Sika Deer Carrying Babesia Parasites Closely Related to B. divergens, Japan

To the Editor: Human babesiosis caused by Babesia divergens, a zoonotic pathogen of bovines in Europe, is an emerging tickborne disease (1). In the United States, a closely related Babesia sp. was identified in persons in Missouri and Kentucky and in eastern cottontail rabbits (Sylvilagus floridanus) on Nantucket Island, Massachusetts (2–5). We report that sika deer (Cervus nippon) in Japan also carry parasites genetically closely related to B. divergens.

During November 2007–February 2008 and November 2008–February 2009, we collected blood samples from 96 wild sika deer throughout Japan. We purified DNA from the blood, which had been stored in a freezer, by phenol/chloroform extraction and performed nested PCR for B. divergens 18S rRNA (rDNA), chaperonin-containing T-complex protein 1 eta subunit (CCT7, formerly described as CCTη) (6), and β-tubulin (7) genes. Primers for rDNA were designed from the sequences of related Babesia spp. (GenBank accession nos. U16370, U16369, and AY046575): dv101F (5′-ACAACTGTTATGTTCTTGG-3′) for the first and dv159F (5′-GCTAATGCAAGTTCAGG-GCTTTTGGGC-3′) and dv1353R (5′-CGGACGAACCTTTTTACG-3′), and dv159F (5′-GCTAATGCAAGTTCAGG-GCTTTTGGGC-3′) and dv1296R (5′-CGGACGAACCTTTTACTAGC-GACATAG-3′) for the first and second rounds, respectively. CCT7 primers were similarly designed (GenBank accession nos. AB367924 and AB367925): Bdv/odoCCTF1 (5′-CAAATGAGYCACTTMC-CAACCTACC-3′) and Bdv/CCTR1 (5′-ATCTCAGCAGTCTACATGTCGACACCTC-3′), and Bdv/odoCCTF2 (5′-CAACCTACCRAT-TCTCITYYTGAAAGGG-3′) and BdivCCTR2 (5′-GGCTATTAGTCGGGGCT-CAGC-3′) for the first and second rounds, respectively. The β-tubulin PCR protocol has been described (7).

Of the 96 blood samples, 12 from 5 prefectures (Hokkaido, Iwate, Tochigi, Nagano, and Miyazaki) were positive for Babesia rDNA (online Technical Appendix, http://wwwnc.cdc.gov/EID/article/20/8/13-0061-TechnicalAppendix.pdf). The sequence for sample 08–22 from Hokkaido (GenBank accession no. KC465978) was distinct from sequences of the other 11 samples (97.5%–97.6% identity in 1,041 bp), which consisted of 7 variant sequences (GenBank accession nos. KC465973–7 and AB857845–6) and 5 identical sequences (GenBank accession nos. KC465977 and AB861504–7) (99.7%–100% identity). The 5 identical sequences varied in only 1 of 909 bp from B. divergens rDNA from an Ixodes persulcatus tick in Russia [GenBank accession no. GU057385] (8).

β-tubulin (900 bp) was also amplified from the 12 Babesia rDNA–positive samples. Ten of the sequences consisted of 3 sequence variants (99.9% identity; GenBank accession nos. KC465969, KC465970, and KC465968/AB861508–14). The 2 divergent sequences (GenBank accession nos. KC465971 [08–22] and KC465989 [08–25]) were most similar to B. odocoilei (GenBank accession no. KC465972; 91% identity) and Theileria orientalis (GenBank accession no. AP011947; 79.9% identity), respectively. Thus, at least 1 deer likely had Babesia and Theileria spp. infections.

CCT7 was amplified from 10 of the 12 Babesia-positive blood samples. The sequences (GenBank accession nos. KC465979–88) were more heterogeneous (98.7%–99.9% identity) than those for rDNA and β-tubulin; this finding was expected because CCT7 evolves more quickly (6).

We generated a neighbor-joining phylogenetic tree (ClustalW, http://clustalw.ddbj.nig.ac.jp/index.php?lang=ja) from the Babesia rDNA sequences from our study and from GenBank (Figure, panel A). The tree shows a distinct lineage (Asian) cluster for the deer parasites, except for 08–22 (GenBank accession no. KC465978), within a clade also holding the B. divergens strains (human and bovine) from Europe (European Union lineage). The tree also shows a cluster encompassing Babesia spp. (human and rabbit) from the United States; B. divergens (deer), B. capreoli (deer), and Babesia sp. (chamois) from Europe; and B. divergens (human) from Portugal. Sequence 08–22 branches with Babesia spp. in Ix. ovatus ticks from Japan (GenBank accession nos. AY190123 and AY190124) (9). The branch lengths indicate clear separation between the isolates from sika deer and ticks, suggesting that the clustering may be attributable to the limited number of available related sequences.

We also generated a phylogenetic tree of β-tubulin sequences (900 bp), which produced similar topology and high bootstrap support (Figure, panel B). However, the limited number of relevant β-tubulin sequences precludes conclusions regarding the phylogeny of Babesia parasites. A CCT7 phylogenetic tree was not generated because of the paucity of sequences. Overall, the phylogenies suggest that B. divergens and B. capreoli–related parasites are found worldwide in temperate zones of the Northern Hemisphere, including Europe (1), the United States (2–5), Russia (8), and Japan.

