Transmission of the Citrus Variegated Chlorosis Bacterium
Xylella fastidiosa with the Sharpshooter Oncometopia nigricans

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ABSTRACT

Citrus variegated chlorosis (CVC) is an economically important, destructive disease in Brazil and is caused by the bacterium Xylella fastidiosa Wells. The bacterium has been found to be transmitted in Brazil by sharpshooter leafhoppers (Cicadellidae). Sharpshooters are present in most citrus growing areas of the United States. The sharpshooter leafhopper, Oncometopia nigricans Walker, is commonly found and an experimental citrus vegetation where sharpshooters were common (6). Various strains or types of X. fastidiosa exist (17,23) and cause many different diseases of commercial crops (24), one of which, Pierce’s disease of grapevines, is well known in California (13,14) and has long been a limiting factor for grape production in Florida (28). Pierce’s disease of grapevines presently is causing increasing losses in California since the introduction of the glassy-winged sharpshooter, Homalodisca coagulata Say (21). CVC causes considerable economic losses in Brazil. In 2000, it was estimated that in the state of Sao Paulo, Brazil, 34% of the 200 million trees have symptoms of the disease and, in 2001, the number of symptomatic trees increased to 36%, with 24% having severe symptoms of leaf chlorosis and small fruit (2). Precise economic figures are not available; however, yields are definitely reduced and the small fruit are usually unmarketable (A. J. Ayres, Fundecitrus, Brazil, personal communication). The bacterium is transmitted by xylem-feeding sharpshooter leafhoppers (Homoptera: Cicadellidae, Cicadellinae; 11,12). Of 16 sharpshooter species tested in Brazil, 12 have been identified as vectors of CVC (16,25). Sharpshooters are present in Florida and feed on citrus (3,4); therefore, we initiated studies on the ability of a sharpshooter from Florida, Oncometopia nigricans Walker, to transmit the CVC bacterium. The methods developed in this study are now being applied to the glassy-winged sharpshooter H. coagulata, another sharpshooter species native to Florida. This information is needed to assess the threat posed by CVC disease to the citrus industries in the United States. Portions of this work have been presented as abstracts previously (5).

MATERIALS AND METHODS
O. nigricans sharpshooters were collected from a Lantana sp. (Lantana camara (L.) Moldenke) in central Florida. The Lantana sp. was planted as a trap crop in field plots between an area of native vegetation where sharpshooters were commonly found and an experimental citrus planting. During the months of August to October, adult sharpshooters were captured from the Lantana plants and put into screened cages which contained L. camera, ‘Madame Vinous’ sweet orange (Citrus sinensis L. ‘Madame Vinous’), grape (Vitis vinifera L. ‘Mission’), and periwinkle (Catharanthus roseus L. (G. Don)). The sharpshooters were allowed to feed on this mixture of plants for a period of 5 days, after which they were put into small plastic cages with cuttings of healthy periwinkle for overnight shipment to the United States Department of Agriculture-Agricultural Research Service, Foreign Disease-Weed Science Research Unit, Fort Detrick, MD. The original plants that the sharpshooters were caged on were observed for any symptoms of Pierce’s disease and were tested for the presence of X. fastidiosa using membrane entrapment immunofluorescence (MEIF) (7) after 6 months.

After arrival at the Fort Detrick lab, the sharpshooters were placed on healthy sweet orange plants for 24 h of acclimation, then transferred onto CVC-infected sweet orange (Citrus sinensis L. ‘Madame Vinous’). Symptomatic source plants had been previously inoculated by grafting buds from CVC-infected sweet orange samples received from Brazil. Source plants were confirmed positive for X. fastidiosa by polymerase chain reaction (PCR)-based assays (18) prior to being used in these experiments. After an acquisition access period (AAP) of 24 to 48 h, surviving sharpshooters were transferred to healthy ‘Madame Vinous’ sweet orange plants for a 48- to 72-h inoculation access period (IAP). Initial experiments were conducted with groups of 29, 47, 48, 50, 51, 54, and 57 sharpshooters that survived and fed on CVC-infected plants and then were transferred to single sweet orange plants. In secondary experiments, groups of 1, 3, 5, 10, and 20 sharpshooters were fed on CVC-infected plants for 24 to 72 h and then were placed on healthy sweet orange plants. The minimum IAP used was 12 h but, in some experiments, sharpshooters survived for up to 7 days.

The sweet orange plants were maintained in the greenhouse and allowed to grow. Assays were performed using PCR with specific primers designed for the CVC strain of X. fastidiosa (18). PCR-based assays were performed on the plants...
3 months later and at various time periods afterward. In the initial experiments, leaf petioles were assayed by immunocapture of the target bacteria followed by nested-PCR, as described by Pooler et al. (19). In subsequent experiments, petioles were assayed without immunocapture by standard PCR using primer pair 272-1-int and 272-2-int (18). In these latter experiments, petioles were sliced and then extracted using the FastPrep system (Qbiogene, Carlsbad, CA). Leaf petioles also were assayed using MEIF designed for detection of the citrus canker bacterium Xanthomonas axonopodis pv. citri and the citrus bacterial spot bacterium X. axonopodis pv. aurantifolia (7).

### RESULTS AND DISCUSSION

#### Plant assays.
No symptoms indicative of *Xylella fastidiosa* caused diseases such as Pierce’s disease of grape, periwinkle wilt, or CVC were found on any of the plant materials that the field-collected sharpshooters fed on prior to transmission experiments. MEIF assays of these materials were negative. Controls using the CVC-inoculated and symptomatic sweet oranges that were previously PCR positive also were positive in MEIF tests. These plants were used as CVC source plants for sharpshooter acquisition.

