ABSTRACT
We are studying the effect of host plant tolerance on insect vector acquisition of Xylella fastidiosa (Xf) from plants tolerant, moderately susceptible, and highly susceptible to Xf infection. We are observing Xf population and distribution in tolerant and susceptible plants, and its relationship to xylem anatomy, symptom development, and bacterial acquisition by sharpshooters. Since host plant resistance is an important component in the long-term goal of curing PD, it is important to know how resistant plants affect PD spread in areas permanently infested with sharpshooter vectors. We also address the short-term goal of controlling PD spread by comparing grape cultivars in their ability to provide inoculum for vine-to-vine spread of Pierce’s disease. Anatomical comparisons of three cultivars, Sylvaner’, ‘Cabernet Sauvignon’ and ‘Pinot Noir’ showed that all three varieties had similar numbers, lengths and distributions of vessels. The only significant difference was that tolerant ‘Sylvaner’ had ~ 20 % more rays than the more susceptible ‘Cabernet Sauvignon’ or ‘Pinot Noir’ (n = 25, P = 0.01) in canes of similar age, length and diameter. In all four alternate hosts, morning glory (Ipomoea purpurea), mugwort (Artemisia douglasiana), sunflower (Helianthus annuus) and annual bur-sage (Ambrosia acanthicarpa), the longest vessels measured were less than 13 cm long, while in grapes the longest vessels averaged 62 cm. Though alternate hosts had various vascular morphologies and stem lengths, all had shorter vessels than grapes. Blue-green sharpshooters failed to efficiently inoculate wild-type Xf and green fluorescent protein-expressing (GFP) Xf into both grapes and alternate hosts; only one of 44 grapes inoculated with BGSS became infected. In order to generate GFP-Xf infected plants for microscopy, we are mechanically inoculating alternate hosts and grapes. Ongoing work focuses on refining microscopic techniques to visualize small numbers of Xf in plant stems, and generating large numbers of Xf infected grapevines to serve as new sources for sharpshooter bacterial acquisition.

INTRODUCTION
Alternate hosts of Xf were selected for their different patterns of Xf colonization after vector inoculation, lack of stem lignification, varying morphology, and absence of green autofluorescence under blue light. In previous experiments, Xf-carrying sharpshooters infected morning glory and sunflower more than 80% of the time. Xf spread systemically throughout both plants and reached populations over 10^5 colony-forming units (CFU)/gram. Quinoa and mugwort were less-frequently infected (32% and 16%, respectively) by Xf and supported lower bacterial populations (10^3 CFU/g for quinoa, 10^4 CFU/g for mugwort). Xf moved systemically to a limited extent in quinoa, but not in mugwort (8, 16). Grape cultivars with varying tolerance to PD selected for evaluation are tolerant ‘Sylvaner’, moderately susceptible ‘Cabernet Sauvignon’ and highly susceptible ‘Pinot Noir’ cultivars of Vitis vinifera (12, 13). Both blue-green sharpshooters (BGSS) and glassy-winged sharpshooters (GWSS) will be used to infect plants and assess the efficiency of insect acquisition of Xf (1, 7, 11).

We are using wild type and transformed isolates of Temecula Xf in our experiment. The transformed isolate continually expresses green fluorescent protein (GFP) when illuminated with blue light. GFP-Xf was transmitted by blue-green sharpshooters, retained typical virulence in grape, and allowed examination of plant tissues without the extensive fixation required with electron microscopy. With confocal microscopy, GFP-expressing Xf can be observed in small and large colonies in vessels, and passing through bordered pits between vessels in symptomatic ‘Cabernet Sauvignon’ petioles (10).

