REDUCTION OF CULISETA MELANURA FITNESS BY EASTERN EQUINE ENCEPHALOMYELOSPHISIS VIRUS

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Abstract. The traditional view of interactions between arboviruses and their arthropod vectors is that vector hosts become increasingly resistant to parasites; parasite attenuation occurs; or through the process of coevolution, resistance and attenuation occur in concert. Detrimental effects from arboviruses are only seen when vector and virus are not yet well adapted. Results from this study indicate that eastern equine encephalomyelitis (EEE) virus reduces survival and reproduction (fitness) of the mosquito Culiseta melanura, which is required for transmission of EEE virus in North America. Mosquito virulence was not measurably attenuated in virus isolates recovered 55 year apart. This virus did not affect the ability of mosquitoes to obtain a blood meal or the rate of mosquito oocyte development. Results from this study support those from earlier investigations with other mosquito-virus interactions and suggest that reproductively successful arboviruses can have detrimental effects on their mosquito vectors.

This way of thinking has been challenged on a theoretical basis. The proposition put forth is that under certain circumstances parasite virulence can be advantageous. Key to this argument is the assumption that virulence increases parasite reproductive success by increasing the efficiency of transmission.

To determine whether EEE virus affects the reproductive fitness of its enzootic mosquito vector, the consequences of virus infection on Cs. melanura survival, reproduction, and blood feeding-success were examined. Results suggest that the relationship between the mosquito vector and virus is not always benign.

MATERIALS AND METHODS

Infection of mosquitoes. The maintenance and care of experimental animals in these studies complied with the National Institutes of Health guidelines for the humane use of laboratory animals.

All experiments were conducted with a colony of Cs. melanura that were provided a 5–10% sucrose solution and maintained in environmental chambers at 25°C, 80% relative humidity, and a 16-hr light and 8-hr dark photoperiod. Five to seven-day-old adult female mosquitoes were exposed to virus by feeding on viremic chickens. Groups of mosquitoes were housed in 3.8-liter plastic cages. Individual mosquitoes were held in 0.5-liter cardboard cages.

Chicken hosts were inoculated with 10^4.5 baby hamster kidney cell (BHK) 50% tissue culture infectious doses (TCID_{50}) of EEE virus 6–24 hr prior to a 1-hr exposure to mosquitoes (Table 1). Two-tenths of a milliliter of blood was collected by venipuncture from chickens immediately before and after mosquitoes fed, mixed with 0.9 ml of avian diluent, centrifuged, and stored at −70°C until assayed for virus.

Serum was assayed for virus on monolayers of BHK cells to estimate the amount of virus imbibed by mosquitoes. Titers are expressed as log_{10} TCID_{50} per 1.0 ml of blood. No virus was detected in blood from any of the control birds.

In two experiments a small number of mosquitoes collected within 1 hr after they fed on viremic birds were collected and frozen at −70°C. To demonstrate that they had imbibed infectious virus, frozen specimens were thawed, triturated in mosquito diluent, and virus in each specimen was titrated on BHK cells.
Fecundity. The first experiment was a comparison of the number of larvae that hatched from the first rafts of eggs that were laid by virus-exposed versus control mosquitoes. Fully engorged and previously mated mosquitoes were individually maintained in cages containing water for oviposition. Within 24 hr of when eggs were laid, hatched larvae were counted using a dissecting microscope to confirm egg viability.

Oogenesis. A follow-up experiment was conducted to determine if virus affects oogenesis. Engorged mosquitoes from control and virus exposed groups were collected and frozen at 12-hr intervals for five days. Ovaries were dissected and made into wet mounts on glass microscope slides. Using bright-field microscopy, ovaries were categorized into one of Christophers’ five stages of ovarian development.23

Blood feeding success. The effect of virus on the ability of mosquitoes to obtain a blood meal, a requirement for Cs. melanura to lay eggs,4 was examined by comparing feeding success of infected versus uninfected mosquitoes. After fully engorging, groups of mosquitoes were held for 19 days and then allowed 1 hr to take a second meal from uninfected chicks. The number of fully and partially engorged mosquitoes was recorded. Mosquitoes categorized as empty were dissected and microscopically examined to determine if their posterior midgut contained a small amount of blood that was undetectable by external examination.

Mosquito survival. In a series of experiments the effect of virus on mosquito survival was examined. Mortality among groups of control and infected mosquitoes was monitored daily until all mosquitoes died. No blood meals, other than the initial one, were provided during the observation period.

