Crimean-Congo Hemorrhagic Fever Virus in Ticks from Migratory Birds, Morocco

Ana M. Palomar, Aránzazu Portillo, Paula Santibáñez, David Mazuelas, Juan Arizaga, Ariñe Crespo, Óscar Gutiérrez, Juan Francisco Cuadrado, and José A. Oteo

Crimean-Congo hemorrhagic fever virus was detected in ticks removed from migratory birds in Morocco. This finding demonstrates the circulation of this virus in northwestern Africa and supports the hypothesis that the virus can be introduced into Europe by infected ticks transported from Africa by migratory birds.

Crimean-Congo hemorrhagic fever virus (CCHFV), the causative agent of Crimean-Congo hemorrhagic fever (CCHF), is an arthropod-borne virus (arbovirus) with clinical relevance worldwide (1). CCHF causes sudden onset of signs and symptoms including headache, high fever, back pain, joint pain, stomach pain, and vomiting, which can progress to severe bruising, severe nosebleeds, and uncontrolled bleeding (www.cdc.gov/ncidod/dvrd/spb/mnpages/dispages/cchf.htm).

CCHFV, belonging to the genus Nairovirus, circulates in an enzootic tick-vertebrate-tick cycle in which ticks can act as vectors and reservoirs. CCHFV has been found in ticks of ≥30 species; *Hyalomma marginatum* ticks are considered the most common vectors. Birds are the main hosts for the immature stages of this tick species (2). Viremia does not develop in most passerine birds (3,4), which are not able to pass the virus to ticks. However, migratory species could carry infected ticks over long distances and thereby disseminate the virus (2).

Since the first descriptions of human infections with this virus in 1944–1955 in Crimea, outbreaks of CCHF have been reported in Africa, Asia, and eastern Europe (1). Only imported cases have been reported in western Europe, although the causal agent has been amplified in *H. lusitanicum* ticks collected from deer in Spain (southwestern Europe) (5). This finding could be explained by the arrival of infected ticks transported by migratory birds coming from Africa (5). To confirm this hypothesis, we investigated the presence of CCHFV in ticks collected from migratory birds in northern Africa.

The Study

In April 2011, bird bandings were conducted in Zouala, Morocco (31°47ʹN, 4°14ʹW) (Figure 1). A total of 546 captured birds were checked for ticks, and parasites were found on 21 birds from 5 passerine bird species (*Phoenicurus phoenicurus*, *Erythropygus galactotes*, *Iduna opaca*, *Acrocephalus scirpaceus*, and *I. pallida*). All but *I. pallida* birds are passerine trans-Saharan migrant species, coming from central and southern Africa and able to reach the Iberian Peninsula.

A total of 52 ticks (19 larvae and 33 nymphs) were processed. Genomic DNA and total RNA from ticks were individually purified by using the AllPrep DNA/RNA Mini Kit (QIAGEN, Hilden, Germany), according to the manufacturer’s instructions. DNA extracts were used as templates for 16S rDNA PCR assays (6), and all specimens were classified as *H. marginatum* ticks. RNA was retrotranscribed by using the Omniscript RT kit (QIAGEN), according to the manufacturer’s instructions. cDNA extracts were distributed in 6 pools used as templates for 2 nested PCRs with EcEF and Gre primer pairs (7) (Table). Negative controls (with template DNA but without primers and with primers and containing water instead of template DNA) were included in all assays.

Nested PCR assays using EcEF primers were positive for 4/6 pools; all samples were negative by Gre PCR primers (7). Three of 4 amplicons (associated with *P. phoenicurus*, *E. galactotes*, and *I. opaca* bird species; Figure 2) could be sequenced, and nucleotide sequences were compared with those available in GenBank by using BLAST (www.ncbi.nlm.nih.gov/blast/Blast.cgi). All nucleotide sequences were identical and showed 100% identity with the Sudan AB1-2009 and Mauritania ArD39554 CCHFV strains (GenBank accession nos. HQ378179 and DQ211641) and 98.9% identity with the sequence amplified from ticks from Spain (5).

