PERORAL SUSCEPTIBILITY OF Aedes albifasciatus AND Culex pipiens COMPLEX MOSQUITOES (DIPTERA: CULICIDAE) FROM ARGENTINA TO WESTERN EQUINE ENCEPHALITIS VIRUS

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ABSTRACT. The transmission cycle of western equine encephalitis (WEE) virus in South America is unknown. A WEE virus strain was isolated from Aedes albifasciatus in Argentina during the WEE epizootic of 1982-83. Also, Culex pipiens from Argentina was reported to be able to transmit WEE virus experimentally, but other results indicate that Cx. pipiens from the USA is refractory to this virus. We determined the susceptibility of Argentina strains of Ae. albifasciatus and Culex pipiens complex mosquitoes to infection by WEE virus by the oral route. Adult females were fed on chicks infected with a WEE virus strain isolated in Cordoba Province, Argentina, or were fed on a blood/virus suspension. Each mosquito ingested between 10^{1.6} to 10^{6.4} vero cell plaque-forming units of virus. Each of 28 Ae. albifasciatus was positive for virus from the fourth day postfeeding, and there was evidence for virus replication. In contrast, 0/44 Cx. p. quinquefasciatus and only 1/15 Cx. p. pipiens was positive. Aedes albifasciatus is susceptible to infection by WEE virus and should be considered a potential vector of this virus in Argentina. Both subspecies of Cx. pipiens are refractory to peroral infection by WEE virus and probably do not play a role in the WEE virus cycle in Argentina.


INTRODUCTION

Western equine encephalitis (WEE) virus causes epidemics and epizootics in North and South America. In the Northern Hemisphere, Culex tarsalis and Aedes melaninomon mosquitoes are important vectors.7,8,15 In South America, severe aperiodic epizootics have been recognized since 1908 in the temperate zone of Argentina and Uruguay, although limited outbreaks in humans have been reported only in Southern Argentina.16 In these areas the natural transmission cycle is unknown.

One hundred and forty-nine thousand mosquitoes collected in Santa Fe Province during the large WEE epizootic of 1982-83, yielded four WEE virus strains from Ae. albifasciatus, Anopheles albifasciatus, Mansonia species and Psorophora pallensces13. The mosquitoes were predominantly Ae. albifasciatus (42.8%) and species of Culex (Culex) (34.4%). In the south, where human cases had occurred, 474 mosquitoes (70% Ae. albifasciatus and 30% Cx. species) were tested for virus with negative results.

Villa et al.17 reported that Cx. pipiens from Buenos Aires Province, Argentina, were able to transmit an Argentine strain of WEE virus under experimental conditions. The virus strain had undergone several laboratory passages in guinea pigs and chick embryos, and a high dose of virus in the blood meal was used. In contrast, attempts to experimentally transmit California and Washington strains of WEE virus by Cx. pipiens in the USA were unsuccessful.18 Subsequent studies showed that USA strains of Cx. pipiens were essentially refractory to peroral infection with WEE virus.8,9

Culex pipiens is represented by two subspecies in Argentina; Cx. p. quinquefasciatus occurs in Central and Northern Argentina and Cx. p. pipiens in Southern Argentina. Hybrids are present in the zone where the subspecies overlap.2,10 The objec-
tive of the present work was to determine whether Argentine Cx. pipiens complex mosquitoes and Ae. albifasciatus were susceptible to infection by WEE virus by the oral route.

MATERIALS AND METHODS

WEE virus strain:

The Cba CIV 180 strain was isolated from the brain of a sick horse in Cordoba Province during the epizootic of 1982-83 and had undergone three passages in suckling mouse brains. The virus stock was made with mouse brains harvested 24 to 48 h after intracerebral inoculation and was prepared as a 10% W/V suspension.

Mosquitoes:

Adult Ae. albifasciatus females were captured by mechanical aspirator from human bait on a farm near Córdoba City (31° 22'S, 64° 14'W), Córdoba Province, Argentina. Attempts to colonize this species in the laboratory were unsuccessful.

Female Cx. p. quinquefasciatus of the second laboratory generation were used at 14 days of age. They originated from a single oviposition from females collected in the urban area of Córdoba City. Culex p. pipiens of the second and the third generations came from two ovipositions of females collected in Puerto Madryn, (42° 50'S, 65° 05'W), Chubut Province, Argentina, and were used as 5-to 10-day old females. The subspecies were determined by examining the male genitalia of thirteen specimens from each colony. No intermediates between pipiens and quinquefasciatus were found.

