LETTERS

were typical of lymphangitis-associated rickettsiosis, and most cases of rickettsioses in southern France in the spring are caused by R. sibirica mongolitimonae. Clustered cases of SFG rickettsiae infection have been reported in Europe, including southern France (3,6). In 2007, R. conorii and R. massiliae infections in humans were reported (3). In 2010, cases for which we were unable to discriminate between R. conorii and R. massiliae infections occurred in a family (6). In these 2 studies, clustered cases of SFG rickettsiosis involved Rh. sanguineus ticks. Clustered cases appeared to be related to an increase in aggressiveness of ticks toward humans during warmer periods (3). In our study, no correlation was identified with warmer weather.

R. sibirica mongolitimonae is most frequently associated with Hya*lomma* spp. ticks (1,2,4). However, 1 case of infection with this bacterium was associated with Rh. pusillus ticks collected in Portugal (7); DNA from this bacteria was also identified in an Rh. pusillus tick collected from a mongoose. The European wild rabbit is the primary host of Rh. pusillus ticks. However, these ticks have been found on wild carnivorous animals, dogs, and domestic cats (8); these ticks can bite humans (8). Moreover, R. massiliae and R. sibirica mongolitimonae were found in Rh. pusillus ticks from Spain (9), and SFG rickettsiae were found in ticks from Sardinia (10). Therefore, Rh. pusillus ticks appear to be an emerging vector for R. sibirica mongolitimonae in Europe.

Sophie Edouard, Philippe Parola, Cristina Socolovschi, Bernard Davoust, Bernard La Scola, and Didier Raoult

Author affiliation: Aix-Marseille Université, Marseille, France

DOI: http://dx.doi.org/eid1902.120863

References

- Fournier PE, Gouriet F, Brouqui P, Lucht F, Raoult D. Lymphangitis-associated rickettsiosis, a new rickettsiosis caused by *Rickettsia sibirica mongolotimonae*: seven new cases and review of the literature. Clin Infect Dis. 2005;40:1435–44. http:// dx.doi.org/10.1086/429625
- Socolovschi C, Barbarot S, Lefebvre M, Parola P, Raoult D. *Rickettsia sibirica* mongolitimonae in traveler from Egypt. Emerg Infect Dis. 2010;16:1495–6. http:// dx.doi.org/10.3201/eid1609.100258
- Parola P, Socolovschi C, Jeanjean L, Bitam I, Fournier PE, Sotto A, et al. Warmer weather linked to tick attack and emergence of severe rickettsioses. PLoS Negl Trop Dis. 2008;2:e338. http://dx.doi. org/10.1371/journal.pntd.0000338
- Mediannikov O, Diatta G, Fenollar F, Sokhna C, Trape JF, Raoult D. Tickborne rickettsioses, neglected emerging diseases in rural Senegal. PLoS Negl Trop Dis. 2010;4:pii:e821 . http://dx.doi. org/10.1371/journal.pntd.0000821
- Renvoisé A, Rolain JM, Socolovschi C, Raoult D. Widespread use of real-time PCR for rickettsial diagnosis. FEMS Immunol Med Microbiol. 2012;64:126–9. http://dx.doi.org/10.1111/j.1574-695X.20 11.00899.x
- Renvoisé A, Delaunay P, Blanchouin E, Cannavo I, Cua C, Socolovschi C, et al. Urban family cluster of spotted fever rickettsiosis linked to *Rhipicephalus sanguineus* infected with *Rickettsia conorii* subsp. *caspia* and *Rickettsia massiliae*. Ticks Tick Borne Dis 2012; 3:389–92.http:// dx.doi.org/10.1016/j.ttbdis.2012.10.008.
- de Sousa R, Barata C, Vitorino L, Santos-Silva M, Carrapato C, Torgal J, et al. *Rickettsia sibirica* isolation from a patient and detection in ticks, Portugal. Emerg Infect Dis. 2006;12:1103–8. http://dx.doi. org/10.3201/eid1207.051494
- Santos-Silva MM, Beati L, Santos AS, de Sousa R, Nuncio MS, Melo P, et al. The hard-tick fauna of mainland Portugal (Acari: Ixodidae): an update on geographical distribution and known associations with hosts and pathogens. Exp Appl Acarol. 2011;55:85–121. http://dx.doi. org/10.1007/s10493-011-9440-x
- Toledo A, Olmeda AS, Escudero R, Jado I, Valcarcel F, Casado-Nistal MA, et al. Tick-borne zoonotic bacteria in ticks collected from central Spain. Am J Trop Med Hyg. 2009;81:67–74.
- Satta G, Chisu V, Cabras P, Fois F, Masala G. Pathogens and symbionts in ticks: a survey on tick species distribution and presence of tick-transmitted micro-organisms in Sardinia, Italy. J Med Microbiol. 2011;60:63–8. http://dx.doi.org/10.1099/ jmm.0.021543-0