Anatomical comparisons between alternate hosts and grape cultivars included measurements of vessel length and number, and vascular bundle number and distribution based on the techniques of Tyson et al. (15), and Ewers and Fischer, modified to infuse the pigment via 100kPa pressure applied to the proximal end of the cutting (5). We evaluated primary vegetative growth rather than secondary xylem due to the difficulties in sectioning, culturing from, and feeding BGSS on partially lignified stems. GFP-Xf inoculation and colonization of all plants will be measured similarly in all plants: groups of four GFP-Xf carrying sharpshooters inoculated a 3-cm stem section, and the plants were evaluated for the presence of GFP-Xf approximately 8 weeks after inoculation. Colonized vessels will be counted, and populations estimated by culture on PWG media (2, 8).
We will measure Xf acquisition by sharpshooters from the alternate hosts and grape cultivars after completing the anatomical comparisons. Insects will be caged on Xf inoculated sites for 4 days to acquire the bacteria, and then be placed on another grape seedling for 2 days to determine their acquisition efficiency. Immediately following sharpshooter feeding, the stem site will be examined with confocal microscopy and tested with culture. Three stem cross-sections and three 1-cm long longitudinal sections per site will be sectioned and suspended in 50% glycerol on a depression slide. When illuminated with blue and ultraviolet light, both GFP-Xf and the individual vessels are visible, and it is possible to determine the proportion of vessels colonized, the extent of bacterial colonization inside them, and the distribution of colonized bundles. Bacterial populations will be determined by culture from remaining plant material of the same site, and symptom development and severity will be assessed. Since acquisition efficiency has been related to bacterial populations (9), we must separate the effects of bacterial distribution and proportion of colonized vessels from the effect of bacterial population. The number of plants we can evaluate via microscopy is a limiting factor. A maximum of 90 observations per experiment will allow examination of 5 inoculation sites for each of three species or cultivars, which should enable detection of a 20% difference in Xf colonization (α = 0.05 and β = 0.10) (14).

OBJECTIVES
1. Describe the bacterial colonization of asymptomatic weed species and grape varieties of varying tolerance to Pierce’s disease using an Xf strain that continuously expresses green fluorescent protein.
2. Determine the relationship between the pattern of colonization of a plant by Xylella fastidiosa (Xf) and the ability of that plant to be a source for bacterial acquisition by sharpshooter vectors.

RESULTS
There were no differences in the total vessel number, the proportion of short vessels, or the longest vessels between resistant and susceptible grape varieties between greenhouse-grown canes of similar length, age, and diameter. The longest vessel measured by paint infusion was 110 cm (Pinot Noir), although most vessels were less than 12 cm long in all cultivars (Figure 1). Cane length had a small but significant influence on longest vessel ($r^2 = 0.20; P = 0.02, n = 27$), but did not relate to the number of very short vessels. There was no relationship between stem length and vessel length in the other plants.

While more replication is needed, the longest vessel measured in any alternate host was 15 cm long (mugwort). In sunflower, 71% of vessels were less than 3 cm long. Other species had a wider range of vessel lengths, with about half their vessels less than 3 cm long (Figure 2). Mugwort had roughly twice as many vessels (592, $n = 3$) at the stem base than morning glory (217), quinoa (251) or sunflower (286) stems of comparable diameter and age. Sunflower, mugwort and quinoa all had vascular tissues in evenly distributed bundles wide interfascicular regions of parenchyma (4). Annual morning glory had large vessels distributed evenly along the cambium.

Table 1: Comparisons between canes of similar length, age, and diameter belonging to 3 grape cultivars.

<table>
<thead>
<tr>
<th>Cultivar</th>
<th>Total # vessels at base of cane (SE)</th>
<th>% Vessels &lt; 3 cm (SE)</th>
<th>Longest vessel (SE)</th>
<th># Rays (SE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cabernet Sauvignon</td>
<td>515 (43)</td>
<td>21 (3)</td>
<td>53 cm (5)</td>
<td>34 (1)</td>
</tr>
<tr>
<td>Pinot Noir</td>
<td>474 (27)</td>
<td>20 (3)</td>
<td>64 (9)</td>
<td>34 (2)</td>
</tr>
<tr>
<td>Sylvaner</td>
<td>514 (38)</td>
<td>18 (5)</td>
<td>69 (9)</td>
<td>40 (2)</td>
</tr>
<tr>
<td><strong>one-way ANOVA</strong></td>
<td>(n = 27 , $P = 0.67$)</td>
<td>(n = 27 , $P = 0.84$)</td>
<td>(n = 27 , $P = 0.35$)</td>
<td>(n = 27 , $P = .01$)</td>
</tr>
</tbody>
</table>

