Mixed Scrub Typhus Genotype, Shandong, China, 2011

To the Editor: Scrub typhus, which is caused by the bacterium Orientia tsutsugamushi, had been considered a disease of the tropical zone in China until the first outbreak in the temperate zone of Shandong Province in 1986 (1). The Sdu-2 genotype had not been reported for human patients in China since its detection in the striped field mouse (Apodemus agrarius) during 2007 (2). We report a case in a person in Shandong Province who was co-infected with the HSX and Sdu-2 genotypes of O. tsutsugamushi.

A female farmer, 55 years of age, was admitted to Mengyin County Hospital on September 29, 2011, with a 5-day history of fever (highest temperature 39°C), headache, back pain, chills, anorexia, and nausea. Her back pain was exacerbated as her fever increased. A black eschar (10 mm × 20 mm) was observed above the left breast, and 2 enlarged lymph nodes (20 mm × 30 mm) that were tender to the touch were found behind the right ear. The patient’s Glasgow Coma Scale score was 15. No comorbidities were reported. Laboratory tests showed slight leukopenia, low potassium level, decreased albumin/globulin ratio, elevated lactate dehydrogenase level, and elevated C-reactive protein level. Results of urine dry chemical analysis were positive for occult blood and leukocytes. The Weil-Felix reaction to the Proteus OXK antigen was negative. Mild abnormalities were observed in results of an electroencephalogram and an electroencephalogram topographic map. The patient was suspected to have scrub typhus complicated by encephalitis and urinary tract infection. Doxycycline (200 mg once daily) and symptomatic treatment were administered immediately. The patient’s symptoms had generally subsided on day 5 of hospitalization. Indirect immunofluorescence assay showed a 4-fold rise in IgG titer against O. tsutsugamushi between acute-phase (128) and convalescent-phase (512) serum samples collected 18 days apart.

Acute-phase whole blood was collected before administration of antimicrobial drugs. The eschar was collected after natural desquamation. DNA from the whole blood and eschar was isolated and screened for O. tsutsugamushi by using PCR primers E and B to target the 56-kDa type-specific antigen gene (3). Type-specific amplification of O. tsutsugamushi was performed (4,5). Normal values for fragment yield are 407, 230, 242, 220, 600, and 523 bp for Gilliam, Karp, Kato, Kuroki, Saitama, and Kawasaki genotypes, respectively.

Three products that had the same molecular weight as those from Kato, Saitama, and Kawasaki genotypes were generated from the blood sample by using type-specific amplification. However, sequence analysis of amplicons yielded by using type-specific primers and those yielded by using primers E and B demonstrated a co-infection in the blood with 2 genotypes of O. tsutsugamushi, which were designated as ZZF-KW (661 bp, GenBank accession no. JX644590) and ZZF-HSB (498 bp, GenBank accession no. JX644591). Co-infection with the 2 genotypes was also detected in the eschar. BLAST (http://blast.ncbi.nlm.nih.gov/) analysis showed that ZZF-KW had 100% sequence similarity to the HSX genotype of O. tsutsugamushi (657 bp, GenBank accession no. JX202566) and that ZZF-HSB had 100% sequence similarity to Sdu-2 genotype (476 bp, GenBank accession no. EF543196). A nucleotide identity of 72% was shown between ZZF-KW and ZZF-HSB. Nucleotide identities of partial 56-kDa type-specific antigen genes among the 2 O. tsutsugamushi genotypes and the reference types are shown in the Table.

Co-infection with HSX and Sdu-2 genotypes of O. tsutsugamushi was detected in a single case. Mixed genotype infections could not be detected by using 1 amplicon, which could amplify only a predominant genotype. Type-specific amplification is a convenient method for the detection of mixed genotype infection, although this method has limitations for identification of novel genotypes. The Sdu-2 genotype was not differentiated from Kato, Saitama, and Kawasaki genotypes by using type-specific primers, but differentiation was achieved by gene sequencing. Considering the genotypic diversity (6) and mixed genotype infection of the agent, type-specific primers require redesign for type designation and subsequent sequencing is recommended for verification.