#### Transmission results.
Fifty-nine sweet orange plants were infested with 1 to 57 *O. nigricans* for a minimum of 24 h of IAP and presumed to be inoculated. However, the overall transmission rates were 20.3% (12 of 59 plants were positive by PCR) and were scattered among insects sets (Table 1). The number of insects did not affect the rate of transmission as much as the length of active feeding. Single insects were good vectors if they were actively feeding during the acquisition and inoculation periods. Initially, transmissions were obtained with high numbers of sharpshooters (47 to 57). These results were obtained even though a limited number of tests were done (Fig. 1) using relatively large plants in three-gallon pots. Successful transmissions also were obtained with 10, 5, 3, and 1 sharpshooters. Of the plants used for single sharpshooter transmissions, 20% tested positive. More transmission tests were conducted using single sharpshooters because small plants could accommodate from 1 to 5 sharpshooters, whereas larger plants had to be used when 10 or more sharpshooters were used. When using high numbers of sharpshooters, from 47 to 57, the bacterium was transmitted to 50% of the test plants. We found that small plants that are actively growing are more susceptible to infection by *X. fastidiosa* than are larger plants, and the bacterium is more easily detected in such plants. This could contribute to the relatively high transmission rates observed in the single insect experiments.

Sharpshooters may acquire and transmit the bacterium without a latent period or without the bacteria colonizing the precibarium or cibarium of the insect. Sharpshooters lose the ability to transmit colonized bacteria after molting (20).

Symptom expression of CVC in greenhouse-grown citrus plants is not easily obtained (10). Even with graft-inoculated or culture-inoculated plants, the expression of the typical chlorotic leaf spots and the variegated chlorosis symptoms seen in field plants often are missing. We have seen typical symptoms vanish in subsequent plant growth flushes. Many of the plants used in sharpshooter transmission

#### Table 1. Transmission of citrus variegated chlorosis to ‘Madame Vinous’ sweet orange by the sharpshooter leafhopper, *Oncometopia nigricans*.

<table>
<thead>
<tr>
<th>No. of sharpshooters per receptor plant</th>
<th>No. of tests</th>
<th>Positive transmissions</th>
<th>Transmission (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>47–57</td>
<td>6</td>
<td>3</td>
<td>50</td>
</tr>
<tr>
<td>20–29</td>
<td>4</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>10</td>
<td>7</td>
<td>2</td>
<td>28</td>
</tr>
<tr>
<td>5</td>
<td>11</td>
<td>1</td>
<td>9</td>
</tr>
<tr>
<td>3</td>
<td>7</td>
<td>1</td>
<td>14</td>
</tr>
<tr>
<td>1</td>
<td>24</td>
<td>5</td>
<td>20</td>
</tr>
<tr>
<td>Totals</td>
<td>59</td>
<td>12</td>
<td>20.3</td>
</tr>
</tbody>
</table>

* Positive by nested polymerase chain reaction (PCR)-based assay (20) or by standard PCR using primer pair 272-1-int and 272-2-int (19).

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**Fig. 1.** Transmission of *Xylella fastidiosa* to sweet orange ‘Madame Vinous’ by the leafhopper *Oncometopia nigricans*. Detection of the specific polymerase chain reaction (PCR) amplification product after immunocapture and nested PCR amplification (20). Upper half gel: Lanes 1–4, healthy plant controls; lanes 5–6, infected source plants; lane 7, negative control; lanes 8–9, bacterial culture positive controls. Lower half gel: Seven test plants representing four separate experiments denoted A, B, C, and D. Lane M is a 100-bp DNA ladder (Life Technologies, Rockville, MD).
work showed good symptoms which often disappeared in the subsequent months. Other plants showed a mild chlorosis that was unlike the symptoms described for CVC. The reason for this is unknown, but it may be that full symptom expression requires nutrient or water stress, which were not present during the course of the experiments reported herein. Symptom expression was inconsistent; therefore, PCR assays specific for the CVC bacterium were a better indication of infection.

In Brazil, 12 of the 16 species of sharpshooters tested have been shown to transmit the CVC Xylella sp. to citrus (15). Most species transmitted at an efficiency rate of lower than 5%; however, two species, Macugonalia leucomelas and a Bucephalogonia sp., transmitted at rates of 11 and 17%, respectively. These transmission rates are similar to our results with O. nigricans (Table 1).

The glassy-winged sharpshooter, H. coagulata, was introduced into California around 1990 and was identified in 1994 (1). This sharpshooter has continued to expand its numbers and range in California (16, 21). It is now found in most of Southern California and has been found south of Bakersfield and in Sacramento. It was found to transmit the X. fastidiosa strain that causes oleander leaf scorch disease (22) and, since then, has been found to transmit the X. fastidiosa strain that causes Pierce’s disease of grape (21). Large populations of this sharpshooter are found in many parts of California, feeding and laying eggs on various types of citrus (lemon and orange) and other host plants (macadamia, apricot, oak, and others). The ability or efficiency of this sharpshooter to transmit the bacterium that causes CVC is unknown. Experimental work is currently underway to determine this. Given the high populations of this insect in California citrus, if the glassy-winged sharpshooter is shown to transmit the CVC strain of X. fastidiosa, the threat of establishment of the CVC strain should be considered serious. Coffee leaf scorch disease recently has been reported in Costa Rica (26). This may mean that CVC also is present because there is a close relationship between CVC and coffee leaf scorch diseases and their causal bacterium.

The ability of the O. nigricans sharpshooter to vector the CVC strain of X. fastidiosa is important to Florida citrus. Should the causal agent of CVC be introduced into Florida, a vector is already present that will spread this important bacterial pathogen.

**LITERATURE CITED**