Experimental Design:

Mosquitoes were infected by feeding on 1-to 12-day-old viremic chicks previously inoculated subcutaneously with approximately 10^{3.0} to 10^{3.7} vero cell culture plaque-forming units (PFU) of virus. Since few Ae. albifasciatus fed, an additional attempt was made to expose mosquitoes to infection by the oral route by using an artificial feeding technique. A heparinized-blood/virus suspension was prepared by mixing 0.5 ml of the stock virus (titer 10^{8.0} PFU/0.1 ml), 3 ml of normal guinea pig blood, and 1.5 ml of Media 199. The suspension contained 10^{3.7} PFU/μl and the mosquitoes were allowed to feed during a 2-h period. Culex pipiens mosquitoes were starved 24 h before the experiments were initiated.

Mosquitoes were allowed to feed on viremic chicks during a 9-to 15-h period at 10 to 24 h postinoculation. Serum specimens were taken from the chicks preexposure and postexposure. Blood was diluted 1:10 in a heparinized diluent and centrifuged at 4°C and 750 x g for 30 minutes. Supernatants were frozen at -80°C until tested for virus. A group of freshly fed mosquitoes also was frozen at -80°C until tested for virus and used as the sample for the first day of incubation.

The remaining fed mosquitoes were incubated in a secure room inside a laboratory with level 2 biosafety (Biosafety in Microbiological and Biomedical Laboratories, 1984), at 26°C maximum-21°C minimum for Ae. albifasciatus, 30°C maximum-24°C minimum for Cx. p. quinquefasciatus and 26°C maximum-15°C minimum for Cx. p. pipiens, in a humid atmosphere and at natural photoperiods with at least 11 h of full light.

Samples of 1 to 11 mosquitoes were frozen on different days of incubation for subsequent titration. They were triturated individually with 1 ml of diluent (Media 199, 20% fetal calf serum and gentamicin 5 mg%). Suspensions were centrifuged at 11,400 x g for 30 minutes at 4°C and supernatants were serially diluted in the same diluent and inoculated in 0.1-ml volumes into C176 vero cell culture under double agar overlay. The second agar overlay was added after 48 h, and the cell cultures were examined for 3 days and characteristic plaques were counted. Virus titers of mosquitoes are expressed as PFU/mosquito. The infection rate of mosquitoes was established taking into account the number of infected mosquitoes and the total number tested from days 3 and 4 to day 21 postinfection.

RESULTS

Culex p. quinquefasciatus fed readily and survived well. Ae. albifasciatus and Cx. p. pipiens showed low feeding rates and high mortality. A total of 41 Ae. albifasciatus, 48 Cx. p. quinquefasciatus, and 21 Cx. p. pipiens were tested (Table 1).

<table>
<thead>
<tr>
<th>Mosquito species/subspecies</th>
<th>No. feeding</th>
<th>No. exposed</th>
<th>No. fed</th>
<th>No. survived</th>
<th>Number tested for virus</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ae. albifasciatus</td>
<td>13</td>
<td>651</td>
<td>86</td>
<td>41</td>
<td></td>
</tr>
<tr>
<td>Cx. p. quinquefasciatus</td>
<td>1</td>
<td>60</td>
<td>52</td>
<td>48</td>
<td></td>
</tr>
<tr>
<td>Cx. p. pipiens</td>
<td>2</td>
<td>85</td>
<td>25</td>
<td>21</td>
<td></td>
</tr>
</tbody>
</table>

Virus titers of the infective meals (chick bloods and viral suspension) and individual Ae. albifasciatus titers are shown in Table 2. Ae. albi-
fasciatus sampled on the first day had virus titers of $10^{3.9}$ to $10^{6.6}$ PFU/mosquito. All Ae. albifasciatus tested after the fourth day were positive with virus titers of $10^{3.3}$ to $10^{6.7}$ PFU/mosquito. 

**TABLE 2**

| Cba CIV 180 WEE virus titers in the infective meals and in individual Aedes albifasciatus mosquitoes infected by peroral route. |
|---|---|---|---|---|---|---|---|---|---|---|---|---|
| Virus pre-post exposure period (log PFU/5 μl) | Virus titers (log PFU/mosquito) of individual mosquitoes per day of extrinsic incubation |
| | 1 | 2 | 4 | 5 | 6 | 8 | 9 | 11 | 13 |
| 1.6 - 2.7 | 4.4 | 5.3 | 6.7 | 6.7 | 5.8 | 6.0 | 4.6 | 6.6 | 6.6 |
| 3.9 - 3.9 | 3.9 | 5.5 | 5.7 | 6.6 | 4.1 - 4.4 | 6.3 | 4.5 | 6.5 | 4.5 | 5.6 |
| 4.7 - 4.9 | 4.5 | 5.5 | 5.9 | 5.0 - 5.4 | 5.3 | 3.5 | 5.9 |
| 4.6 - 5.7 | 4.8 | 5.6 | 5.4 - 5.7 | 5.8 | 6.5 | 6.0 | 5.8 |
| 5.2* | 5.8 | 5.3 | 5.0 | 5.0 - 5.8 | 4.9 | 3.3 | 5.2 - 6.0 | 4.8 | 5.0 | 5.3 - 6.3 | 4.6 | 5.0 |
| 3.7 - 6.4 | 5.9 | 3.3 | 5.3 |

