Mosquitoes and Mosquito-Borne Arboviruses in the Qinghai-Tibet Plateau—Focused on the Qinghai Area, China

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Abstract. An investigation was conducted to identify the distribution of mosquitoes and mosquito-borne arboviruses in the Qinghai-Tibet Plateau, China from July to August in 2007. A total of 8,147 mosquitoes representing six species from three genera (Aedes, Culex, and Anopheles) were collected in three locations (Geermu city, altitude of 2,780 m; Xining city, 2,200 m; Minhe county, 1,700 m). Six virus isolates were obtained including Tahyna virus (TAHV), Liaoning virus, and Culex pipiens pallens Densovirus. A serosurvey showed immunoglobulin G antibodies by immunofluorescence assay (IFA) against TAHV in residents of all three locations. The IFA-positive human samples were confirmed by 90% plaque-reduction neutralization tests (PRNT₉₀) against TAHV with titers ranging from 1:20 to 1:10,240. In addition, TAHV sero-positive cows, sheep, and swine were found in these locations. This investigation represents the first isolation of TAHV from Aedes (Ochlerotatus) detritus and the first evidence of TAHV infection in residents and livestock in the Qinghai-Tibet Plateau.

INTRODUCTION

The Qinghai area is located in the western region of mainland China, and most of its area is located in the Qinghai-Tibet Plateau. Qinghai lies between latitude 31°32’N and 39°20’N and longitude 98°24’E and 103°04’E, and it covers a 721,000-km² area. The average altitude of Qinghai is over 3,000 m, and it reaches a high of 6,880 m (Bukadaban Peak of Kunlun Mountains) and a low of 1,650 m (Xiachuankou village in Minhe county). Qinghai borders Gansu Province on the northeast, Xinjiang Uygur Autonomous Region (Xinjiang) on the northwest, Sichuan Province on the southeast, and Tibet Autonomous Region (Tibet) on the southwest. In recent years, communication and trade among Tibet and the other western provinces in China has intensified as a result of the construction of the Qingzang Railway that connects Xining city (capital of Qinghai) to Lhasa city (capital of Tibet). Consequently, Qinghai has become a crucial region that connects inland provinces of China with countries in South Asia such as Nepal, India, and Bangladesh. As a result, trade and travel across Qinghai are increasing rapidly, and with them, the likelihood of introduction and exportation of vectors and vector-borne diseases increases. For example, plague bacillus was identified in a woodchuck captured on the train from Geermu city in September 2000, and two human plague cases occurred in Wulan county along the Qingzang Railway in October 2004. Therefore, it is important to strengthen the detection and monitoring of vector-borne diseases in the region.

Previous research showed that bacterial vector-borne diseases such as plague, typhus fever, brucellosis, and Lyme disease have occurred in Qinghai. Since the 1950s, there was reported to be no Japanese encephalitis in this area, and as a result, no systematic investigations of mosquito-borne arboviruses have been carried out. However, seasonal fever and viral encephalitis cases of unknown etiology are frequently reported in the summer and autumn in Qinghai. Here, we describe a study of the distribution of mosquitoes and mosquito-borne viruses and show evidence of arbovirus infection in humans and livestock in three locations in Qinghai (Geermu city, Xining city, and Minhe county).

MATERIALS AND METHODS

Collection of mosquitoes and site descriptions. During the period from July 30 to August 14 in 2007, three collection sites were chosen in Geermu city (Xinhua village, Baoku village, and Xititan village), Xining city (Datonghe village, Taobei village, and Jujiawan village), and Minhe county (Caotan village, Jintian village, and Laqia village) within Qinghai (Figure 1). Geermu city is located in the western region of Qinghai with an average altitude of 2,780 m, an annual average temperature of 4.3°C, and an annual average precipitation of 42.5 mm. Xining city is located in the eastern region of Qinghai with an average altitude of 2,200 m, an annual average temperature of 7.7°C, and an annual average precipitation of 520.3 mm. Minhe county is located in the most eastern region of Qinghai with an average altitude of 1,700 m, an annual average temperature of 9°C, and an annual average precipitation of 350 mm.

