La Crosse Virus in *Aedes japonicus japonicus* Mosquitoes in the Appalachian Region, United States

M. Camille Harris,¹ Eric J. Dotseth, Bryan T. Jackson, Steven D. Zink, Paul E. Marek, Laura D. Kramer, Sally L. Paulson, Dana M. Hawley

La Crosse virus (LACV), a leading cause of arboviral encephalitis in children in the United States, is emerging in Appalachia. For local arboviral surveillance, mosquitoes were tested. LACV RNA was detected and isolated from *Aedes japonicus* mosquitoes. These invasive mosquitoes may significantly affect LACV range expansion and dynamics.

La Crosse virus (LACV; family Bunyaviridae, genus Orthobunyavirus), in the California serogroup, is the major cause of arboviral encephalitis among children in the United States (1). Since its 1963 discovery in Wisconsin, LACV has been identified in 30 other US states (2). These include states within the Appalachian Mountain region (West Virginia, Virginia, Ohio, Tennessee, and North Carolina), which is an emerging focus of LACV (3).

The primary vectors of LACV, *Aedes triseriatus* mosquitoes, are present in southwestern Virginia and West Virginia, but 2 invasive congeners—*Ae. albopictus* and *Ae. japonicus*—have recently emerged (3). Both species have been shown to be competent experimental LACV vectors (4,5). Although LACV has been isolated from *Ae. albopictus* mosquitoes (6), previously it had only been detected in the Asian bush mosquito (*Ae. japonicus japonicus*) in Tennessee (7). *Ae. japonicus* mosquitoes are marmalophilic container breeders that co-occur with the primary LACV vector (*Ae. triseriatus* mosquitoes). Known to feed on humans (8), *Ae. japonicus* mosquitoes are found in woodlands (where this “rural encephalitis” virus is endemic) and urban areas (9).

To ascertain the public health risk that *Ae. japonicus* mosquito vectors represent for LACV transmission, we examined mosquitoes from West Virginia and Virginia for presence of this arbovirus. We report 2 independent isolations of LACV from adult *Ae. japonicus* mosquitoes in southwestern Virginia and 7 field detections of LACV RNA from adults (Virginia and West Virginia) and adults reared from eggs (Virginia). Our findings suggest a potential role of this invasive vector in the ecology of LACV in Appalachia (Figure 1).

**The Study**

In 2005, mosquito eggs were collected weekly in Wise County, Virginia, by using ovitraps. The resultant larvae were reared to adults in a Biosafety Level 2 insectary at 24°C, 75% relative humidity, and a photoperiod of 16 hours of light and 8 hours of dark. In 2008 and 2009, adult mosquitoes were collected weekly from infusion-baited gravid traps in Montgomery and Craig Counties, Virginia. Mosquitoes were identified to the species level according to morphology and grouped in pools of ≤50 individuals according to species, collection location, and collection date. Adults were stored at −80°C until testing. Reverse transcription PCR (RT-PCR) was used for LACV detection in mosquitoes collected in 2005 and 2008. In 2009, mosquito pools were homogenized by using previously described methods for LACV isolation (6). Homogenate supernatant (150 μL) was inoculated onto Vero cells, incubated at 37°C, and monitored daily for cytopathic effect. Isolates that showed marked cytopathic effect were harvested and submitted to the Centers for Disease Control and Prevention (CDC) in Fort Collins, Colorado, USA, and the Wadsworth Center in Slingerlands, New York, USA, for quantitative RT-PCR (qRT-PCR).

In 2013, mosquito surveillance was conducted as part of the West Virginia Department of Health and Human Resources Mosquito Surveillance Program. Gravid traps, carbon dioxide–emitting light traps, and BG Sentinel (Biogents AG, Regensburg, Germany) traps baited with octenol lures were used to collect adult mosquitoes weekly from counties with high (Nicholas, Fayette, Raleigh) and low (Kanawha, Jackson, Wood) incidence of LACV among humans as defined (10) and on a somewhat regular basis in additional counties. Specimens were pooled by species, county, and collection date for qRT-PCR at the West Virginia Office of Laboratory Services, by use of previously described methods (11).

LACV RNA was detected in a pool of *Ae. japonicus* mosquitoes collected as eggs in August 2005 from Wise

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County, Virginia. LACV was also detected in a pool of *Ae. japonicus* mosquitoes from Montgomery County, Virginia, in July 2008. LACV RNA was detected in 5 separate pools of *Ae. japonicus* mosquitoes collected in West Virginia in 2013, representing 3 counties over a 4-month period. Of 3,529 *Ae. japonicus* mosquitoes collected from Montgomery County in 2009, we isolated LACV from 1 pool (n = 3). In that same year, of 796 *Ae. japonicus* mosquitoes from Craig County tested, LACV was isolated from 1 pool (n = 50) (Table). These isolations were verified by qRT-PCR at CDC (Montgomery County isolate) and the Wadsworth Center (Craig County isolate).

Nucleotide sequencing and a BLAST (http://blast.ncbi.nlm.nih.gov/Blast.cgi) query were performed on the amplified cDNA from both isolates. The LACV medium segment was used to infer phylogeny (Figure 2). Coding sequences were aligned in Mesquite version 2.75 with Opalescent version 2.10 (12,13). Phylogenetic trees for the polyprotein genes were estimated by using a maximum likelihood–based method and assuming a general time-reversible (GTR) model with gamma-distributed rate heterogeneity of nucleotide substitution GTR + Γ in RAXML version 8.0.0 (14). Support values for each clade were generated in RAXML by using 1,000 rapid bootstrap replicates. The Virginia 2009 isolates were within the previously described lineage I, which contains the reference LACV strain isolated from the brain tissue of a child in Wisconsin (National Center for Biotechnology Information accession no. U18979) (15). The 2 isolates from mosquitoes collected in Virginia in 2009 differed by only 1 bp.