Address for correspondence: Didier Raoult, Unité des Rickettsies, Unité de Recherche sur les Maladies Infectieuses et Tropicales Émergentes, Centre National de la Recherche Scientifique–Institut de Recherche pour le Développement, Unités Mixtes de Recherché 6236, Faculté de Médecine et de Pharmacie, 27 Blvd Jean Moulin 13385 Marseille Cedex 05, France; email: didier.raoult@gmail.com

Rickettsiae in Ticks, Japan, 2007–2011

To the Editor: Japanese spotted fever (JSF), caused by Rickettsia japonica, is the most prevalent tickborne infectious disease in Japan (1), occurring most frequently in central and western regions (http://idsc.nih. go.jp/idwr/CDROM/Main.html Γin Japanese]). Cases of unknown fever with rickettsiosis-like symptoms not associated with JSF have been reported in JSF-endemic regions of Japan (2). Several spotted fever group (SFG) rickettsiae (R. japonica, R. heilongjiangensis, R. helvetica, R. tamurae, R. asiatica, Candidatus R. tarasevichiae) and other related *Rickettsia* spp. have been identified in Japan (1,3-6). Human infections with R. heilongjiangensis and R. tamurae have been confirmed (3,5), and Anaplasma phagocytophilum and Ehrlichia chaffeensis, known human pathogens, have been detected in ticks and deer in Japan. We conducted this study to determine the risk in central and western Japan for human exposure to ticks harboring SFG rickettsiae, A. phagocytophilum, or Ehrlichia spp.

In 2007–2011, we collected 827 *Haemaphysalis*, *Amblyomma*, and *Ixodes* spp. ticks (392 adults, 435 nymphs) by flagging vegetation in the prefectures of Shizuoka, Mie, Wakayama, Kagoshima, Nagasaki (Goto Island), and Okinawa (the main island and Yonaguni Island) (Technical

LETTERS

Appendix Figure 1, wwwnc.cdc.gov/ EID/article1/12-0856-Techapp/.pdf). We extracted DNA from the salivary glands of each tick and performed PCR to amplify *gltA*, 16S rDNA, and *ompA* of SFG rickettsiae. To detect *A. phagocytophilum* and *Ehrlichia* spp., we performed nested PCR targeting the *p44/msp2* and *p28/omp-1* multigenes, respectively.

PCR gltA screening revealed SFG rickettsiae in 181 (21.9%) of the 827 ticks (Table). We obtained nearly full-length (1.1-kb) gltA sequences and classified them into 5 groups by phylogenetic analyses (Technical Appendix Figure 2). Sequences for groups 1 (prevalence 1.0%) and 2 (prevalence 3.2%) were identified as R. japonica YH (GenBank accession no. AP011533) and R. tamurae (GenBank accession no. AF394896), respectively (Table). Group 3 (prevalence 15.1%) sequences were identical to that of *Rickettsia* sp. LON (GenBank accession no. AB516964). The sequence for group 4 (prevalence 1.6%) was closely related to that for R. raoultii strain Khabarovsk (98.8% similarity), and a part of the sequence (342 bp) was identical to that of Rickettsia sp. Hf 151 (GenBank accession no. AB114815). Group 5 consisted of 4 newly identified rickettsiae (Technical Appendix Figure 2). Of these 4 rickettsiae, 3 (Mie311, Goto13, and Mie334) were closely related to *R. raoultii* strain Khabarovsk (98.0% identity) and 1 (Mie201) was similar to *Candidatus* R. principis (99.7% identity).

