LETTERS

humans. Additionally, Mycobacteria spp. can occasionally cause disease in humans through contact with fish (M. marinum), and pedicure treatments have previously been associated with M. fortuitum infections (10).

Recently, the risks associated with exposure to G. rufa fish were reported to be low (1). To date, there are only a limited number of reports of patients who might have been infected by this exposure route (1). However, our study raises some concerns over the extent that these fish, or their transport water, might harbor potential zoonotic disease pathogens of clinical relevance. In particular, patients with underlying conditions (such as diabetes mellitus immunosuppression) should be discouraged from undertaking such treatments, especially if they have obvious breaks in the skin or abrasions. This risk can probably be reduced by use of certified diseasefree fish reared in controlled facilities under high standards of husbandry and welfare.

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References

- 1. Health Protection Agency Fish Spa Working Group. Guidance on the management of the public health risks from fish pedicures: draft for consultation. 2011 Aug 31 [cited 2012 Mar 21]. http:// www.hpa.org.uk/webc/HPAwebFile/ HPAweb C/1317131045549
- Jones N, Bohnsack JF, Takahashi S, Oliver KA, Chan MS, Kunst F, et al. Multilocus sequence typing system for group B Streptococcus. J Clin Micro-2003;41:2530-6. http://dx.doi. org/10.1128/JCM.41.6.2530-2536.2003
- Evans JJ, Bohnsack JF, Klesius PH, Whiting AA, Garcia JC, Shoemaker CA, et al. Phylogenetic relationships among Streptococcus agalactiae isolated from piscine, dolphin, bovine and human sources: a dolphin and piscine lineage associated with a fish epidemic in Kuwait is also associated with human neonatal infections in Japan. J Med Microbiol. 2008:57:1369-76. http:// dx.doi.org/10.1099/jmm.0.47815-0
- Janda JM, Abbott SL. The genus Aeromonas: taxonomy, pathogenicity, and infection. Clin Microbiol Rev. 2010;23:35-73. http://dx.doi.org/10.1128/CMR.00039-09
- Jones MK. Oliver, JD. Vibrio vulnificus: disease and pathogenesis. Infect Immun. 2009;77:1723-33. http://dx.doi. org/10.1128/IAI.01046-08
- Morris JG. Non-O group-1 Vibrio cholera-a look at the epidemiology of an occasional pathogen. Epidemiol Rev. 1990;12:179-91.
- Wagner D, Young LS. Nontuberculous mycobacterial infections: a clinical review. Infection. 2004;32:257-70. http:// dx.doi.org/10.1007/s15010-004-4001-4
- Skoff TH, Farley MM, Petit S, Craig AS, Schaffner W, Gershman K, et al. Increasing burden of invasive group B streptococcal disease in non-pregnant adults, 1990-2007. Clin Infect Dis. 2009;49:85-92. http://dx.doi.org/10.1086/599369
- Verner-Jeffreys DW, Welch TJ, Schwarz T. Pond MJ. Woodward MJ. Haig SJ. et al. High prevalence of multidrug-tolerant bacteria and associated antimicrobial resistance genes isolated from ornamental fish and their carriage water. PLoS ONE. 2009;4:e8388. http://dx.doi.org/10.1371/ journal.pone.0008388
- De Groote MA, Huitt G. Infections due to rapidly growing mycobacteria. Clin Infect Dis. 2006;42:1756-63. http://dx.doi. org/10.1086/504381

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Rickettsia conorii **Indian Tick Typhus** Strain and R. slovaca in **Humans, Sicily**

To the Editor: Rickettsiae are vector-borne pathogens that affect humans and animals worldwide (1). Pathogens in the Rickettsia conorii complex are known to cause Mediterranean spotted fever (MSF) (R. conorii Malish strain), Astrakhan fever (R. conorii Astrakhan strain), Israeli spotted fever (R. conorii Israeli spotted fever strain), and Indian tick typhus (R. conorii Indian tick typhus strain) in the Mediterranean basin and Africa. southern Russia, the Middle East, and India and Pakistan, respectively (2). These rickettsioses share some clinical features, such as febrile illness generalized cutaneous rash, and are transmitted to humans by Rhipicephalus spp. ticks (2).

MSF is endemic to Sicily (Italy); fatal cases occur each year, and the prevalence of R. conorii in dogs is high (3-6). Recently, R. conorii Malish strain and R. conorii Israeli spotted fever strain were confirmed in humans in Sicily in whom MSF was diagnosed (4), which suggests that other R. conorii strains might be present and diagnosed as causing MSF. The rickettsiae within the R. conorii complex, which are relevant for the study of bacterial evolution and epidemiology, can be properly identified only by appropriate genetic analyses.