Each village was sampled for one night from 19:30 to 21:30 hours. Adult mosquitoes were collected using light traps baited with CO₂. Three different kinds of habitat were sampled in each location: wheat field, grass river bank, and pond with heavy reed vegetation. Female mosquitoes were identified in the species using morphologic characteristics; they were pooled into groups of approximately 50–100 mosquitoes and stored in liquid nitrogen.

Isolation of viruses. Mosquitoes were removed from the liquid nitrogen and immediately homogenized and centrifuged as reported previously. The supernatants were inoculated into monolayers of BHK-21 and C6/36 cells and incubated at 37°C and 28°C, respectively. Cells were observed for cytopathic effect (CPE) daily from day 1 through day 7 post-inoculation. A specimen was regarded as a positive isolate if it caused CPE in three successive cell passages. Suckling mice were inoculated with 0.02 ml supernatant of positive isolates intracerebrally and observed for 14 days thereafter.

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Indirect immunofluorescence assay. Virus isolates in cell culture were identified initially by indirect immunofluorescence assay (IFA). Infected and uninfected cell suspensions were applied to Teflon-coated, 10-well slides; then, they were air dried and fixed in acetone. IFA was conducted with a panel of arbovirus immune ascites as described previously. The panel of antisera included broadly reactive immune ascites against Alphaviruses, specific antisera to Sindbis virus, Getah virus, Mayaro virus, Chikungunya virus, and Smiliki Forest virus, Bunyavirus broadly reactive antisera, and antisera against the flaviviruses Japanese Encephalitis virus, dengue virus, Bangkok virus, and antisera against Alphaviruses, specific antisera to Sindbis virus, Getah virus, Mayaro virus, Chikungunya virus, and Smiliki Forest virus.

RNA-polyacrylamide gel electrophoresis. The virus RNA was extracted and processed using RNA-polyacrylamide gel electrophoresis (RNA-PAGE) as previously described.

RNA extraction, complementary DNA synthesis, reverse transcription–polymerase chain reaction, and sequence analysis. Extraction of viral RNA was done with cell-culture supernatants using the QIAamp Viral RNA kit (Qiagen, Valencia, CA), and first-strand complementary DNA (cDNA) was synthesized using the Ready-To-Go You Prime First Strand Beads (GE Healthcare, Uppsala, Sweden) according to the manufacturer’s procedure. Reverse transcription–polymerase chain reaction (RT-PCR) was conducted with primers designed for *Bunyavirus genus*, Tahyna virus (TAHV), the 10th fragment of Liaoning virus, and a partial NS1 gene of *Culex pipiens pallens* densovirus (CppDNV) (Table 1). Amplified DNA fragments were visualized by electrophoresis in 1% agarose gels. Positive DNA fragments were extracted by a TaKaRa DNA fragment Purification Kit and sequenced by a commercial provider (Beijing Genome Institute, Beijing, China).

Initial sequence assembly and analysis was conducted using SeqMan (DNASTAR, Madison, WI; http://www.dnastar.com). Homology and alignment analysis was carried out using the Clustal X (version 1.8, http://www.clustal.org) and MegAlign (DNASTAR, Madison, WI; http://www.dnastar.com). MEGA 3.1 (http://www.megasoftware.net) was used for phylogenetic analysis and tree construction based on neighbor-joining (N-J) assay, and bootstrap value was 1,000.

Sera collection of local residents. Serum samples of local residents were collected from the same villages where mosquitoes were collected in Geermu city, Xining city, and Minhe county during the period from July 22 to July 27. In each site, human subjects were selected to obtain samples from several age classes by systematically visiting every other house and soliciting participants. Approximately 350–360 samples were obtained from each location. After receiving informed consent, trained personnel performed venipuncture to obtain 2-ml blood samples and recorded the age, sex, and village of residence for each subject. The separated serum samples were stored in −20°C until tested. This project was reviewed and approved by the local human subjects committee. Sera collection of local livestock. During the period from July 22 to July 27, serum samples were obtained from local livestock (cows, sheep, and swine) at the local abattoirs in Xining city and Minhe county. There were no local swine in Geermu city, so only cows and sheep were bled. The species, sex, and village or origin was obtained for each animal that was bled. The serum samples were stored in −20°C before tested.

