Coxiella burnetii in Rodents on Heixiazi Island at the Sino-Russian Border

Lijuan Liu, Xu Baoliang, Fu Yingyun, Li Ming, Yang Yu, Hou Yong, Wang Shasha, Hu Manxia, Guo Tianyu, Jiang Chao, Sun Xiaohong, and Wang Jing*

Institute of Health Quarantine, Chinese Academy of Inspection and Quarantine, Beijing, People’s Republic of China; Department of Health and Quarantine, Heilongjiang Entry-Exit Inspection and Quarantine Bureau, Harbin, People’s Republic of China; Department of Zoology, Shenyang Agriculture University, Shenyang, People’s Republic of China; Department of Health and Quarantine, Suifenhe Entry-Exit Inspection and Quarantine Bureau, People’s Republic of China

Abstract. This work is a molecular epidemiologic study to detect the incidence of Coxiella burnetii in rodents on Heixiazi Island at the Sino-Russian border. Liver tissues were extracted and processed to test the incidence of C. burnetii infection using polymerase chain reaction analysis. In total, 18% (66 of 370) of rodents tested positive for infection. The results of logistic regression analysis indicated that infection with C. burnetii was associated significantly with weight and month of capture, and infection was found in all rodent species that were observed; there was no significant difference of sex on the infection of C. burnetii. Though phylogenetic analysis disclosed heterogeneity in the nucleotide sequences isolated from the island rodents, the majority of observed strains were among the most common strains found worldwide. This is the first report on the incidence of C. burnetii in rodents on Heixiazi Island at the Sino-Russian border.

INTRODUCTION

Recently, several reported outbreaks of Q fever in European countries have raised public concerns regarding this disease, which for decades has been known to occur throughout the world.1,2 Humans can be infected through inhalation of contaminated aerosols, ingestion of contaminated food, or skin trauma. The manifest symptoms of Q fever in humans include both acute and chronic illnesses, where chronic illness occurs in ~1–3% of patients. Acute cases occur as an influenza-like illness, hepatitis, and/or pneumonia, which occasionally lead to a lethal respiratory distress syndrome. Endocarditis and hepatitis are the most frequent and serious manifestations of illness in Q fever.3

Q fever is caused by Coxiella burnetii, an intracellular organism with reservoirs in birds, arthropods, wild and domestic mammals.4 The organism has a spore-like morphology that is extremely resistant to heat, pressure, desiccation, and other antiseptic compounds. It can survive in the ambient environment for long periods. Because of its characteristics, C. burnetii aerosols can be used in biological warfare, and it is considered a potential terrorist threat.5

In addition, with economic development, increased habitat loss and fragmentation have made human contact with wild species more frequent. Parks often serve as refuges for wildlife, but they may also be important transmission zones of diseases from wildlife to humans.6 Investigations that attempt to discover wild reservoir species of zoonotic diseases are critically important for understanding the risk of pathogen exchange between wild and human populations.

Heixiazi Island, located at the junction of Heilong River (called Amur in Russia) and the Wusuli River, was once occupied by the former Soviet Union during a 1929 border skirmish. After long years of negotiations, 174 km², or about half the island, was returned to China after 2008. Because of its special location, Heixiazi Island is of great strategic importance. Its natural environment and cultural history has allowed for the establishment of a distinctive Heixiazi Island tourist attraction, which displays a diverse landscape and abundant wildlife.

Our main objectives for this study were to determine whether the island was the endemic area for Q fever and whether the wild rodents inhabiting the island were naturally infected with C. burnetii; we could then assess the risk of Q fever for visitors to the island.

MATERIALS AND METHODS

Ethics statement. The handling of rodents was conducted in compliance with the Animal Welfare Provision of the Chinese Academy of Inspection and Quarantine (CAIQ). All animal experiments were performed following the guidelines of CAIQ.

Trapping of rodents. Rodents were collected from April through October, 2011, on a monthly basis, from different landscapes, including woodland, grassland, and edge of bush in Heixiazi Island.

Using fresh peanuts as bait, rodents were trapped by snap traps. Traps were set in the evening and checked early in the morning. Generally, each rodent was placed in an individual cloth bag in the field and then transported to the local laboratory. A zoologist identified each rodent according to morphologic features specific to its species and developmental stage. After identification of their species and sex and developmental stage, the rodents were dissected. Liver tissues from the rodents were removed and stored immediately in liquid nitrogen and then transported to CAIQ laboratory for further processing.