We analyzed 15 blood and 19 inoculation eschar samples collected during 2005-2009 from 31 patients in Palermo Province and 2 in Catania Province, none of whom had recently traveled. None were severely ill, but all 33 had clinical manifestations and laboratory results compatible with MSF: 1-week incubation after tick bite, fever, headache, myalgia, papulonodular rash that on the upper limbs and spread centripetally with or without tache noire, and detection of antibody titers >180 to R. conorii by indirect immunofluorescence antibody test (bioMérieux, Marcy L'Etoile, France).

Total DNA was extracted by using the GeneElute Mammalian Genomic DNA Miniprep Kit (Sigma-Aldrich, Milan, Italy) and used to analyze Rickettsia spp. sequences by PCR, cloning, and sequence analysis of the amplicons. At least 3 clones were sequenced for each amplicon. Genes targeted by PCR included ATP synthase α subunit (atpA) (7), heatshock protein 70 (dnaK) (7), outer membrane protein A (*ompA*) (primers Rr190.70p and 190-701 [8]), outer membrane protein B (*ompB*) (primers rompBSFGIF and rompBSFG/TGIR [9]), citrate synthase (gltA) (2), and 17-kDa protein (primers TZ15-19 TZ16–20 [6]). Nucleotide sequence identity to reference strains (2), multilocus analysis by atpAdnaK-ompA-ompB-gltA-17-kDa and ompA-ompB sequences and in silico PstI-RsaI restriction analysis of ompA sequences (8) were used to characterize *Rickettsia* spp. and *R*. conorii strains.

Results for 15 (45%) patients were positive for *Rickettsia* spp. Thirteen isolates were confirmed as *R. conorii* Malish strain (identification [ID] nos. 44, 45, 47, 49, 54, 55, 57, 59, 61, 66, 68, 92, 112) and 1 each as *R. conorii* Indian tick typhus strain (ID no. 58) and *R. slovaca* (ID no. 50). *R. slovaca* DNA was also found in a *Dermacentor marginatus* tick removed from the

patient who had confirmed *R. slovaca* infection. *R. conorii* Malish strains showed 99.9%–100%, 100%, 100%, 98.7%–100%, 100%, and 97.8%–100% pairwise nt sequence identity to reference strain Malish 7 (AE006914) *atpA*, *dnaK*, *ompA*, *ompB*, *gltA*, and 17-kDa protein, respectively.

The *R. conorii* Indian tick typhus strain showed 100%, 100%, 99.4%, 100%, 100%, and 99.9% pairwise nt sequence identity to *R. conorii* strain Malish 7 (AE006914) *atpA*, *dnaK*, 17-kDa protein, and *R. conorii* Indian tick typhus reference strain *ompA* (U43794), *ompB* (AF123726), and

gltA (U59730), respectively. The R. slovaca strain showed 99.4%, 97.8%, 100%, 93.7%, 99.7%, and 99.4% pairwise nt sequence identity to R. slovaca atpA (AY124734), dnaK (DQ821824), ompA (HM149286), ompB (HQ232242), gltA (AY129301), and R. conorii strain Malish 7 (AE006914) 17-kDa protein, respectively. The sequences were deposited in GeneBank under accession nos. JN182782–JN182804.

Multilocus sequence analysis (Figure, panel A) and in silico *PstI-RsaI* restriction analysis of *ompA* sequences also confirmed the identity

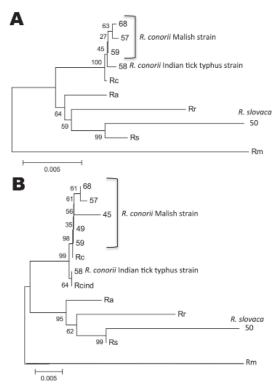


Figure. Multilocus sequence analysis of Rickettsia spp. Evolutionary history was inferred by using the neighbor-joining method for ATP synthase α subunit (atpA)-heat shock protein 70 (dnaK)-outer membrane protein A (ompA)-ompB-citrate synthase (gltA)-17-kDa (A) and ompA-ompB sequences (B). The optimal tree with the sum of branch length = 0.06205323 (A) and 0.11097561 (B) is shown. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1,000 replicates) are shown next to the branches. The tree is drawn to scale, with branch lengths in the same units as those of the evolutionary distances used to infer the phylogenetic relationship. Evolutionary distances were computed by using the Kimura 2-parameter method and are in the units of the number of base substitutions per site. All ambiguous positions were removed for each sequence pair. Evolutionary analyses were conducted in MEGA5 (www. megasoftware.net). Identification numbers of strains detected are shown with the species/ strain name next to them. Rc, R. conorii strain Malish 7; Ra, R. africae strain ESF-5; Rr, R. rickettsii strain lowa; Rs, R. slovaca; Rm, R. massiliae strain MTU5; Rcind, R. conorii Indian tick typhus strain.

of the *Rickettsia* spp. we identified. As shown (2), multilocus analysis with *ompA-ompB* sequences was highly informative about the phylogenetic relationship between *Rickettsia* spp. and *R. conorii* strains (Figure, panel B).