Serological detection of antibodies against TAHV isolate QH07060. After it was determined that the isolate QH07060 from mosquitoes collected in Geermu city was a TAHV strain, this isolate was used to infect BHK-21 cells for use in the IFA assay. Infected cells were applied to Teflon-coated, 10-well slides as antigen with uninfected cells as controls. The serum samples (human, cow, sheep, and swine) were tested at a dilution of 1:50 for antibodies against QH07060 by IFA as reported previously.

Plaque-reduction neutralization tests. Seropositive samples detected by IFA were tested for neutralizing antibodies to TAHV isolate QH07060 by 90% plaque-reduction neutralization tests (PRNT<sub>90</sub>) using standard methods. Sera were tested with serial 2-fold dilutions from 1 to 5. Diluted sera were mixed with equal volumes of culture medium containing QH07060 (100 plaque forming unit [PFU]) and incubated at 37°C for 1 hour. Six-well plates were used to determine the 50% plaque-reduction end-point titers.

Table 1

<table>
<thead>
<tr>
<th>Primer</th>
<th>Amplicon region</th>
<th>Sequence data</th>
<th>Site in genome</th>
<th>Size (base pair)</th>
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<td>BUP&lt;sup&gt;56&lt;/sup&gt;</td>
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<td>ATGACTGAGTGGACTTTCAGTTTGCTGC</td>
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<td>TS141f</td>
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<td>Tahyna virus</td>
<td>CTAACCTCTATTCC</td>
<td>919–936</td>
<td>903</td>
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<td>LNV309REV&lt;sup&gt;17&lt;/sup&gt;</td>
<td>Liaoning virus</td>
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<td>1,176–1,198</td>
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<td>ATGAGTAAAGTGAGGGTCAGG</td>
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<td>DNV 3F&lt;sup&gt;18&lt;/sup&gt;</td>
<td>CppDNV&lt;sup&gt;*&lt;/sup&gt;</td>
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<tr>
<td>DNV 3R</td>
<td>CppDNV</td>
<td>CATACACATGGTCTCCTACAC</td>
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</table>

*CppDNV = *Culex pipiens pallens* densovirus.
plates of confluent BHK-21 cells were inoculated with the serum–
virus mixtures and incubated at 37°C in a 5% CO₂ incubator for
1 hour. Plates were overlaid with 3 mL of the medium containing
0.8% agarose and again with 2.5 mL of second overlay medium
containing neutral red vital stain (Sigma-Aldrich, Inc., CA)
as described previously.¹⁹ The neutralizing antibody titer was
identified as the highest serum dilution that reduced the number
of virus plaques in the test by 90% or greater. The samples were
considered to be positive when titers ≥ 20.¹⁹

RESULTS

Collection of mosquitoes. From July 30 to August 14 in 2007, a total of 8,147 mosquitoes representing six species from
three genera (Aedes, Culex, and Anopheles) were collected
in Geermu city, Xining city, and Minhe county (Table 2). In
Geermu city, 3,520 mosquitoes representing only two species
were collected (Ae. [Och.] detritus Holiday, 53.2, 1,873/3,520; Cx. pипiens Linnaeus, 46.8%, 1,647/3,520). In Xining city,
1,329 mosquitoes representing two species were collected (Ae. vexans Meigen, 96.3%, 1,280/1,329; Cx. Pipiens, 3.7%,
49/1,329). In Minhe county, 3,298 mosquitoes in four species
were collected (Cx. modestus Ficalbi, 56.0%, 1,848/3,298; Cx. Pipiens, 10.4%, 343/3,298; Ae. Vexans, 26.4%, 871/3,298). The fourth species was identified as An. nigerrimus Giles,²⁰,²¹ which constituted 1.2% of the total collection in Minhe county. This
represents the first detection of this species this far north in
China. Of the total collected in Minhe county, 6% were Aedes
that could not be identified to species.