Detection of C. burnetii infection. Total genomic DNA was extracted from the liver tissue samples by using a Tissue DNA Extract kit (Tiangen Biotech Inc., Beijing, China), following the instructions of the manufacturer. Nested polymerase chain reaction (PCR) was performed to amplify the coml gene as previously described.7 The primers used in the first- and second-round PCR reactions are listed in Table 1.

To avoid possible contamination, DNA extraction, the reagent setup, amplification, and agarose gel electrophoresis were performed in separate rooms, and negative control samples (distilled water) was included in all amplifications.

Amplifications were performed in a total volume of 50 µL containing 5 µL of DNA template, 0.5 µM MgCl₂, 0.2 µM

*Address correspondence to Wang Jing, No. A 3, Gaobeidian North Road, Chaoyang District, Beijing 100123, China. E-mail: wangjing0115@126.com
analyzed using either the accession nos. JX522479 generated in the study were deposited in GenBank under pseudo-replicate data sets. Some of the nucleotide sequences logenies was estimated using bootstrap analysis with 1,000 (version3.1). The statistical significance of the inferred phy-
sion 1.83) followed by phylogenetic analysis using MEGA
The resulting sequences were analyzed using Clustal X (ver-
sent to Sangon Biotech (Shanghai) Co., Ltd., for sequencing.
Instruments, Inc., Winooski, VT). The purified products were
were purified using the Omega Gel extraction kit (BioTek
PCR product on a partial
Coxiella burnetii by nested
gene, with an overall positive rate of
(18.4%), whereas the minority of rodents included members from Apodemus peninsulae (4.1%), Eutamias sibiricus (1.4%), and Rattus norvegicus (0.8%), respectively. In total, 66 rodents were positive for C. burnetii infection by nested PCR on a partial Com1 gene, with an overall positive rate of 18%, all observed species included rodents infected with the organism, and the infection rates of C. burnetii among various species were not significantly different (χ^2 = 3.294, P > 0.05). The average weight of C. burnetii infection in rodents was 31.71 ± 17.86 g, and was not significantly higher than 30.57 ± 15.24 g in non-infection rodents (F = 0.280, P > 0.05). Rodents were captured in seven continuous months from April to October. There was a significant difference in the prevalence of C. burnetii between the months of capture (χ^2 = 73.768, P < 0.05). The highest prevalence of C. burnetii infection in rodents was observed in September, followed by July and May; no positive samples were detected in August (see in Table 3).
Multivariable logistic regression analysis was used to detect the risk factors of C. burnetii, including month of capture, weight, gender, and species. To detect whether there was a difference of prevalence among the species, Eutamias sibiricus was used for the indicator index and compared with the other species (shown in Table 4), the results showed there was not a significant difference among the rodent species. In this model, month of capture and weight were found to be associated with the infection of C. burnetii in the rodents (P < 0.05).
The 438-nt fragment corresponding to the Com1 gene of the positive species was sequenced. Considering sequences of some positive samples from the same study site were nearly identical, 11 representative sequences of the positive samples were used for alignment and phylogenetic analysis. The identities of the nucleotide sequences isolated from rats on the island ranged from 85.9% to 100% (see Figure 1). Figure 2 shows the variations of amino acid sequences of C. burnetii isolated in the Island. The results of phylogenetic analysis

<table>
<thead>
<tr>
<th>Month of capture</th>
<th>Prevalence of C. burnetii in rodents</th>
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<tbody>
<tr>
<td>April</td>
<td>3% (2/72)</td>
</tr>
<tr>
<td>May</td>
<td>26% (19/72)</td>
</tr>
<tr>
<td>June</td>
<td>16% (10/63)</td>
</tr>
<tr>
<td>July</td>
<td>28% (11/39)</td>
</tr>
<tr>
<td>August</td>
<td>0% (0/51)</td>
</tr>
<tr>
<td>September</td>
<td>58% (19/33)</td>
</tr>
<tr>
<td>October</td>
<td>13% (5/40)</td>
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</table>

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<tr>
<th>Variables</th>
<th>OR and 95% CI</th>
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</thead>
<tbody>
<tr>
<td>Month of capture</td>
<td>0.002*</td>
</tr>
<tr>
<td>Species</td>
<td>0.449</td>
</tr>
<tr>
<td>Species (1)</td>
<td>0.747</td>
</tr>
<tr>
<td>Species (2)</td>
<td>0.814</td>
</tr>
<tr>
<td>Species (3)</td>
<td>0.536</td>
</tr>
<tr>
<td>Species (4)</td>
<td>0.403</td>
</tr>
<tr>
<td>Species (5)</td>
<td>0.828</td>
</tr>
<tr>
<td>Sex</td>
<td>0.378</td>
</tr>
<tr>
<td>Weight</td>
<td>0.048*</td>
</tr>
</tbody>
</table>

*Means P < 0.05. OR = odds ratio. CI = confidence interval.
showed that 10 of the strains clustered in a clade with others isolated from the United States, Japan, China, and Russia, and No.221 were placed in a separate clade, which is distinct from the other sequences (see Figure 3).