| n | 12 | 1 | 11 | 6 | 1 | 3 | 3 | 1 | 3 | 3 | 1 | 3 |

Culex p. quinquefasciatus were fed an infective meal that titered $10^{4.3}$ to $10^{4.8}$ PFU/ml. Four freshly-fed females contained $10^{2.2}$ to $10^{5.5}$ PFU/mosquito immediately after feeding. Thereafter, none of 44 females tested on days 3 through 20 postfeeding were positive for virus. Culex p. pipiens were fed on an infective meal that titered $10^{4.5}$ to $10^{7.5}$ PFU/ml. Six freshly-fed females contained $10^{2.1}$ to $10^{4.1}$ PFU/mosquito immediately after feeding. Thereafter, only 1 of 15 mosquitoes tested on days 4 through 21 postfeeding was positive for virus. The positive female was tested on day 4 postfeeding and contained $10^{2.1}$ PFU of virus.

Using an estimated blood meal volume of 5 μl, it was calculated that each Ae. albifasciatus ingested between $10^{1.4}$ to $10^{6.4}$ PFU, each Cx. p. quinquefasciatus between $10^{2.2}$ to $10^{2.5}$ PFU, and each Cx. p. pipiens between $10^{3.1}$ to $10^{5.1}$ PFU of virus.

**DISCUSSION**

Virus titers in freshly-fed mosquitoes (day one of incubation) represent remnants of ingested virus and correlate with titers of the infective meals. The results of Villa et al. suggested that WEE virus could infect and be transmitted by Cx. pipiens after ingestion of large doses of virus ($10^{7.3}$ LD50/5 μl for 3-week-old mice). According to our results, Argentinian Cx. pipiens of both subspecies have a low capacity to become infected with Argentinian WEE virus, and the infection threshold is $>10^{3}$ PFU for Cx. p. quinquefasciatus and $>10^{2}$ PFU for Cx. p. pipiens. These titers are sufficient to infect known vector species of Culex and Aedes. The lack of isolations of WEE virus from Culex in nature during WEE epizootics in Argentina, combined with our results, suggest that Cx. pipiens does not play an important role in the WEE virus cycle in Argentina. Our results concerning the vector competence of Cx. pipiens for WEE virus agree with those from the USA.

The demonstrated susceptibility of Ae. albifasciatus to infection with WEE virus by the oral route is further evidence for considering this species to be a potential vector of WEE virus in Argentina. Aedes albifasciatus has been found naturally infected and it was an abundant mosquito during the 1982-83 epizootic. This mosquito feeds mainly on mammals, especially bovines and equines. It has been suggested that Ae. albifasciatus may serve as a vector in a WEE virus transmission cycle involving the European hare (Lepus europaeus) in Argentina. This would correspond to a similar cycle involving Aedes melanom and Lepus californicus in California. Further incrimination of Ae. albifasciatus as a vector of WEE virus must await more definitive experiments on vector competence that include determination of transmission efficiency. This is difficult to accomplish because the species has not been colonized, and it is difficult to get field-collected specimens to feed, survive, oviposit, and refeed in the laboratory.

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RESUMO: Desconhece-se o ciclo de transmissão da encefalite equina tipo oeste (WEE) na América do Sul. Uma cepa do vírus foi isolada na Argentina, durante a epizootia de 1982-1983, a partir de *Aedes albifasciatus*. Sob o ponto de vista experimental, o *Culex pipiens* da Argentina revelou-se capaz de transmitir o vírus WEE, porém outros resultados têm indicado que o *Cx. pipiens* dos Estados Unidos é refratário a esse vírus. Assim, procurou-se determinar a suscetibilidade de cepas argentinas de *Ae. albifasciatus* e complexo *Culex pipiens*, à infecção do vírus WEE por via oral. As fêmeas adultas foram alimentadas em pintos infectados com cepa do vírus isolada na Província de Córdoba, Argentina, ou então alimentadas em suspensão do vírus e sangue. Cada mosquito ingeriu entre 10^3 e 10^4 unidades virais formadoras de plaques de cultura de célula (“ vero cell”). Cada um dos 28 *Ae. albifasciatus* mostrou-se a partir do quarto dia pós-prandial e houve evidência de replicação viral. Em contraposição, 0/44 *Cx. quiquefasciatus* e apenas 1/15 *Cx. p. pipiens* revelou-se positivo. *Aedes albifasciatus* é suscetível à infecção pelo vírus WEE e deveria ser considerado vetor potencial desse agente na Argentina. Ambas subespécies de *Cx. pipiens* são refratárias à infecção por via oral e provavelmente não desempenham papel do ciclo do vírus WEE na Argentina.


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