Isolation of viruses. A total of 8,147 mosquitoes were
processed in 166 pools to isolate viruses by BHK-21 and C6/36
cells. CPE was observed in six pools. The mosquito species
producing the isolates and the location where the mosquitoes
were collected are shown in Table 3. Isolates QH07029 and QH07060 caused CPE only in BHK-21 cells and produced
no CPE on C6/36 cells. The BHK-21 cells became round by
48 hours post-infection and were lysed by 72 hours. Sucking
mice inoculated with QH07029 and QH07060 intracerebrally
showed signs of tremor and stiff neck at 24 hours and died
within 48 hours. The other four isolates caused CPE in C6/36
cells; however, no CPE was observed in BHK-21 cells, and no
illness was observed in mice. C6/36 cells infected with QH07130
became round, and they showed enlarged intercellular space
and shedding by 96 hours post-infection. The CPE of QH07022,
QH07092, and QH07150 resembled QH07130 except for formation of syncytia by 96 hours post-infection.

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Mosquito species collected in Qinghai in 2007</th>
</tr>
</thead>
<tbody>
<tr>
<td>Collection sites</td>
<td>Species</td>
</tr>
<tr>
<td></td>
<td>Ae. (Och.) detritus</td>
</tr>
<tr>
<td></td>
<td>Ae. vexans</td>
</tr>
<tr>
<td></td>
<td>Ae. sp²</td>
</tr>
<tr>
<td>Culex</td>
<td>Cx. pipiens pellens</td>
</tr>
<tr>
<td></td>
<td>Cx. modestus</td>
</tr>
<tr>
<td>Anopheles</td>
<td>An. nigerrimus</td>
</tr>
<tr>
<td></td>
<td>Total</td>
</tr>
</tbody>
</table>

*Species could not be determined.
†This is the first report of An. nigerrimus found in the northern region of China.

Identification of TAHV. Both QH07029 and QH07060 reacted with immune ascites against Bunyaviruses and prototype
TAHV (Bardos 92 strain), indicating that both isolates were TAHV or a closely related Bunyavirus. RT-PCR amplification
using Bunyavirus-reactive primers amplified the nucleotide sequences of both QH07029 and QH07060. Sequence analysis
showed the highest homology (90%) with TAHV Bardos 92 strain (U47142) by blast (http://blast.ncbi.nlm.nih.gov/),
indicating that QH07029 and QH07060 were TAHV. The S genes
of QH07029 and QH07060 were amplified, and the nucleotide
sequence homology between QH07029 and QH07060 was
99.9%. The homologies of QH07029 and QH07060 with Bardos
92 strain were 90.1% and 90%, respectively, confirming that
QH07029 and QH07060 were TAHV.

Phylogenetic analysis showed that both QH07029 and QH07060 were grouped in the same evolutionary branch
with TAHV Bardos 92 strain and were closely related to that strain, which was first isolated in the former Czechoslovakia
(Figure 2).²² This is the first time that TAHV has been iso-
dated in the Qinghai-Tibet Plateau and represents only the
second isolation in China after isolation of TAHV in Xinjiang
Province in 2006.¹²

Identification of Liaoning virus. RNA-PAGE showed that
QH07130 was a 12-fragment double-stranded RNA virus, and
the fragments could be divided into three groups, 6-5-1, which
are similar to those of Liaoning virus (LNV; NE97-31 strain);
this indicates that QH07130 was LNV (Figure 3A). RT-PCR
was done by applying primers designed for the 10th fragment
of LNV. The nucleotide sequences of QH07130 amplified showed
the highest homology with LNV (NE97-12 strain, NC007745,
98%; NE97-31 strain, NY05217, 97%); and LNV was isolated in
the northeast of China in 1999.²³ Phylogenetic analysis showed
that QH07130 distributed in the same evolutionary branch
with LNV NE97-12 strain and NE97-31 strain, and it showed the
closest relationship of QH07130 with the LNV NE97-12
strain, confirming that QH07130 was LNV (Figure 3B).