We showed the presence of B. divergens–like rDNA, β-tubulin, and CCT7 genes in sika deer from different Japanese prefectures, confirming the presence of this parasite in Japan. B. capreoli, which is serologically indistinguishable from B. divergens, was previously reported in sika deer (10). However, no molecular data for the B. capreoli isolate exist, so no
conclusion may be drawn regarding its relationship to the *B. divergens*–like parasites from sika deer in our study.

There is an overabundance of wild sika deer in Japan because these animals easily adapt to a variety of climates, vegetation, and geography. Increased human exposure to deer habitats increases the risk of exposure to tickborne zoonoses, such as those caused by *Babesia* spp. In humans, infections caused by *B. divergens* and *B. divergens*–like parasites can be life threatening; fatality rates of 42% and 33% have been reported in infected asplenic patients in Europe and the United States, respectively (1–3).

The findings from our study emphasize the need for increased clinical awareness of babesiosis in Japan and globally. They also emphasize the need for the swift diagnosis of suspected cases and prompt treatment of confirmed cases, especially in asplenic patients at high risk for the potentially deadly consequences of babesiosis.

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References


Transcontinental Movement of Asian Genotype Chikungunya Virus

To the Editor: Chikungunya virus (CHIKV), a mosquito–transmitted virus (family Togaviridae, genus Alphavirus), was first isolated >60 years ago in Africa and is responsible for epidemics of acute polyarthralgia. During CHIKV epidemics, the transmission cycle is from humans to mosquitoes, with no intervening amplifying host, and the virus can rapidly disseminate, infecting large numbers of persons. Epidemics have been described in Africa, the Middle East, Europe, India, and Southeast Asia. On the basis of detailed clinical descriptions of the disease, chikungunya fever, it appears that CHIKV caused epidemics in the Caribbean (St. Thomas, US Virgin Islands) and the southeastern coastal United States during the early 19th century (1).

Genetic studies show that the virus has evolved into 3 distinct genotypes: West African, East/Central/South African (ECSA), and Asian (2). The genotypes likely indicate independent evolution of the virus in historically isolated areas. Phenotypic differences have been described between genotypes and between individual strains, most notably an E1 mutation among some ECSA strains, which facilitates replication in Aedes albopictus mosquitoes (3). However, more recently, the movement of virus genotypes has increased dramatically, probably as a direct result of increased movement of humans and increased commercial trade. Beginning in 2005 and through 2006, the ECSA genotype virus was responsible for an explosive epidemic, during which the virus moved from coastal Kenya to islands adjacent to southeastern Africa and then to India, where >1 million cases were recorded (2). During this time, imported cases were reported worldwide, and in some instances, autochthonous transmission was detected in distal locations (4,5).

In October 2013, the arbovirus diagnostic laboratory at the Centers for Disease Control and Prevention (CDC; Fort Collins, CO, USA) detected CHIKV in human serum specimens from Yap State, Federated States of Micronesia; the specimens were collected during an epidemic of disease clinically compatible with chikungunya fever. In December 2013, the French National Reference Centre for arboviruses confirmed that CHIKV was responsible for an epidemic occurring on St. Martin Island, French West Indies, in the Caribbean (6). In January 2014, the Caribbean Public Health Agency detected CHIKV in 2 human serum specimens from the British Virgin Islands (BVI); the samples were subsequently confirmed by CDC to be positive for CHIKV.

By using next-generation sequencing, we determined the complete nucleotide sequence for 1 of the CHIKV specimens detected in BVI and for 2 of the CHIKV specimens detected in Yap. DNA libraries for next-generation sequencing were prepared directly from RNA extracted from serum, and the amplified libraries were sequenced by using the Ion Torrent Personal Genome Machine (Life Technologies, Grand Island, NY, USA). The CLC Genomics Workbench (CLC Bio, Aarhus, Denmark) and Lasergene NextGen (DNASTAR, Madison, WI, USA) were used to analyze and assemble raw sequence reads. ClustalW (http://www.ebi.ac.uk/Tools/msa/clustalw2/) was used to align the complete genome sequences with a variety of CHIKV sequences, representing the 3 genotypes, from GenBank. Nearly identical phylogenetic trees were generated by several methods (i.e., minimum evolution, maximum likelihood, neighbor joining); a representative neighbor-joining tree generated and analyzed with 1,000 replicates for bootstrap testing is shown in the Figure.
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**Technical Appendix**

Prefectures with no sika deer positive for *Babesia* spp. (N): Chiba (5), Shizuoka (3), Hyogo (1), Hiroshima (5), Tokushima (5), Oita (3), Kumamoto (4), Kagoshima (4)

Technical Appendix Figure. Map of Japan showing prefectures where rDNA genes of *Babesia divergens*-like parasites were detected by PCR in blood samples from sika deer (black areas). Sample identification numbers are listed under prefecture names; numbers in parentheses indicate the number of positive samples/total number tested samples. A list of prefectures where no positive samples were found is shown at the bottom of the figure; the number of animals tested is shown in parentheses.