We further analyzed the 16S rDNA and *ompA* in *gltA*-positive tick samples. The 16S rDNA and *ompA* for group 1 samples shared 100% identity with 16S rDNA and *ompA* of *R. japonica* YH (AP011533). The 16S rDNA of group 2 was identical to that of *R. tamurae* (AY049981). In groups 3–5, some of the specific amplicons in 16S rDNA or *ompA* could be detected; their sequences were confirmed to be similar (but not identical) to those of several known rickettsial sequences.

We amplified the p44/msp2 amplicons of *A. phagocytophilum* from 25 (3%) of 827 ticks (Table). By cloning (TA Cloning Kit; Life Technologies, Carlsbad, CA, USA) and sequencing these amplicons, we obtained and identified 60 new TA-clone sequences (366–507 bp) for p44/msp2 (GenBank accession nos. JQ697880-JQ697950); these sequences may include a potentially novel Anaplasma species. (7). Ehrlichia p28/omp-1 was detected from 2 (0.2%) of the 827 ticks. Of 5 TA-clone sequences (284-315 bp) obtained from the 2 ticks, 2 from an A. testudinarium tick (GenBank accession nos. JQ697886 and JQ697887) shared 83.3%-86.7% similarity with E. ruminantium Gardel Map-1 (Gen-Bank accession no. YP196842), and 3 from an H. longicornis tick (GenBank accession nos. JQ697888-JQ697890) showed the closest relationship to E. ewingii omp-1-15 (67%-73% similarity: GenBank accession no. EF116932).

We identified the tick species associated with *R. japonica* as *H. formo*sensis, *H. hystricis*, and *H. cornigera*, and another study reported an association with *Dermacentor taiwanensis*, *H. flava*, *H. longicornis*, and *I. ovatus* (4). In our study and previous studies, the tick species associated with *A. phagocytophilum* in Japan were identified as *H. formosensis*, *H. lon*gicornis, *H. megaspinosa*, *A. testudi*narium, *I. ovatus*, and *I. persulcatus* (8). Thus, it appears that 3 tick species (*H. formosensis*, *H. longicornis*, and *I.*

Table. PCR survey results for *Haemaphysalis*, *Amblyomma*, and *Ixodes* spp. ticks tested for rickettsiae, central and western Japan, 2007–2011*

	No. ticks	Total no. (%) ticks	No. (%) ticks positive for						
			Rickettsia gltA, by species group†					A. phagocytophilum	Ehrlichia
Tick species	tested	positive	Group 1	Group 2	Group 3	Group 4	Group 5	p44/msp2	p28/omp-1§
H. formosensis	224	6 (2.7)	1 (0.4)	0	0	0	5 (2.2)	18 (8)	0
H. hystricis	97	19 (19.6)	6 (6.1)	0	0	13 (13.4)	0	0	0
H. longicornis	294	119 (40.5)	0	0	119 (40.5)	0	0	2 (0.7)	1 (0.4)‡
H. flava	55	6 (10.9)	0	0	2 (3.6)	0	4 (7.3)	0	0
H. kitaokai	10	0	0	0	0	0	0	0	0
H. megaspinosa	18	4 (22.2)	0	0	4 (22.2)	0	0	1 (5.6)	0
H. cornigera	11	1 (9.1)	1 (9.1)	0	Ò	0	0	О́	0
A. testudinarium	112	26 (23.2)	0	26 (23.2)	0	0	0	3 (2.7)	1 (0.9)
A. geoemydae	1	0	0	0	0	0	0	0	0
I. ovatus	5	0	0	0	0	0	0	1 (20.0)	0
Total	827	181 (21.9)	8 (1.0)	26 (3.1)	125 (15.1)	13 (1.6)	9 (1.1)	25 (3.0)	2 (0.2)

