1/2014 (GenBank accession nos. KT592516–KT592522) and D/bovine/Italy/46484/2015 (GenBank accession nos. KT592523–KT592529) were obtained directly from the nasal swab samples. The 7 genomic segments of each of the 3 influenza D virus genomes encode the proteins of polymerase basic subunits 1 and 2, polymerase 3, glycoprotein, nucleoprotein, matrix 1, matrix 2, and nonstructural proteins 1 and 2. These segments contain 772, 755, 710, 664, 552, 387, 246, 243, and 184 aa residues, respectively, similar to viruses of their counterparts of the isolates documented in Asia and America. The predicted amino acid sequence of the hemagglutinin gene shows unique features for the strains isolated in Italy: V in position 289, K409R, I563L, and A652V. In the apex of the hemagglutinin 1 receptor-binding domain of the glycoprotein-predicted proteins, position 212 is occupied by K, as previously observed for D/swine/Oklahoma/1334/2011 (5). Moreover, the 3 isolates from Italy share unique mutations in the polymerase basic 1 gene (R191G, F278S, R444G) and in the polymerase 3 predicted proteins (I194V, M596V). D/swine/Italy/199723-3/2015 shows no unique amino acid difference to bovine strains, and its gene segments cluster with influenza D viruses isolated from cattle, suggesting the circulation of this virus among cattle and swine in Italy. Phylogenetically, all 7 segments of the strains isolated in Italy clustered with D/swine/Oklahoma/1334/2011, showing no sign of reassortment (Figure).

Our findings show that influenza D viruses circulate among swine and bovine herds in Italy affected by respiratory disease. Genetic analysis highlights that the swine and bovine influenza D viruses are very closely related, both belonging to the D/swine/Oklahoma/1334/2011 cluster. Further studies are ongoing to better understand the epidemiology, virology, and pathobiology of influenza D virus in swine and cattle, especially concerning the evidence that Koch’s postulates are fulfilled for this agent. Implications in zoonotic aspects of influenza D virus infections will be also considered in ongoing research.

**Figure.** Phylogenetic trees of the 7 genes of influenza D viruses obtained from 1 sow and 2 cattle in Italy (vertical bars) and comparison isolates retrieved from GenBank. A) Polymerase basic (PB) 2: 2,319 nt; B) PB1: 1,434 nt; C) P3: 2,133 nt; D) glycoprotein hemagglutinin-esterase: 1,995 nt; E) nucleoprotein: 1,659 nt; F) polymerase 42: 1,164 nt; G) nonstructural: 732 nt. Genes were trimmed and aligned, then phylogenetically analyzed by using the maximum-likelihood method. Sequences are listed by their host, country, strain name, and collection year. Scale bars indicate nucleotide substitutions per site.
AP92-like Crimean-Congo Hemorrhagic Fever Virus in Hyalomma aegyptium Ticks, Algeria

Matej Kautman, Ghoulem Tiar, Anna Papa, Pavel Široký

Affiliation information: University of Veterinary and Pharmaceutical Sciences Brno, Brno, Czech Republic (M. Kautman, P. Široký); University Chadli Bendjedid, El Tarf, Algeria (G. Tiar); Aristotle University of Thessaloniki, Thessaloniki, Greece (A. Papa); Central European Institute of Technology, Brno (P. Široký)

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To the Editor: Crimean-Congo hemorrhagic fever virus (CCHFV) (Nairovirus, Bunyaviridae), the causative agent of Crimean-Congo hemorrhagic fever, has been detected in sub-Saharan Africa, southeastern Europe, the Middle East, and central Asia. The virus has been detected in >31 species of ticks and is transmitted to humans by bite of infected ticks (mainly of the genus Hyalomma) or by contact with body fluids or tissue of viremic patients or livestock. The disease is characterized by fever, myalgia, headache, vomiting, and sometimes hemorrhage; reported mortality rate is 10%–50% (1).

CCHFV strains currently constitute 7 evolutionary lineages, 1 of which (Europe 2) contains the prototype strain AP92, which was isolated in 1975 from Rhipicephalus bursa ticks collected from goats in Greece (2). This strain seems to have low or no pathogenicity for humans; only a few mild cases have been reported (3). This observation is supported by the relatively high (14.4%) seroprevalence but no clinical cases in humans in northwestern Greece (4). The documented tick carriers of this strain are R. bursa and Hyalomma marginatum (5).

Hyalomma aegyptium ticks are highly host specific; adults feed almost entirely on tortoises of the genus Testudo (6) and occasionally on hedgehogs and hares. Unlike adult ticks, the larvae and nymphs are less host specific and feed on a wide spectrum of hosts (e.g., other reptiles, birds, and mammals [including humans]) (7). This trait elevates the epidemiologic role of the tick as a possible bridge vector connecting wildlife, domestic animals, and humans.

To determine the biological and epidemiologic role of H. aegyptium ticks, during 2009–2010, we collected 56 adult ticks from 12 Testudo graeca tortoises at a locality near the city of Aflou in Laghouat Province, Algeria. We tested the ticks for probable CCHFV infection by using nested reverse transcription PCR (8), which amplifies a partial fragment of the CCHFV small RNA segment. We slightly modified the assay: reverse transcription time was 60 minutes and annealing temperature was 52°C (9).

In total, 16 (28.6%) ticks were positive for CCHFV. The PCR products of 15 (26.8%) positive samples were sequenced. BLAST (http://blast.ncbi.nlm.nih.gov/Blast.cgi) analysis identified all 15 sequences as CCHFV with 98%–100% identity to the AP92 strain (GenBank accession no. DQ211638). Two variants of AP92 were detected and differed by 0.6%. A phylogenetic tree was constructed by Bayesian inference, using MrBayes version 3.1.2 (http://mrbayes.sourceforge.net/index/php) under a general time-reversible plus gamma distribution plus invariant site model with 107 generations setup (Figure).

Our findings demonstrate the presence of CCHFV in Algeria, either recently introduced or overlooked. The nearest location where CCHFV has been reported is the Zouala region in Morocco, where the virus was detected in H. marginatum tick larvae and nymphs collected from

References

Address for correspondence: Emanuela Foni, Istituto Zoonopatologico Sperimentale della Lombardia ed Emilia Romagna, Via Dei Mercatì 13A, 43126 Parma, Italy; email: emanuela.foni@izsler.it