Conclusions

The detection of CCHFV in ticks from migratory birds in Zouala demonstrates the circulation of this virus...
in Morocco. This country has optimal conditions for the establishment of CCHF, including populations of *H. marginatum* ticks and reservoirs of the virus, such as livestock. Furthermore, autochthonous cases of the disease have been reported in neighboring Mauritania (8).

Our finding of 3 positive tick pools demonstrates the potential dispersion of the virus through infected ticks transported by migratory birds. Several bird species, including *P. phoenicurus*, *E. galactotes*, *I. opaca*, and *A. scirpaceus*, migrate in the spring from southern or central Africa to northern Europe. The Iberian Peninsula may be a stopover or breeding site along those routes (9), which suggests that migratory birds may transport *H. marginatum* ticks from Africa to Europe (2).

Ticks of this species are found in Africa, southern Asia, and southern Europe (10,11); some specimens have been collected in northern European regions such as Germany or England (12,13). Birds are commonly parasitized by immature stages of *H. marginatum*, a 2-host tick that has the same host for larva and nymph stages. Thus, ticks may be attached to the bird for >2–3 weeks (11). An average ground speed of 50 km/h for migratory birds crossing Sahara has been reported (14); this speed would enable birds to cover the distance between Morocco and the Iberian Peninsula in less time than it would take ticks to develop from immature to adult.

In early May 2012, our team captured *A. scirpaceus* birds parasitized by *H. marginatum* ticks in northern Spain (42°48ʹN; 2°39ʹW); some ticks were fully engorged nymphs (A.M. Palomar, unpub. data). A high probability exists that the larvae were attached to birds in Africa and molted and engorged during migration, which supports the possibility of the arrival of migratory birds with CCHFV-infected ticks.

A study conducted in England of CCHFV in ticks collected from migratory birds found negative real-time PCR results for CCHFV, although the parasitization rate of birds with *H. marginatum* ticks was low (13). In addition, a group of experts has reported that migratory birds may not be sufficient to establish new foci of CCHFV infection in Europe and may not represent a high risk for its implantation because adult ticks are necessary and immature specimens cannot find optimal climate conditions to molt in spring (10). However, this report stated that Spain’s average spring temperatures are lower than those needed for birds to molt; in April 2011, the weather station located in northern Spain (La Rioja) (42°27ʹN, 2°19ʹW) recorded average temperatures >14°C for 20/30 days. In addition, *H. marginatum* tick populations are established on the Iberian Peninsula.

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**Table. Distribution in pools of *Hyalomma marginatum* ticks collected from birds in Zouala, Morocco, and PCR results for detection of Crimean-Congo hemorrhagic fever virus**

| Pool | Bird species (no. specimens) | Larvae | Nymphs | Eecf primers | Gre primers |
|------|------------------------------|--------|--------|--------------|-------------|
| A    | *Erythropygia galactotes* (1) |        |        | +            | –           |
|      | *Phoenicurus phoenicurus* (1) | 2 FE   | 6 FE   |              |             |
| B    | *E. galactotes* (2)           | 1 SE   | 7 FE   | +            | –           |
|      | *Iduna opaca* (1)             | 3 FE   |        |              |             |
| C    | *I. opaca* (2)                | 6 FE   | 1 FE   |              |             |
|      | *P. phoenicurus* (1)          |        | 1 SE   |              |             |
| D    | *Acrocephalus scirpaceus* (1) | 4 SE   | 1 FE   | +†           | –           |
|      | *I. opaca* (1)                |        | 1 FE   |              |             |
|      | *I. pallida* (1)              |        | 1 SE, 1 FE |             |             |
| E    | *A. scirpaceus* (1)           | 2 FE   |        |              |             |
|      | *E. galactotes* (1)           | 1 FE   | 2 FE   |             | 1 FE        |
|      | *I. opaca* (4)                |        | 2 SE, 1 FE |             |             |
|      | *P. phoenicurus* (1)          |        | 1 FE   |             |             |
| F    | *E. galactotes* (1)           | 2 SE   | 2 FE   | –            |             |
|      | *I. opaca* (2)                |        | 1 SE, 1 FE |             |             |
|      | *P. phoenicurus* (1)          |        | 1 SE, 1 FE |             |             |