Coexistence of multiple genotypes O. tsutsugamushi was reported for 25% of scrub typhus patients in a study population in Thailand (7). Mixed genotype infections of a single pathogen species in the ecosystem are common; these mixed genotypes provide possibilities for recombination events and prompts the coevolution of host–parasite interactions (8). Disease severity and epidemiologic characteristics could be influenced by

---

Table. Nucleotide sequence homologies of partial 56-kDa type-specific antigen gene sequences among Orientia tsutsugamushi genotypes and reference strains, China, 2011

<table>
<thead>
<tr>
<th>Genotype</th>
<th>HSX</th>
<th>Sdu-2</th>
<th>Gilliam</th>
<th>Karp</th>
<th>Kato</th>
<th>Kawasaki</th>
<th>TA686</th>
<th>HSB</th>
</tr>
</thead>
<tbody>
<tr>
<td>ZZF-KW</td>
<td>100</td>
<td>68.8</td>
<td>90.3</td>
<td>72.7</td>
<td>73.8</td>
<td>96.0</td>
<td>67.3</td>
<td>74.7</td>
</tr>
<tr>
<td>ZZF-HSB</td>
<td>68.8</td>
<td>100</td>
<td>68.1</td>
<td>72.1</td>
<td>66.6</td>
<td>67.7</td>
<td>77.8</td>
<td>74.3</td>
</tr>
</tbody>
</table>
ecologic interactions between genetically diverse strains (8). In this report, we describe a single case; comparison of disease severity of scrub typhus caused by mixed and single genotype infections should be studied further.

Simultaneous infection with multiple antigenic strains of *O. tsutsugamushi* was detected in an individual mite, *Leptotrombidium arenicola* (9), a probable vector of scrub typhus. Infection with multiple *O. tsutsugamushi* strains may be caused by being bitten by multiple mites or by multiple genotypes coexisting within individual mites (7). We ascribed the co-infection to the second cause because the 2 genotypes were simultaneously detected from an eschar sample associated with the bite of 1 mite examined in this study. There may be diverse genotypic co-infection patterns of *O. tsutsugamushi*. Mechanisms of in-host interactions between genetically diverse strains of *O. tsutsugamushi* and the initiated host response require the establishment of animal models for further research.

Acknowledgment

We thank Shu-Xia Li for her assistance with sample collection.

The work was supported by grants from the National Natural Science Foundation of China (no. 81273133 and no. 30972515).

Meng Zhang, Zhong-Tang Zhao, Xian-Jun Wang, Zhong Li, Lei Ding, Shu-Jun Ding, and Li-Ping Yang

Author affiliations: Shandong University, Jinan, China (M. Zhang, Z.-T. Zhao, L. Ding, L.P. Yang); and Shandong Center for Disease Control and Prevention, Jinan (X.-J. Wang, Z. Li, S.-J. Ding)

DOI: http://dx.doi.org/10.3201/eid2003.121349

References


Address for correspondence: Zhong-Tang Zhao, Department of Epidemiology and Health Statistics, School of Public Health, Shandong University, 44 Wenhua Rd, Jinan 250012, Shandong Province, PR China; email: ztzhao@sdu.edu.cn

The opinions expressed by authors contributing to this journal do not necessarily reflect the opinions of the Centers for Disease Control and Prevention or the institutions with which the authors are affiliated.

Staphylococcus delphinus and Methicillin-Resistant *S. pseudintermedius* in Horses, Canada

To the Editor: *Staphylococcus aureus* is a well-known pathogen of horses (1), but the role of other coagulase-positive staphylococcal species in these animals is unclear. *S. pseudintermedius* and *S. delphinus*, members of the *S. intermedius* group (SIG), cause infections in some companion animals and equids (2), can be multidrug resistant, and could be a concern in horses. Members of SIG are difficult to differentiate by using biochemical methods and require molecular techniques for accurate species-level identification (3); therefore, misidentification of these pathogens could occur.

Methicillin-resistant or unusual staphylococci that are isolated at the Ontario Veterinary College Health Sciences Centre by the University of Guelph Animal Health Laboratory (AHL) routinely undergo further characterization. During 2011, the laboratory tested 5 isolates from different horses that were coagulase-positive staphylococci other than methicillin-resistant *S. aureus* (MRSA). Isolates were identified by using matrix-assisted laser desorption/ionization–time of flight (MALDI-TOF) mass spectrometry, *S. pseudintermedius* or *S. delphinus* PCR (4), and sodA sequence analysis (5). Isolates were further characterized, as indicated, by direct repeat unit typing (5), pulsed-field gel electrophoresis (PFGE) (6), meca PCR (7), penicillin-binding protein 2a latex agglutination test, and antimicrobial drug susceptibility testing by broth microdilution and/ or disk diffusion. A search of AHL’s database was performed to identify other *S. pseudintermedius* and *S. delphini* isolates for all submissions of samples from equids during January 2011–August 2012.