Life table statistics. The effect of virus on mosquito fitness was examined by comparing life table statistics for virus-exposed versus control mosquitoes. Mosquitoes were allowed to randomly mate and then were exposed to either an EEE virus infected or control chick. Each fully engorged mosquito was held in a separate cage and provided an oviposition substrate. After laying her eggs, each female was provided a 1-hr feeding opportunity each day until she imbibed another blood meal from an uninfected chick. The number of first instar larvae that hatched from her eggs were counted as described in the methods for fecundity studies. This process of offering blood meals, allowing mosquitoes to lay eggs, and counting larvae that hatched from eggs was repeated daily until all mosquitoes died. The methods of Price23 were used to calculate life table statistics.

Statistical analysis. Statistical analyses were done using SAS.25 Survival data, expected number of daughters, reproductive expectation, and net replacement values were compared using a Kruskal-Wallace k-sample test.26 A one-way analysis of variance26 was used to analyze data on blood feeding success and fecundity. Results from the oogenesis study were compared by chi-square analysis.26

RESULTS

Mosquito infection. Virus infection rates for a subsample of mosquitoes from each experiment were not determined because 1) virus assay of whole insects requires killing mosquitoes and would reduce already small sample sizes and 2) determining infection status by removing and titering a leg might have undefined effects, potentially detrimental, on mosquitoes and would compromise the interpretation of data. Even though it is possible that not all mosquitoes in virus-exposed groups became infected, when the titer of EEE virus in avian host blood exceeds 1010 TCID50 per 1.0 ml, as it did in all experimental infections discussed below, results from past experiments show that infection rates of Cs. melanura are 100%.21,27-30 Viremias in experimental chicken hosts were similar to those previously determined for natural avian hosts.4

Fecundity. Mosquitoes that fed on the control chick produced more larvae (n = 29, χ² = 93 ± 34 (mean ± SD); degrees of freedom [df] = 1, F = 5.84, P < 0.02) than did mosquitoes that fed on the infected chick (n = 25, χ² = 73 ± 28). Titers of virus in the blood of the viremic chick (strain ME-77132) were 106.4 to 106.8 before and 106.5 to 106.8 after exposure to mosquitoes. All six mosquitoes collected and assayed for virus within 1 hr after feeding on the viremic chick contained detectable amounts of virus. The virus titer of triturated mosquitoes ranged from 104.8 to 105.8.

Oogenesis. Chi-square analysis by time of collection and stage of ovarian development revealed no significant differences in ovarian development between the two treatment groups (χ² < 0.20, P > 0.50), except for collections at 120 hr (Table 2). At that time more mosquitoes that fed on the infected chicks had laid more of their eggs than had controls (χ² = 6.4, P < 0.025). Titers of virus in the blood of the viremic chick (strain ME-77132) were 107.3 before and 108.1 after exposure to mosquitoes.

Blood-feeding success. Feeding successes of virus exposed and control groups were not different (Table 3), whether the analysis was based on mosquitoes that were fully engorged, partially engorged, contained a trace of blood in their midgut or some combination of these three categories of engorgement (F ≤ 0.12, P ≥ 0.74). Across five paired replicates of 13–63 mosquitoes that fed on different chicks, feeding rates were 67% for virus exposed versus 65% for control mosquitoes. Mean virus titers for blood from five viremic chicks (strain ME-77132) were 108.03 before and 107.5 before exposure to mosquitoes.

Effect of virus on mosquito survival. When a single strain of EEE virus was studied (ME-77132, Figure 1a), con-

<table>
<thead>
<tr>
<th>Strain</th>
<th>Passage history*</th>
<th>Isolation location</th>
<th>Year</th>
<th>Host</th>
</tr>
</thead>
<tbody>
<tr>
<td>TenBroeck</td>
<td>sm-12, Vero-2</td>
<td>Virginia</td>
<td>1933</td>
<td>Horse</td>
</tr>
<tr>
<td>ME-77132</td>
<td>sm-1, C6/36-2</td>
<td>Massachusetts</td>
<td>1977</td>
<td>Culiseta melanura</td>
</tr>
<tr>
<td>2061-88</td>
<td>C6/36-1</td>
<td>Maryland</td>
<td>1988</td>
<td>Culiseta melanura</td>
</tr>
</tbody>
</table>

* sm = suckling mice; Vero = African green monkey kidney cell line; C6/36 = Aedes albopictus cell line.
mosquitoes lived longer than those exposed to virus (df = 8, F = 2.04, P < 0.05). Titers of virus in the blood of the viremic chick were $10^{7.0}$ before and $10^{7.3}$ after exposure to mosquitoes. All six mosquitoes collected and assayed for virus within 1 hr after feeding on the viremic chick contained detectable amounts of virus. The titer of triturated mosquitoes ranged from $10^{4.3}$ to $10^{7.1}$. The number of mosquitoes in six replicates (six groups of mosquitoes that had been exposed to the same viremic chick) for the virus-exposed and three control replicates (three groups of mosquitoes that had been exposed to the same control chick) ranged from 8 to 15 and 9 to 21, respectively.