Figure 1: Vessel length distribution in greenhouse-grown Pinot Noir (PN), Sylvaner (SYL) and Cabernet Sauvignon (CS).
Blue-green sharpshooters failed to efficiently inoculate \( Xf \) into both grapes and alternate hosts in three separate attempts from 7/03 to 4/04; only one of 44 grapes became infected. Though the \( Xf \)-infected source plants had fully developed symptoms and were positive for \( Xf \) by culture, there may have been nutritional or physiological factors that prevented them from being good sources of bacterial acquisition. We are mechanically inoculating alternate hosts and grapes to generate GFP-\( Xf \) infected plants for microscopy practice. Because the distribution of \( Xf \) in an insect-inoculated stem is likely different from a mechanically inoculated stem, we still plan to use insect-inoculated plants when we compare sharpshooter acquisition and bacterial distribution in alternate host stems. Ongoing work focuses on refining microscopic techniques to visualize small numbers of \( Xf \) in alternate host stems, and generating large numbers of \( Xf \)-infected grapevines to serve as new sources for sharpshooter bacterial acquisition.

**CONCLUSIONS**

Three things are required for the development of Pierce’s disease in grape: the pathogen *Xylella*, a sharpshooter insect vector, and a susceptible plant host. We are systematically examining the interactions between plants and the pathogen, and the role that host resistance plays in the ability of the vector to acquire \( Xf \) and spread Pierce’s disease. The only significant difference between grape varieties was that tolerant ‘Sylvaner’ had approximately 20% more rays per stem compared with susceptible ‘Cabernet Sauvignon’ or ‘Pinot Noir’. In grapes, rays are composed of dense parenchyma cells, without tracheids or vessels, and separate the water-conducting xylem into longitudinal zones (3). Perhaps this limits the lateral spread of \( Xf \) to the zone it is originally inoculated into. While additional work is needed, the vessels of other hosts were approximately 75% shorter than vessels of grapes, limiting the passive spread of \( Xf \) via xylem sap movement, and are found in bundles separated by parenchyma cells, which may also limit the lateral spread of \( Xf \). Additionally, it is likely \( Xf \) movement between bordered pits is an active process (10); anatomical and biochemical differences in pit structure may explain differences between cultivar susceptibility to \( Xf \).

In grapes, electron and confocal microscopy showed \( Xf \) densely packed in individual vessels, with adjacent vessels empty or containing a few cells (10, 15). Alternate hosts or tolerant grape cultivars with low overall populations may have just a few vessels with bacteria, so acquisition would be highly variable and dependant upon sharpshooters encountering the few colonized vessels while feeding. In symptomatic grape petioles, 13% of vessels were colonized to some extent with GFP-\( Xf \), though only 2.1% of all vessels were completely blocked with bacteria (10). Though it is not known how many probes a sharpshooter makes in a given feeding session, glassy-winged sharpshooters can generate multiple salivary sheaths in one insertion, adjacent to vessels and xylem parenchyma cells (6). Sharpshooter acquisition of \( Xf \) increased along with bacterial populations in infected grapes (9), and a similar positive relationship is expected if the proportion if colonized vessels increases insect acquisition of *Xylella*.

**REFERENCES**


**FUNDING AGENCIES**

Funding for this project was provided by grants from the Viticulture Consortium, the California Agricultural Experiment Station (at College of Natural Resources, University of California, Berkeley), and the CDFA Pierce’s Disease and Glassy-winged Sharpshooter Board.