*DNA was extracted from the salivary glands of each tick by using the DNeasy Mini Kit (QIAGEN Sciences, Germantown, MD, USA) and used as a template for PCR. The newly identified sequences of *gltA*, 16S rDNA, *ompA*, *p44/msp2*, and *p28/omp*-1 in this study were deposited into GenBank under accession nos. JQ697880–JQ697959. A. phagocytophilum, Anaplasma phagocytophilum.

†The PCR primers used, gltA–Fc (5'-CGAACTTACCGCTATTAGAATG-3') and gltA–Rc (5'-CTTTAGAGGCGATAGCTTCAAG-3'), were designed in this study. Groups: 1, Rickettsia japonica YH (GenBank accession no. AP011533); 2, R. tamurae (GenBank accession no. AF394896); 3, Rickettsia sp. LON-13 (GenBank accession no. AB516964); 4, Rickettsia sp. Hf151; 5, other rickettsiae.

‡PCR primers of p3726 (5'-GCTAAGGAGTTAGCTTATGA-3'), p3761 (5'-CTGCTCT[T/G]GCCAA(AG)ACCTC-3', p4183 (5'-

CAATAGT[C/T]TTAGCTAGTAACC-3'), and p4257 (5'-AGAAGATCATAACAAGCATTG-3') were used for detection of p44/msp2.

§PCR primers conP28-F1 (5'-AT[C/T]AGTG[G/C]AAA[A/G]TA[T/C][A/G]T[G/A]CCAA-3'), conP28-F2 (5'-

CAATGG[A/G][T/A]GG[T/C]CC[A/C]AGA[A/G]TAG-3'), conP28-R1 (5'-TTA[G/A]AA[A/G]G[C/T]AAA[C/T]CT[T/G]CCTCC-3'), and conP28-R2 (5'-TTCC[T/C]TG[A/G]TA[A/G]G[A/C]AA[T/G]TTTAGG-3') were used to detect p28/omp-1.

LETTERS

ovatus) are associated with *R. japonica* and *A. phagocytophilum*.

In addition, in an H. formosensis tick, we detected an SFG rickettsia that is closely related to R. raoultii, the etiologic agent of Dermacentorborne necrosis erythema and lymphadenopathy in Europe and Russia (9). We detected Candidatus R. principis in H. flava in Japan; this species was previously detected in H. japonica douglasi and H. danieli ticks in Russia and China, respectively, (10). And, we found a high prevalence of R. tamurae in A. testidinarium ticks; Imaoka et al. (5) recently reported that R. tamurae causes local skin inflammation without general JSP-like symptoms. We did not detect the human pathogen E. chaffeensis, but we identified 2 potentially new Ehrlichia species.

Our findings contribute to the known risks for exposure to *Rickettsia*-related pathogens in central and western Japan. Further studies may be required for the surveillance of additional pathogens, such as *Candidatus* Neoehrlichia mikurensis (2), which was recently recognized as a human pathogen.

This work was supported by the Research on Emerging and Reemerging Infectious Diseases grant from the Association for Preventive Medicine of Japan; grants for Research on Emerging and Reemerging Infectious Diseases from the Japanese Ministry of Health, Labor and Welfare (H18-Shinkou-Ippan-014, H21-Shinkou-Ippan-006, and H24-Shinkou-Ippan-008); and a Global Center of Excellence Program grant from Japanese Ministry of Education, Culture, Sports, Science and Technology (to N.O.).

