Rickettsia aeschlimannii in Hyalomma marginatum Ticks, Germany

To the Editor: Rickettsia spp. of the spotted fever group cause world-wide emerging human infections known as tick-borne rickettsioses (1). Data on the occurrence and prevalence of Rickettsia in Germany are still limited (2). Six Rickettsia species have been reported to date (2). R. helvetica, R. felis, R. massiliae, and R. monacensis were detected with a relatively low prevalence in Ixodes ricinus ticks collected in southern Germany (2); R. raoultii was identified with high prevalence in the rapidly expanding areas where D. reticulatus ticks are found (2). R. raoultii was recently recognized as an agent of tick-borne lymphadenopathy/Dermacentor-borne necrosis and erythema lymphadenopathy (3). Low prevalence of another tick-borne lymphadenopathy agent, R. slovaca, in Dermacentor marginatus ticks collected in southern Germany was recently reported (4).

We report the detection in Germany of the pathogenic SFG species R. aeschlimannii (1), which is phylogenetically close to R. raoultii and causes an infection with clinical signs similar to those of Mediterranean spotted fever (1). To determine the prevalence of R. raoultii in the ticks in Berlin/Brandenburg and neighboring regions, we collected 294 ticks; 288 had been collected either from vegetation or domestic animals and morphologically identified as adult D. reticulatus ticks. The remaining 6 ticks were delivered by an ornithologist who had removed them from a bird (belonging to the genus Acrocephalus sp.) that he had captured in the reeds near Pakendorf and Zerbst, Saxony-Anhalt, in May 2007. These 6 ticks were reported as D. reticulatus–like adults but were damaged in the collection process, making an exact morphologic evaluation impossible.

DNA was isolated from the complete tick body by homogenization in the SpeedMill (Analytik Jena Biosolutions, Jena, Germany) followed by purification by RapideStripe tick DNA/RNA Extraction Kit (Analytik Jena Biosolutions). Multispacer typing (5) was used for molecular detection and determination of Rickettsia spp. (Figure). DNA sequencing and analysis were performed as described (Figure).

In 51.3% of the intact tick isolates, R. raoultii was detected. In each of the 3 damaged isolates, the multispacer type pattern was detected, which was 100% identical to that of R. aeschlimannii (5) (Figure). Moreover, PCR, which we routinely use as a positive control for molecular identification of D. reticulatus, yielded no product in the damaged isolates (Figure).

To determine the species of the damaged ticks, we used 3 tick-specific PCRs. One amplified a 16S rRNA fragment used for phylogenetic studies of ticks (6). Use of the other 2 PCRs was

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based on the consideration that \textit{R. aeschlimannii} is usually found in ticks of the genus \textit{Hyalomma}, primarily in \textit{H. marginatum} (1). Therefore, 1 of the PCRs amplified a fragment of the \textit{Hyalomma} tick mitochondrial cytochrome oxidase I gene and the other a fragment of the internal transcribed spacer 2 (7).

The ITS2 fragment displayed the highest (99%) similarity with the respective fragment of \textit{H. marginatum}, \textit{H. dromedarii}, \textit{H. truncatum}, and \textit{H. lusitanicum}. Cytochrome oxidase subunit I fragment was 99% identical to \textit{H. marginatum}, \textit{H. dromedarii}, and \textit{H. truncatum}. The 16S RNA fragment was 98% identical to \textit{H. marginatum}; its identity to the second closest sequence belonging to \textit{H. lusitanicum} was 96%.

Earlier, \textit{R. aeschlimannii} had been detected in sub-Saharan and North Africa, southern Europe, and southwestern Russia (8). Therefore, the area of Zerbst, the middle of Germany, marks the northernmost point of \textit{R. aeschlimannii} detection. \textit{Hyalomma} spp. ticks are distributed in Africa, the Mediterranean climatic zone of southern Europe, and in Asia. The only documented \textit{Hyalomma} spp. tick in Germany was found on a human in the southern part of the country (Lake Constance area) in May 2006, but the possibility of tick transportation from Spain was not ruled out (9).

\textit{Acrocephalus scirpaceus} birds are migratory birds and live in central Europe from April to October and winter in sub-Saharan Africa in the region inhabited by \textit{Hyalomma} spp. ticks. Therefore, it is reasonable to suggest that the \textit{Hyalomma} spp. ticks that we examined had been transported by the birds from Africa. The fact that a randomly caught bird was infested with \textit{R. aeschlimannii}–infected ticks is suggestive of the intensive stream of new pathogens transported through Europe by migrating birds. The first possible implication of a bird as a vector of infected ticks was proposed for \textit{Hyalomma} spp. ticks infected by \textit{R. sibirica mongolitimonae} (10). Until now, the role of migrating birds in distribution of tick-borne pathogens has been poorly understood (9). The changing climate and environment in central Europe may facilitate the establishment of pathogen-carrying tick species transported by birds. These new pathogens can be directly transmitted from infected birds to the species of the local fauna.

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\textbf{Dogs as Reservoirs for \textit{Leishmania braziliensis}}

To the Editor: I have read the review by Sousa and Pearson (1), which provides a fascinating historical account of the Great Drought and the smallpox epidemic of the 1870s and their association with the emergence of cutaneous leishmaniasis in Ceará, Brazil. In their review, the authors went back to the 19th century, remembering the hard years experienced by those who faced the Great Drought,