In Sicily, R. conorii Malish strain has been characterized in MSF patients (4), and R. slovaca DNA was identified in ixodid ticks (5). However, to our knowledge, R. slovaca in humans in Sicily and R. conorii Indian tick typhus strain infection in Sicily and Europe have not been reported. The only previous report outside India and Pakistan was documented in a traveler with severe clinical manifestations in France (10). Differences were not observed between R. conorii Indian tick typhus strain and R. slovacainfected patients. Both patients had similar clinical symptoms compatible with MSF; in both, only IgM for rickettsiae was detected at hospital admission, but IgM and IgG were detected during convalescence. Tache noire were detected in the neck and right arm of patients with R. conorii Indian tick typhus strain and R. slovaca, respectively.

These results demonstrated that new rickettsiae, such as *R. conorii* Indian tick typhus strain, of public health relevance are emerging in Europe. The widespread distribution of tick vectors in Europe and the transtadial and transovarial transmission of the pathogen in ticks might favor transmission to humans.

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References

- Nicholson WL, Allen KE, McQuiston JH, Breitschwerdt EB, Little SE. The increasing recognition of rickettsial pathogens in dogs and people. Trends Parasitol. 2010;26:205–12. http://dx.doi. org/10.1016/j.pt.2010.01.007
- Zhu Y, Fournier PE, Eremeeva M, Raoult D. Proposal to create subspecies of *Rickettsia conorii* based on multilocus sequence typing and an emended description of *Rickettsia conorii*. BMC Microbiol. 2005;5:11. http://dx.doi. org/10.1186/1471-2180-5-11
- Ciceroni L, Pinto A, Ciarrocchi S, Ciervo A. Current knowledge of rickettsial diseases in Italy. Ann N Y Acad Sci. 2006;1078:143–9. http://dx.doi. org/10.1196/annals.1374.024
- Giammanco GM, Vitale G, Mansueto S, Capra G, Caleca MP, Ammatuna P. Presence of *Rickettsia conorii* subsp. *israelensis*, the causative agent of Israeli spotted fever, in Sicily, Italy, ascertained in a retrospective study. J Clin Microbiol. 2005;43:6027–31. http://dx.doi.org/10.1128/JCM.43.12.6027-6031.2005
- Beninati T, Genchi C, Torina A, Caracappa S, Bandi C, Lo N. Rickettsiae in ixodid ticks, Sicily. Emerg Infect Dis. 2005;11:509–11. http://dx.doi.org/10.3201/eid1103.040812
- Tzianabos T, Anderson BE, McDade JE.
 Detection of *Rickettsia rickettsii* DNA in clinical specimens by using polymerase chain reaction technology. J Clin Microbiol. 1989;27:2866–8.
- Fernández de Mera IG, Zivkovic Z, Bolaños M, Carranza C, Pérez-Arellano JL, Gutiérrez C, et al. *Rickettsia massiliae* in the Canary Islands. Emerg Infect Dis. 2009;15:1869–70.

- Roux V, Fournier PE, Raoult D. Differentiation of spotted fever group rickettsiae by sequencing and analysis of restriction fragment length polymorphism of PCRamplified DNA of the gene encoding the protein rOmpA. J Clin Microbiol. 1996;34:2058–65.
- Choi YJ, Jang WJ, Ryu JS, Lee SH, Park KH, Paik HS, et al. Spotted fever group and typhus group rickettsioses in humans, South Korea. Emerg Infect Dis. 2005;11:237–44. http://dx.doi. org/10.3201/eid1102.040603
- Parola P, Fenollar F, Badiaga S, Brouqui P, Raoult D. First documentation of *Rick-ettsia conorii* infection (strain Indian tick typhus) in a traveler. Emerg Infect Dis. 2001;7:909–10. http://dx.doi.org/10.3201/eid0705.010527

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Detection of European Strain of Echinococcus multilocularis in North America

To the Editor: In 2009, an alveolar hydatid cyst, the intermediate stage of the cestode Echinococcus multilocularis, was detected in the liver of a dog from Quesnel, British Columbia (BC), Canada (1), 600 km west of the nearest known record of this parasite in central North America (Figure). Alveolar hydatid cysts normally occur in rodent intermediate hosts. However, humans can serve as aberrant intermediate hosts; cysts generally originate in the liver and, in about one third of cases, metastasize throughout the body (2). Detection of the larval stage of this pathogen in an unusual host in a new geographic region required application of multiple molecular epidemi-