Conclusions

The isolation of LACV from field-collected *Ae. japonicus* mosquitoes, and particularly from mosquitoes collected as eggs, is highly significant because of the pervasiveness of this species in the United States. The large number of LACV detections in this invasive species highlights the need for LACV mosquito surveillance and control efforts to include *Ae. japonicus* in addition to *Ae. triseriatus* mosquito populations. Most (3/5) detections of LACV in West Virginia were in mosquitoes from Fayette County, where incidence of LACV among humans is high, suggesting that *Ae. japonicus* mosquitoes may play a major role in transmission of LACV to humans. Our detection of LACV in *Ae. japonicus* mosquitoes from field-collected eggs in 2005 and the ability of LACV to be transmitted transovarially in *Ae. triseriatus* mosquitoes suggests that future research should examine the possibility of vertical transmission of LACV in invasive *Ae. japonicus* mosquitoes. In states east of the Mississippi River, where *Ae. japonicus* mosquitoes (9) and LACV (2) co-exist, this mosquito may play a major role in the maintenance, transmission, and range expansion of LACV.

Acknowledgments

We thank Amy Lambert for conducting RT-PCR and sequencing of the 2009 isolate from Montgomery County, Virginia; Dee Petit and Andrew Luna for technical assistance; and Nate

<table>
<thead>
<tr>
<th>Collection date</th>
<th>County, state</th>
<th>Trap type†</th>
<th>Pool size‡</th>
<th>LACV detection method</th>
<th>C value</th>
<th>MLE (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2005 Aug</td>
<td>Wise, VA</td>
<td>Ovitrap</td>
<td>9</td>
<td>RT-PCR</td>
<td>38.04</td>
<td>8.59 (0.54 –41.00)</td>
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<tr>
<td>2008 Jul</td>
<td>Montgomery, VA</td>
<td>Gravid</td>
<td>22</td>
<td>RT-PCR</td>
<td>37.57</td>
<td>4.51 (0.26–22.00)</td>
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<td>2009 Jul</td>
<td>Montgomery, VA</td>
<td>Gravid</td>
<td>3</td>
<td>Isolation, RT-PCR</td>
<td>14.00</td>
<td>0.23 (0.01–1.11)</td>
</tr>
<tr>
<td>2009 Jul</td>
<td>Craig, VA</td>
<td>Gravid</td>
<td>50</td>
<td>Isolation, RT-PCR</td>
<td>23.00</td>
<td>1.28 (0.07–6.29)</td>
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<td>2013 Jun</td>
<td>Fayette, WV</td>
<td>Multiple adult</td>
<td>36</td>
<td>RT-PCR</td>
<td>37.66</td>
<td>13.41 (5.18–29.14)</td>
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<td>Cabell, WV</td>
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<td>1</td>
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<td>RT-PCR</td>
<td>37.43</td>
<td>13.41 (5.18–29.14)</td>
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</tbody>
</table>

†C, cycle threshold; LACV, La Crosse virus; MLE, maximum-likelihood estimate of the proportion of infected mosquitoes; RT-PCR, reverse transcription PCR.
‡All mosquitoes were collected from the field as adults except when ovitraps were used to collect eggs. Adults were reared from the field-collected eggs before testing for arboviruses. Multiple adult traps were gravid traps, carbon dioxide–emitting light traps, and BG Sentinel (Biogents AG, Regensburg, Germany) traps baited with octenol lures.
‡Pool size indicates no. adult mosquitoes tested for LACV.
Lambert, Allen Patton, Bonnie Fairbanks, Jennifer Miller, and Laila Kirkpatrick for field and laboratory assistance. In West Virginia, Christi Clark, Chris Boner, and Lindsay Kuncher provided laboratory assistance, and field surveillance was conducted by Kristin Alexander, Stephen Catlett, Hannah Cavender, Robert Deneer, Jennifer Beamer Hutson, Mickey King-Fowler, Dustin Mills, Daniel Payne, and Courtney Stamm. We thank the Wadsworth Center sequencing core for sequencing 1 isolate.

Mosquito surveillance in West Virginia was supported in part by a CDC Epidemiology and Laboratory Capacity for Infectious Diseases grant. Contributions of M.C.H. were supported in part by a National Institutes of Health Ruth L. Kirschstein National Research Service Award for Individual Predoctoral Fellows (no. 1F31AI080160-01A1).

Dr. Harris is a wildlife veterinarian and disease ecologist. Her primary research interest is understanding how anthropogenic environmental changes affect disease dynamics, especially zoonotic and vectorborne diseases.

References


Figure 2. Phylogeny of La Crosse virus (LACV) based on the medium (M) segment of the viral polyprotein gene. State of isolate origin, isolation year, mosquito, or vertebrate isolate source and the National Center for Biotechnology Information (NCBI) accession numbers are listed for each isolate within the tree. The scale bar represents the number of nucleotide substitutions per site. LACV historical lineages are identified by vertical bars. The 2009 isolates from Virginia (NCBI accession nos. KP226847, KP226848) group with lineage 1 viruses. Ae., Aedes; Ps., Psorophora.
La Crosse Virus in Appalachian Region, USA


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Developed with support from the CDC Foundation through an educational grant from Pfizer Inc.