Recent Findings of Potentially Lethal Salamander Fungus *Batrachochytrium salamandrivorans*

David Lastra González, Vojtech Baláž, Milič Solský, Barbora Thumsová, Krzysztof Kolenda, Anna Najbar, Bartłomiej Najbar, Matej Kautman, Petr Chajma, Monika Balogová, Jiří Vojar

Author affiliations: Czech University of Life Sciences, Prague, Czech Republic (D. Lastra González, M. Solský, B. Thumsová, P. Chajma, J. Vojar); University of Veterinary and Pharmaceutical Sciences, Brno, Czech Republic (V. Baláž, M. Kautman); University of Wrocławski, Wrocław, Poland (K. Kolenda, A. Najbar); University of Zielona Góra, Lubuskie, Poland (B. Najbar); Slovak Academy of Sciences, Košice, Slovakia (M. Kautman); Pavol Jozef Šafárik University in Košice, Košice (M. Balogová)

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The distribution of the chytrid fungus *Batrachochytrium salamandrivorans* continues to expand in Europe. During 2014–2018, we collected 1,135 samples from salamanders and newts in 6 countries in Europe. We identified 5 cases of *B. salamandrivorans* in a wild population in Spain but none in central Europe or the Balkan Peninsula.
Republic) by standard PCR with *B. salamandrivorans*-specific primers STerF and STerR, as described by Martel et al. (2), with subsequent electrophoresis on the amplified target. We reanalyzed samples that produced positive or equivocal results by using duplex quantitative PCR (qPCR) for *B. dendrobatidis* and *B. salamandrivorans* (9) at the University of Veterinary and Pharmaceutical Sciences (Brno, Czech Republic). Trenton Garner of the Institute of Zoology, Zoological Society of London (London, England), provided DNA for quantification standards of the *B. dendrobatidis* GPL lineage, strain IA042, and An Martel of Ghent University (Ghent, Belgium) provided quantification standards of *B. salamandrivorans*.

We directly analyzed samples from other countries by qPCR. We used negative and positive controls for standard PCR analyses and quantification standards for qPCR analyses. For *B. dendrobatidis* or *B. salamandrivorans*-positive sites, we estimated prevalence and Bayesian 95% CIs using 3 parallel Markov chains with 2,000 iterations each, a burn-in of 1,000 iterations, and no thinning (Appendix Table 1). We performed all statistical analyses in R 3.3.1 using the R2WinBUGS package and WinBUGS 1.4.3 (10).

Samples from 5 *L. helveticus* newts tested positive for *B. salamandrivorans*, implying that this species is not resistant to this fungus as previously indicated by experimental exposures (3). The positive cases were found in populations from an isolated area encompassing 2 different regions in northern Spain, Cantabria and Asturias, with remote human populations. Four cases were found in livestock drinking troughs located 150–1,000 m above sea level, and 1 case was found in a pond in a private garden, 30 km from the nearest recorded case. We did not find *B. salamandrivorans*-positive cases in consecutive locations during our monitoring.

Although *B. salamandrivorans* cases have been reported in captive salamanders (6), our reported cases were >1,000 km from any area of known *B. salamandrivorans* occurrence (7). We also detected *B. dendrobatidis* by duplex qPCR in 11 samples from 3 newt species (*L. helveticus*, *L. vulgaris*, and *Triturus cristatus*) from Spain and Montenegro and 1 captive *Cynops ensicauda* newt from the Czech Republic. The *B. dendrobatidis*-positive cases did not involve co-infection with *B. salamandrivorans*.

We confirmed that the known distribution of *B. salamandrivorans* continues to expand in Europe, indicating that this fungus might be capable of dispersing over long distances (4), might be introduced by humans, or might even have been circulating in this geographic range with no detected deaths. Our results should alert the research and conservation community and motivate urgent action to identify regions with early emergence of the disease and implement mitigation measures to prevent further spread of this deadly pathogen.

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**About the Author**

Mr. Lastra González is a PhD candidate at Czech University of Life Sciences, Prague. His research focuses on amphibian conservation and emerging infectious diseases that affect them.

**References**


Address for correspondence: David Lastra González, Faculty of Environmental Sciences, Czech University of Life Sciences, Kamýcká 129, 165 21 Prague–Suchdol, Czech Republic; email: lastra_gonzalez@fzp.czu.cz

## Crimean-Congo Hemorrhagic Fever Virus Genome in Tick from Migratory Bird, Italy

Elisa Mancuso, Luciano Toma, Andrea Polci, Silvio G. d’Alessio, Marco Di Luca, Massimiliano Orsini, Marco Di Domenico, Maurilia Marcacci, Giuseppe Mancini, Fernando Spina, Maria Goffredo, Federica Monaco

Author affiliations: Istituto Zooprofiliattico Sperimentale dell’Abruzzo e del Molise “G. Caporale,” Teramo, Italy (E. Mancuso, A. Polci, S.G. d’Alessio, M. Orsini, M. Di Domenico, M. Marcacci, G. Mancini, M. Goffredo, F. Monaco); Istituto Superiore di Sanità, Rome, Italy (L. Toma, M. Di Luca); Istituto Superiore per la Protezione e la Ricerca Ambientale, Bologna, Italy (F. Spina)

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We detected Crimean-Congo hemorrhagic fever virus in a *Hyalomma rufipes* nymph collected from a whinchat (*Saxicola rubetra*) on the island of Ventotene in April 2017. Partial genome sequences suggest the virus originated in Africa. Detection of the genome of this virus in Italy confirms its potential dispersion through migratory birds.

Crimean-Congo hemorrhagic fever virus (CCHFV) is a vectorborne virus responsible for severe illness in humans, whereas other mammals usually act as asymptomatic reservoirs. The virus is transmitted through tick bites or by direct contact with blood or body fluids of infected vertebrate hosts. CCHFV, an *Orthonairovirus* within the Nairoviridae family, has a negative-sense tripartite RNA genome characterized by high genetic diversity. The sequences of the circulating strains cluster in 6 genotypes (I–VI) reflecting their geographic origin; worldwide distribution is the result of efficient dispersion through migratory birds, human travelers, and the trade and movement of livestock and wildlife (I,2). In Europe, CCHFV distribution was limited to the Balkan region until 2010, when the virus was identified in ticks collected from a red deer (*Cervus elaphus*) and, 6 years later, in 2 autochthonous human cases in the same region of Spain (3). Sequences from the Iberia strains clustered in the Africa genotype III (4), supporting the hypothesis of CCHFV dispersion through ticks hosted by migratory birds.

The role of birds in the potential spread of the virus was confirmed by CCHFV detection in ticks collected from migratory birds in Greece in 2009 (5) and Morocco in 2011 (6). Because Italy hosts an intense passage of birds migrating along major routes connecting winter quarters in Africa and breeding areas in Europe, the country is potentially exposed to the risk for virus introduction. We report the detection of CCHFV RNA in a tick collected in Italy from a migratory bird.

We conducted tick sampling during March–May 2017 on the island of Ventotene, where a ringing station has been operating since 1988 as part of the Small Islands Project, a large-scale and long-term effort to monitor spring migrations of birds across the central and western Mediterranean. We ringed 5,095 birds and checked ≈80% for ectoparasites. We collected 14 adults, 330 nymphs, and 276 larvae from 268 passerines belonging to 28 species; 18 species were trans-Saharan migrants. We stored ticks in 70% ethanol until morphologic identification and assignment to a genus or, whenever possible, a species (7). We then individually

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