Identification of CppDNV. Nucleotides of QH07022,
QH07092, and QH07150 were extracted from supernatants and
electrophoresed in 1% agarose gels. All showed a 4k
genome, and then, RT-PCR was performed using primers
designed for NS1 genes of CppDNV. The amplified nucleotide
sequences of QH07022, QH07092, and QH07150 showed the
highest homology (more than 99%) with CppDNV (such as
JZ-16, EF579756; YN05217, EF579771; YN05169, EF579770;
XJ0545, EF579764; GZWN1, EF579757). These were members
of genus Brevidensovirus, subfamily Densovirinae, family
Parvoviridae, confirming that QH07022, QH07092, and
QH07150 were members of CppDNV.

Prevalence of TAHV antibodies in local residents. A total
of 1,078 serum samples of local residents were collected from
Of 1,078 serum samples, 19 (1.8%) were immunoglobulin G (IgG) positive. Among the 19 positive samples, 16 were collected from Geermu city where seroprevalence was 4.4% (16/366), 2 were from Xining city where seroprevalence was 0.6% (2/352), and 1 was from Minhe county where seroprevalence was 0.3% (1/360). All of the IgG positive samples came from the under 30 age groups, and no TAHV IgG positives were found in the >30 age groups in any of the locations. The highest seroprevalence was found in Geermu city in the 5–9 (7.1% positive) and the 10–14 (6.5% positive) age groups. The seroprevalence was 6.8% for the 5- to 14-year-old age group in Geermu city.

Of the 19 IgG positive samples, 18 were confirmed by PRNT with titers of 1:20 or greater. One sample produced a titer of 1:10.

Prevalence of TAHV antibodies in local livestock. A total of 240 serum samples from local livestock were collected in Geermu city, Xining city, and Minhe county. Species included cow, sheep, and swine. Of 90 cow sera, 3 were IgM positive, and 6 were IgG positive. Of 90 sheep sera, 7 were IgM positive, and 9 were IgG positive. Of 60 swine sera, 3 were IgM positive, and 2 were IgG positive.

Most of the positive livestock sera were collected from Geermu city where 13.7% (5/30) of cows and 26.7% (8/30) of sheep were positive. There were also positive sera collected from Xining city and Minhe county, but positive rates in both locations were lower than in Geermu city.

**DISCUSSION**

Although mosquito sampling was limited to one collection conducted during the summer season in each site in the Qinghai-Tibet Plateau, the results were consistent with previously published observations describing species distribution and relative abundance. A relatively low species richness is not unexpected in this area, which is characterized by a high-altitude arid steppe interspersed with mountain ranges. Our results suggest that the predominant species varied by location with *Ae. (Och.) detritus* being the most abundant species in sites we sampled in Geermu city, *Ae. vexans* being the most abundant species in sites we sampled in Xining City, and *Cx. modestus* being the most abundant species in sites we sampled in Minhe county. The difference in mosquito-species distribution is associated with the local physical geography. In Geermu city, there are numerous salt lakes, alkali flats, and marshes, including Qarham Salt Lake, the biggest salt lake in China. *Ae. (Och.) detritus* larvae can develop in high salinity, leading to the predominance of this species in this area. It is important to note that no *Cx. tritaeniorhynchus* Giles were collected in this study. This species is the major vector of Japanese Encephalitis virus in China. The absence of *Cx. tritaeniorhynchus* is consistent with the absence of Japanese Encephalitis cases reported from Qinghai.

The virus isolates obtained from mosquitoes collected in Qinghai were strains of TAHV, LNV, and CppDNV. Previous
reports indicated that few arboviruses have been isolated at elevations above 2,500 m. However, we identified TAHV, LNV, and CppDNV from Geermu city, Xining City, and Minhe County, indicating that multiple arboviruses could be isolated in Qinghai up to altitudes of 2,780 m.