Two additional experiments were conducted to examine the effect of variation in virus strain on mosquito longevity. In the first (Figure 1b), mosquitoes that were exposed to viruses recovered 11 years apart (strains 2061-88 and ME-77132) did not live as long as controls (df = 5, F = 4.12, P < 0.002) but not different from one another (P > 0.32). Titers of virus in the blood of the viremic chicks before and after exposure to mosquitoes were $10^{6.5}$ and $10^{6.9}$ for the Ten Broeck strain and $10^{5.5}$ and $10^{6.7}$ for strain 2061-88. The number of mosquitoes in each treatment group were as follows: Ten Broeck n = 21, 21; 2061-88 n = 23, 24; and controls n = 20, 21.

**Life table statistics.** Two groups of 15 randomly mated *Cs. melanura* fed on a virus infected or on a control chick. Titers of virus in the blood of the viremic chick (strain ME-77132) were $10^{6.8}$ before and $10^{7.0}$ after exposure to mosquitoes. Expected number of daughters ($m_1$, Figure 2b; df = 1, F = 6.88, P < 0.01), reproductive expectation values ($1,m_1$, Figure 2c; df = 1, F = 8.59, P < 0.005), and cumulative reproductive expectation values (net replacement, $\Sigma_1m_1$, Figure 2d; df = 1, F = 492.66, P < 0.0001) were significantly higher for control mosquitoes than for mosquitoes that imbibed viremic blood. Differences in reproductive output were evident throughout the time that mosquitoes laid eggs (Figure 2d). The net replacement rate for control mosquitoes ($R_0 = 62$) was 82% greater than that of mosquitoes that fed on the viremic chick ($R_0 = 34$). Net replacement values should be viewed in a relative sense because they do not account for mortality of immature forms before they became reproductively competent. There was no difference in age specific survivorship between the two groups ($n$, Figure 2a; df = 1, F = 0.61, P > 0.4). Periodic decreases in survival were associated with oviposition (Figure 2a), indicating that elements connected with laying eggs are a mortality factor for *Cs. melanura*.

**DISCUSSION**

Results from this study indicate that survival and reproduction of *Cs. melanura* are reduced after the mosquito imbibes North American strains of EEE virus. Variation in virulence, as determined by significantly different survival, was not detectable among different virus strains isolated over a

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**Table 2**

A comparison of ovarian development in *Culiseta melanura* that fed on eastern equine encephalomyelitis (EEE) virus (strain ME-77132) infected versus control chicks

<table>
<thead>
<tr>
<th>Stage of ovarian development</th>
<th>Hours after imbibing blood</th>
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<tbody>
<tr>
<td>Stage 1</td>
<td>12 24 36 48 60 72 84 96 120</td>
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<tr>
<td>Stage 2</td>
<td>2 9 1</td>
</tr>
<tr>
<td>Stage 3</td>
<td>9 10 10 10 2</td>
</tr>
<tr>
<td>Stage 4</td>
<td>8</td>
</tr>
<tr>
<td>Stage 5</td>
<td>10 9</td>
</tr>
</tbody>
</table>

**Table 3**

Blood feeding success of *Culiseta melanura* that had 19 days earlier fed on an eastern equine encephalomyelitis virus infected (strain ME-77132) versus a control chick