*FE, fully engorged; †, positive; –, negative; SE, semi-engorged.
†No sequence was obtained.
The finding of fragments of the small segment of CCHFV identical to fragments of the Mauritania and Sudan strains and closely related to the sequence previously amplified by our team (5), in pools of immature specimens of different engorged states (Figure 2), may explain 2 hypotheses. First, infected ticks from CCHF-endemic areas, such as Mauritania or Sudan, could have attached to birds and transported to Morocco over them. Second, these ticks could have been carried from Africa to the Iberian Peninsula, thus explaining the circulation of CCHFV in southwestern Europe.

Autochthonous cases of CCHF have not been reported in southwestern Europe, and CCHFV has not been detected in ticks from migratory birds in Europe (13). However, some authors support the limited role of migratory birds harboring CCHFV-infected ticks for the establishment of the virus (10,15). Nevertheless, the finding of CCHFV from migratory birds in Morocco, along with the previous detection of the virus in Spain (5), where H. marginatum tick populations and CCHFV reservoirs occur, call for further study of the distribution of CCHFV in southwestern Europe. Health care workers should be informed about the possibility of CCHF and the clinical picture in the potential disease-endemic areas, and groups potentially at risk for infection (e.g., health care workers, hunters, farmers) and reservoirs for the virus (livestock) should be investigated further.

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Dr Palomar has worked in the Center of Rickettsioses and Arthropod-borne Diseases at the Infectious Diseases Area, Hospital San Pedro–Center for Biomedical Research of La Rioja since March 2009. Her research interests are the taxonomy of ticks and their associated pathogens.

References
1. Ergonul O. Crimean-Congo hemorrhagic fever virus: new outbreaks, new discoveries. Curr Opin Virol. 2012;2:215–20. http://dx.doi.org/10.1016/j.coviro.2012.03.001
2. Hoogstraal H, Kaiser MN, Traylor MA, Gaber S, Guindy E. Ticks (Ixodidea) on birds migrating from Africa to Europe and Asia. Bull World Health Organ. 1961;24:197–212.
3. Shepherd AJ, Swanepoel R, Leman PA, Shepherd SP. Field and laboratory investigation of Crimean-Congo haemorrhagic fever virus (Nairovirus, family Bunyaviridae) infection in birds. Trans R Soc Trop Med Hyg. 1987;81:1004–7. http://dx.doi.org/10.1016/0035-9203(87)90379-8
4. Zeller HG, Cornet JP, Camicas JL. Experimental transmission of Crimean-Congo hemorrhagic fever virus by African wild ground-feeding birds to Hyalomma marginatum rufipes ticks. Am J Trop Med Hyg. 1994;50:676–81.
5. Estrada-Peña A, Palomar AM, Santibáñez P, Sánchez N, Habela MA, Portillo A, et al. Crimean-Congo hemorrhagic fever virus in ticks, southwestern Europe, 2010. Emerg Infect Dis. 2012;18:79–80. http://dx.doi.org/10.3201/eid1801.110104
6. Black WC, Piesman J. Phylogeny of hard and soft tick taxa (Acari:Ixodida) based on mitochondrial 16s rDNA sequences. Proc Natl Acad Sci U S A. 1994;91:10034–8. http://dx.doi.org/10.1073/pnas.91.21.10034
7. Midilli K, Gargili A, Ergonul O, Elevli M, Ergin S, Turan N, et al. The first clinical case due to AP92 like strain of Crimean-Congo hemorrhagic fever virus and a field survey. BMC Infect Dis. 2009;9:90. http://dx.doi.org/10.1186/1471-2334-9-90
8. Nabeth P, Cheikh DO, Lo B, Faye O, Vail JO, Niang M, et al. Crimean-Congo hemorrhagic fever, Mauritania. Emerg Infect Dis. 2004;10:2143–9. http://dx.doi.org/10.3201/eid1012.040455
9. Hoyo J, Elliott A, Christie DA. Handbook of the birds of the world, vol. 10–11. Barcelona (Spain): Lynx Edicions; 2004.
10. European Food Safety Authority Panel on Animal Health and Welfare. Scientific opinion on the role of tick vectors in the epidemiology of Crimean-Congo hemorrhagic fever and African swine fever in Eurasia. EFSA Journal. 2010;8:1703 [cited 2012 Jul 22]. [http://www.efsa.europa.eu/en/efsaJournal/doc/1703.pdf]
11. Estrada-Peña A, Martínez M, Muñoz MJ. A population model to describe the distribution and seasonal dynamics of the tick *Hyalomma marginatum* in the Mediterranean basin. Transbound Emerg Dis. 2011;58:213–23. http://dx.doi.org/10.1111/j.1865-1682.2010.01198.x