Gaowa,¹ Norio Ohashi,¹ Minami Aochi, Wuritu, Dongxing Wu, Yuko Yoshikawa, Fumihiko Kawamori, Toshiro Honda, Hiromi Fujita,² Nobuhiro Takada, Yosaburo Oikawa, Hiroki Kawabata, Shuji Ando, and Toshio Kishimoto Author affiliations: University of Shizuoka Global Center of Excellence Program, Shizuoka, Japan (Gaowa, N. Ohashi, M. Aochi, Wuritu, D. Wu, Y. Yoshikawa, F. Kawamori); Shizuoka Institute of Environment and Hygiene, Shizuoka (F. Kawamori); Kagoshima Prefectural Institute for Environment Research and Public Health, Kaqoshima, Japan (T. Honda); Ohara General Hospital, Fukushima, Japan (H. Fujita); Fukui University, Fukui, Japan (N. Takada); Kanazawa Medical University, Ishikawa, Japan (Y. Oikawa); National Institute of Infectious Diseases, Tokyo, Japan (H. Kawabata, S. Ando); and Okayama Prefectural Institute for Environmental Science and Public Health, Okayama, Japan (T. Kishimoto)

¹These authors contributed equally to this article.

²Current affiliation: Mahara Institute of Medical Acarology, Anan, Japan.

DOI: http://dx.doi.org/10.3201/eid1902.120856

References

- National Institute of Infectious Diseases and Tuberculosis and Infectious Diseases Control Division, Ministry of Health, Labour and Welfare. Scrub typhus and Japanese spotted fever in Japan, as of December 2006. Infectious Agents Surveillance Report. 2006;27–2.
- Tabara K, Arai S, Kawabuchi T, Itagaki A, Ishihara C, Satoh H, et al. Molecular survey of *Babesia microti, Ehrlichia* species and *Candidatus* Neoehrlichia mikurensis in wild rodents from Shimane Prefecture, Japan. Microbiol Immunol. 2007;51:359–67.
- Ando S, Kurosawa M, Sakata A, Fujita H, Sakai K, Sekine M, et al. Human *Rickettsia heilongjiangensis* infection, Japan. Emerg Infect Dis. 2010;16:1306–8. http:// dx.doi.org/10.3201/eid1608.100049
- Fournier PE, Fujita H, Takada N, Raoult D. Genetic identification of rickettsiae isolated from ticks in Japan. J Clin Microbiol. 2002;40:2176–81. http:// dx.doi.org/10.1128/JCM.40.6.2176-2181.2002
- Imaoka K, Kaneko S, Tabara K, Kusatake K, Morita E. The first human case of *Rickettsia tamurae* infection in Japan. Case Rep Dermatol. 2011;3:68–73. http:// dx.doi.org/10.1159/000326941

- Inokuma H, Ohashi M. Jilintai, Tanabe S, Miyahara K. Prevalence of tick-borne *Rickettsia* and *Ehrlichia* in *Ixodes persulcatus* and *Ixodes ovatus* in Tokachi district, Eastern Hokkaido, Japan. J Vet Med Sci. 2007;69:661–4. http://dx.doi. org/10.1292/jvms.69.661
- Ybañez AP, Matsumoto K, Kishimoto T, Inokuma H. Molecular analyses of a potentially novel *Anaplasma* species closely related to *Anaplasma phagocytophilum* detected in sika deer (*Cervus nippon yesoensis*) in Japan. Vet Microbiol. 2012;157:232–6. http://dx.doi. org/10.1016/j.vetmic.2011.12.001
- Gaowa, Wuritu, Wu D, Yoshikawa Y, Ohashi N, Kawamori F, et al. Detection and characterization of *p44/msp2* transcript variants of *Anaplasma phagocytophilum* from naturally infected ticks and wild deer in Japan. Jpn J Infect Dis. 2012;65:79–83.
- Parola P, Rovery C, Rolain JM, Brouqui P, Davoust B, Raoult D. *Rickettsia slovaca* and *R. raoultii* in tick-borne rickettsioses. Emerg Infect Dis. 2009;15:1105–8. http:// dx.doi.org/10.3201/eid1507.081449
- Chahan B, Jian Z, Jilintai, Miyahara K, Tanabe S, Xuan X, et al. Detection of DNA closely related to '*Candidatus* Rickettsia principis' in *Haemaphysalis danieli* recovered from cattle in Xinjiang Uygur Autonomous Region Area, China. Vet Parasitol. 2007;144:184–7. http://dx.doi. org/10.1016/j.vetpar.2006.09.019