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Number exposed</th>
<th>Fully engorged</th>
<th>Partially engorged</th>
<th>Trace blood</th>
<th>Empty</th>
<th>% fed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control-1</td>
<td>26</td>
<td>18</td>
<td>2</td>
<td>3</td>
<td>3</td>
<td>89</td>
</tr>
<tr>
<td>Control-2</td>
<td>37</td>
<td>25</td>
<td>3</td>
<td>2</td>
<td>7</td>
<td>81</td>
</tr>
<tr>
<td>Control-3</td>
<td>34</td>
<td>15</td>
<td>0</td>
<td>3</td>
<td>16</td>
<td>53</td>
</tr>
<tr>
<td>Control-4</td>
<td>63</td>
<td>26</td>
<td>2</td>
<td>0</td>
<td>35</td>
<td>44</td>
</tr>
<tr>
<td>Control-5</td>
<td>35</td>
<td>17</td>
<td>7</td>
<td>3</td>
<td>8</td>
<td>77</td>
</tr>
<tr>
<td>Total</td>
<td>195</td>
<td>101</td>
<td>14</td>
<td>11</td>
<td>69</td>
<td>65</td>
</tr>
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To determine if virulence had been attenuated over a longer period of time, a second experiment (Figure 1c) was carried out using strains isolated 55 years apart (Ten Broeck & 2061-88). Again, survival of mosquitoes exposed to the two viruses was less than that of controls (df = 5, F = 4.12, P < 0.002) but not different from one another (P > 0.32). Titers of virus in the blood of the viremic chicks before and after exposure to mosquitoes were $10^{6.8}$ and $10^{7.0}$ for the Ten Broeck strain and $10^{5.5}$ and $10^{6.7}$ for strain 2061-88. The number of mosquitoes in each treatment group were as follows: Ten Broeck n = 21, 21; 2061-88 n = 23, 24; and controls n = 20, 21.
FIGURE 1. Reductions in survival of Culiseta melanura due to infections with eastern equine encephalomyelitis virus. 

- **a**. a single virus strain. 
- **b**. two virus strains isolated 11 years apart. 
- **c**. two virus strains isolated 55 years apart.

55-year time span. Virus-induced mortality was only detected when mosquitoes were not given subsequent blood meals and, therefore, were not subject to oviposition-associated mortality. No effect of virus was detected on the ability of mosquitoes to obtain a blood meal or on the rate of oocyte development. Significantly, however, virus-associated reductions in mosquito fecundity and total fitness were observed (Figure 2). Because virus did not affect the rate of oogenesis, it was concluded that the effect on fecundity was associated with variation in the number of oocytes that developed, rather than a delay in reproductive output.

Effects of virus on mosquito survival were not apparent until approximately 14–21 days after imbibing a viremic blood meal (Figure 1). The probability of Cs. melanura refeeding on vertebrate blood and transmit virus is highest 7–10 days after imbibing an infected blood meal. Reductions in mosquito survival, therefore, occur after the probability of virus transmission is highest and would not be expected to reduce the probability of virus transmission.

Unlike effects on survival, reduction in fecundity (Figure 2) occurred within days after imbibing viremic blood, indicating that fitness reductions for EEE virus exposed Cs. melanura occur early in adult mosquito life and confer a selective disadvantage. Reduction in the number of mosquito progeny would be expected to affect the efficiency of virus transmission only if it reduced the number of potential vectors for virus progeny. This virus is transmitted horizontally and infection rates for wild mosquitoes are low (≤ 3/1,000 tested). Virulence may not have been measurably attenuated because relatively few mosquitoes in the population are virus-infected, infected vectors are not related to one another, and virus-induced reductions in mosquito survival occur after the time when the probability of transmission is highest. Results from experiments described herein did not determine whether virulence is directly related to a virus fitness advantage but suggest that such a relationship may exist. Additional research is needed to clarify the connection between mosquito virulence and efficiency of EEE virus transmission.

Results from these experiments support previous studies on mosquito-virus interactions and indicate that reproducingly successful arboviruses can have deleterious effects on their mosquito vectors. There were measurable reductions in vector fitness (Figure 2) and thus, a selective disadvantage for virus-infected mosquito hosts that are required for EEE virus transmission in North America. In earlier studies, Aedes triseriatus that were orally infected with La Crosse virus probed a vertebrate host more and imbibed less blood than uninfected cohorts. Larval Aedes dorsalis and Ae. melanimex that were vertically infected with California encephalitis virus took longer to develop to adults than did uninfected siblings. Culex pipiens that were horizontally infected with Rift Valley fever virus had a reduced ability to obtain a blood meal, to lay eggs, and to survive as adults than uninfected controls. Future investigations of arbovirus-vector interactions should consider 1) that some degree of virulence in vectors may be advantageous for the virus and 2) specifically test the assumption that virulence is associated with a virus fitness advantage.

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FIGURE 2. Effect of eastern equine encephalomyelitis (EEE) virus (strain ME-77132) on *Culiseta melanura* fitness. *a*, age-specific survivorship ($l_x$). *b*, expected number of daughters ($m_x$). *c*, reproductive expectation ($l_xm_x$). *d*, net replacement rate ($R_x$).

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REFERENCES