TAHV, a member of California serogroup, genus Orthobu-

nyavirus, family Bunyaviridae, was first isolated in the former Czechoslovakia in 1958. TAHV is widely distributed in central Europe, and human illness from infection with TAHV has been reported as manifesting undifferentiated fever and influenza-like symptoms as well as occasionally pneumonia and central nervous system involvement. In 2006, TAHV was isolated from Cx. sp. mosquitoes collected in Xinjiang, and this was the first report of TAHV isolation in China. In this study, TAHV was isolated from Ae. (Och.) detritus collected in Qinghai, which is the second report of TAHV isolation in China, suggesting that TAHV is widely distributed in the western regions of China. TAHV has been isolated from 12 mosquito species in four genera: Ae. vexans Meigen, Ae. cinereus Meigen, Ae. caspius Meigen, Ae. punctor Kirby, Ae. communis De Geer, Ae. flavescens Mueller, Ae. excrucians Walker, Cs. annulata Schrank, Cx. modestus Ficalbi, Cx. pipiens Linnaeus, and An. hyrcanus Pallas. This is the first time that TAHV has been isolated from Ae. (Och.) detritus, suggesting that this species could serve as a vector for TAHV. In this study, the minimum infection rate of TAHV in Ae. (Och.) detritus in Geermu city was 1.07/1,000 (2 positive pools/1,873 total Ae. (Och.) detritus tested).

LNV, a member of genus Seadornavirus, family Reoviridae, was first isolated from Ae. dorsalis mosquitoes in China, and subsequent isolations from mosquitoes have been made in Xinjiang and Shanxi Provinces. Our results show the first isolation of LNV from Cx. modestus and the first isolation of LNV in Qinghai. This represents not only expansion of the vector associations of this virus but also suggests widespread distribution of LNV in China. Although experimental infections show that LNV-infected adult mice die with signs of hemorrhage, there is currently no evidence of human infection with this virus, and more investigation is needed to determine if LNV is an etiology of human disease.

CppDNV, a member of genus Brevidensovirus, subfamily Densovirinae, family Paroviridae, was first isolated from multiple mosquito species in China. The host range of CppDNV is limited to a few closely related invertebrates. There is no evidence suggesting a relationship of CppDNV with human infection or disease. In this study, three strains of CppDNV were isolated from Ae. (Och.) detritus, Ae. Vexans, and Cx. modestus that were collected in Geermu city, Xining city, and Minhe county, suggesting that CppDNV is widely distributed in Qinghai in multiple mosquito species.

The results of serological testing showed that TAHV infection in humans occurs in several areas within Qinghai and is relatively common in Geermu city, particularly Xinhua village where both of the TAHV isolates were found and where human seroprevalence was 11.6% (11/95). The investigation of local livestock indicated that TAHV antibodies (IgM and IgG) were detected in serum samples of cows and sheep with high seroprevalence in Geermu city.

The investigation also showed that IgG positive residents were all under the age of 30, and the high seroprevalence found in the 5–9 and 10–14 age groups indicates that TAHV readily infects children in Geermu city. A serosurvey in the Czech Republic showed that older residents had higher seroprevalence for TAHV antibodies, a pattern that was also seen in wild boars in the area. Such a pattern is typical for humans and animals living in a long-term enzootic focus, and the older age classes experience a higher probability of infection and a higher rate of antibodies. Our observation that seroprevalence was higher in young age groups may indicate that TAHV was recently imported into the Qinghai-Tibet Plateau, although this could not explain the lack of antibodies in older age classes. Larger sample sizes are required to resolve this question.

This investigation showed that TAHV is present in Geermu city at an altitude of 2,780 m, indicating that TAHV transmission cycles can be maintained in areas of high altitude in the
Qinghai-Tibet Plateau. In addition, this represents the first time that TAHV has been isolated from Ae. (Och.) detritus. This shows that TAHV cycles naturally and infects people in the Qinghai-Tibet Plateau, and further investigations of the distribution and public health impact of TAHV and other arboviruses are warranted in the Qinghai-Tibet Plateau.

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