12. Rumer L, Graser E, Hillebrand T, Talaska T, Dautel H, Medianikov O, et al. *Rickettsia aeschlimannii* in *Hyalomma marginatum* ticks, Germany. Emerg Infect Dis. 2011;17:325–6. http://dx.doi.org/10.3201/eid1702.100308

13. Jameson LJ, Morgan PJ, Medlock JM, Watola G, Vaux AG. Importation of *Hyalomma marginatum*, vector of Crimean-Congo haemorrhagic fever virus, into the United Kingdom by migratory birds.Ticks Tick Borne Dis. 2012;3:95–9. http://dx.doi.org/10.1016/j.ttbdis.2011.12.002

14. Schmaljohann H, Liechti F, Bruderer B. Songbird migration across the Sahara: the non-stop hypothesis rejected! Proc Biol Sci. 2007;274:735–9. http://dx.doi.org/10.1098/rspb.2006.0011

15. Gale P, Stephenson B, Brouwer A, Martinez M, de la Torre A, Bosch J, et al. Impact of climate change on risk of incursion of Crimean-Congo haemorrhagic fever virus in livestock in Europe through migratory birds. J Appl Microbiol. 2012;112:246–57. http://dx.doi.org/10.1111/j.1365-2672.2011.05203.x

Address for correspondence: José A. Oteo, Área de Enfermedades Infecciosas, Hospital San Pedro-CIBIR, C/Piqueras 98-7ª NE, 26006 Logroño (La Rioja), Spain; email: jaoteo@riojasalud.es

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**etymologia**

**Quarantine [kworˈən-tən]**

From the Italian quaranta (forty), “quarantine” refers to the practice established in European port cities during the Black Death requiring vessels to lie at anchor for 40 days before landing. Isolation (from the Latin insula or island), the practice of separating sick persons from those who are healthy to prevent spread of disease, goes back a long time. “As long as they have the disease they remain unclean. They must live alone; they must live outside the camp” (Leviticus 13:46).

Quarantine, on the other hand, is the practice of separating persons who appear to be healthy but may have been exposed to a disease. In 1377, the Great Council of Ragusa (modern-day Dubrovnik) established a 30-day separation period (trentino) for visitors from plague-endemic areas. In the following decades, the practice spread to other cities and was extended from 30 to 40 days (quarantino). The longer period may have been more effective at preventing disease or just a nod to the 40-day duration of Biblical events—the Great Flood or Jesus’ fast in the wilderness.

**Sources**

1. Dorland’s Illustrated Medical Dictionary. 32nd ed. Philadelphia: Elsevier Saunders; 2012.
2. Holy Bible, New International Version. Colorado Springs (CO): Biblica; 2011.
3. Sehdev PS. The origin of quarantine. Clin Infect Dis. 2002;35:1071–2. PubMed http://dx.doi.org/10.1086/344062

Address for correspondence: Ronnie Henry, Centers for Disease Control and Prevention, 1600 Clifton Rd NE, Mailstop E03, Atlanta, GA 30333, USA; email: boq3@cdc.gov

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