**DISCUSSION**

Emerging zoonoses are linked to increasing globalization. These diseases impact not only animal populations but also humans who are in close contact with animal species. Q fever is a zoonotic disease caused by *Coxiella burnetii* that has been widely distributed in nature. *Coxiella burnetii* in domestic animals, such as cattle, sheep, and goats, has been widely reported. However, there is little detailed epidemiological data regarding the distribution and determinants of *C. burnetii* infection in rodents. Recent findings in Germany indicated regular sightings of wild rodents as risk factors for *C. burnetii* infection in humans, suggesting wild rodents as a direct source for human infection. In this study, *C. burnetii* was identified by PCR in 18% of the rodents captured on Heixiazi Island at the Sino-Russian border, from April to October in 2011. However, it is difficult to compare these findings with those reported in other works, which present information on the presence of antibodies to *C. burnetii* in rodents. The statistical results showed that weight was associated with *C. burnetii* infection, possibly because larger rodents become matured and move widely and have more chances to be infected with *C. burnetii*. The reason for the association between month of capture and *C. burnetii* infection may be the temperature differences between months. Previous studies showed that Hantaviruses coevolved with specific rodents which means certain pathogens dominate in certain species. However, in this study, no significant association of specific rodent species and *C. burnetii* infection was found, and all observed rodent species incidences of infection with *C. burnetii*. This result was partly consistent with the previous study that identified *Rattus norvegicus* as a host for *C. burnetii* infection. In a human-based study, it was found that sex was related with *C. burnetii* infection, where men were 2.5 times more likely to be infected with the pathogen than women; the same result was also reported in mice in a laboratory-based study. However, the results of our study were different; no significant association was found between *C. burnetii* infection and sex in wild rodents.

The results of the molecular epidemiologic analysis showed that although discrepancy existed in the nucleotide sequences of *C. burnetii* in the tested rodents, the majority of sequences showed high shared identity in the region of the *com1* gene, and the sequences aligned with other strains isolated elsewhere in the world. This sequence heterogeneity led us to continue surveillance of the pathogen to describe the variants of the pathogen that occur in the island rodent population.

In conclusion, this study confirms that rodents are reservoirs of *C. burnetii*, and it is the first evidence of *C. burnetii* infection in rodents on Heixiazi Island at the Sino-Russian border of Heilongjiang Province, China. As reservoirs,
Figure 3. Phylogenetic analysis of the coml gene of Coxiella burnetii detected in the study from April to October 2011. The accession nos. included in the tree were 208(JX522479), 209(JX522480), 212(JX522481), 213(JX522482), 214(JX522483), 216(JX522484), 221(JX522485), 225(JX522486), 233(JX522487), 240(JX522488), 242(JX522489), JX131364, HM804027, AF317646, AF318148, AB004709, HM237793, and GU797241, respectively.

however, rodents have not yet been shown as a vector for transmission of Q fever to humans, and further studies need to address pathogen maintenance in and transmission to humans.

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Disclosure: Dr. Liu is an epidemiologist at the Institute of Health Quarantine of CAIQ. Her primary research interest is public health in the field of Quarantine.

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Authors’ addresses: Lijuan Liu, Xu Baoliang, Guo Tianyu, Yang Yu, Sun Xiaohong, and Wang Jing, Institute of Health Quarantine, Chinese Academy of Inspection and Quarantine, Beijing, PRC. E-mails: lijyhx@126.com, xubaol@yahoo.com.cn, gty200411@sina.com, redyy99@gmail.com, dotsunny@hotmail.com, and wangjing0115@126.com. Fu Yingqun, Li Ming, and Hou Yong, Department of Health and Quarantine, Heilongjiang Entry-Exit Inspection and Quarantine Bureau, Harbin, PRC, E-mails: fuyingqun109@yahoo.com.cn, chff2002@163.com, lemon_ly2003@126.com. Jiang Chao and Wang Shasha, Department of Zoology, Shenyang Agriculture University, Shenyang, PRC, E-mails: jc_0043@163.com and wangsha_327@163.com. Hu Manxia, Department of Health and Quarantine, Suifenhe Entry-Exit Inspection and Quarantine Bureau, PRC, E-mail: hmx1@163.com.

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