Address for correspondence: Norio Ohashi, Laboratory of Microbiology, Department of Food and Nutritional Sciences, School of Food and Nutritional Sciences, Graduate School of Integrated Pharmaceutical and Nutritional Sciences, University of Shizuoka, 52–1 Yada, Suruga-ku, Shizuoka 422–8526, Japan; email: ohashi@u-shizuoka-ken.ac.jp

Letters

Letters commenting on recent articles as well as letters reporting cases, outbreaks, or original research are welcome. Letters commenting on articles should contain no more than 300 words and 5 references; they are more likely to be published if submitted within 4 weeks of the original article's publication. Letters reporting cases, outbreaks, or original research should contain no more than 800 words and 10 references. They may have 1 Figure or Table and should not be divided into sections. All letters should contain material not previously published and include a word count.

Rickettsiae in Ticks, Japan, 2007–2011

Technical Appendix

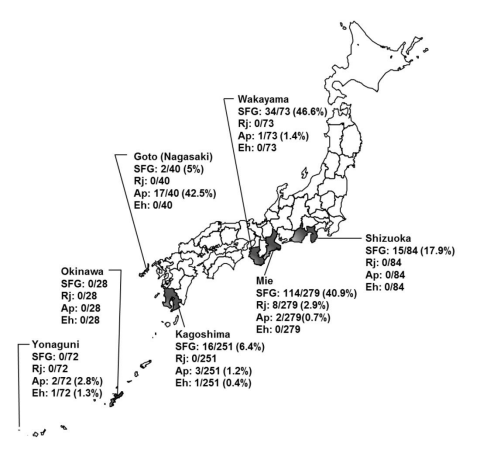


Figure 1. Prefectures where ticks were collected for a study of *Rickettsia* spp.–related pathogens in ticks in central and western Japan, 2007–2011. SFG, spotted fever group rickettsiae; Rj, *R. japonica*; Ap, *A. phagocytophilum*; Eh, *Ehrlichia* spp. Numbers indicate the number of positive ticks/number of ticks tested. The prevalence of respective *Rickettsiales* bacteria in each area is shown in parentheses.

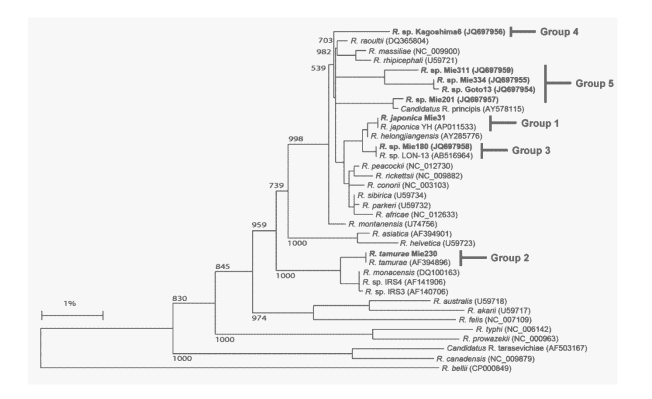


Figure 2. Phylogenetic classification of *Rickettsia spp. gltA* sequences detected in ticks inhabiting central and western Japan, 2007–2011. The tree, based on the *gltA* sequences (1,115–1,123 bp), was constructed by using the neighbor-joining method with 1,000 bootstrap resamplings. **Boldface** font indicates *Rickettsia* spp. identified in this study. Numbers in parentheses indicate GenBank accession numbers for the respective sequences. The scale bar indicates nucleotide substitutions per site.