FUNDING AGENCIES
Funding for this project was provided by the CDFA Pierce's Disease and Glassy-winged Sharpshooter Board.

Table 1. Summary of physiological measurements for water relations and photosynthesis.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>plants</th>
<th>( \Upsilon_p )</th>
<th>( \text{A}_{\text{max}} )</th>
<th>( g_s )</th>
<th>( C_i )</th>
<th>( E )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Non-inoculated</td>
<td>6</td>
<td>0.27</td>
<td>25.11</td>
<td>0.2612</td>
<td>1239.17</td>
<td>4.34</td>
</tr>
<tr>
<td>Mock-inoculated</td>
<td>6</td>
<td>0.317</td>
<td>25.75</td>
<td>0.2211</td>
<td>1188.34</td>
<td>3.79</td>
</tr>
<tr>
<td>( Xf )-inoculated</td>
<td>8</td>
<td>0.49</td>
<td>16.22</td>
<td>0.0721</td>
<td>817.00</td>
<td>1.50</td>
</tr>
<tr>
<td>Mild stress</td>
<td>7</td>
<td>0.434</td>
<td>19.98</td>
<td>0.2196</td>
<td>1245.00</td>
<td>3.78</td>
</tr>
<tr>
<td>Double treatment</td>
<td>8</td>
<td>0.583</td>
<td>12.14</td>
<td>0.0261</td>
<td>556.5</td>
<td>0.64</td>
</tr>
</tbody>
</table>

\( \Upsilon_p \): pre-dawn water potential (-MPa), \( \text{A}_{\text{max}} \): net assimilation (mmol CO\(_2\)/m\(^2\)/s) (measured at saturating CO\(_2\) and light), \( g_s \): Stomatal conductance (umol H\(_2\)O/m\(^2\)/s), \( C_i \): Internal CO\(_2\) concentration (mmol CO\(_2\)/mol air), \( E \): transpiration rate (mmol H\(_2\)O/m\(^2\)/s).

Figure 1. 2-Dimensional hierarchical cluster analysis of 24 microarrays from the moderate drought stress condition. 238 transcripts were identified with a minimum of 2-fold induction and a T-test score of \( a=0.05 \). Red = increased expression; Blue = decreased expression; White = no change in expression. I=infection; D=drought; N=healthy; E= prior to symptom development; L=subsequent to symptom development.

Figure 2. In situ localization of candidate gene 8946. Note intense staining in phloem and xylem associated pyrenchyma, indicating \textit{Xylella}-induced gene expression in living tissue adjacent to differentiated xylem.
**V. Osmotic stress and cell wall modifying enzymes.**

Eleven *Xylella*-associated transcripts have predicted roles in cell wall metabolism (e.g., expansins, enzymes involved in pectin degradation and pectin modification) or osmotic stress (e.g., galactinol synthase, dehydrin proteins and several aquaporins). These genes were induced an average of 5-fold in *Xylella* infected tissues. None of these 11 genes were up-regulated in response to water stress alone, and only the dehydrin and galactinol synthase genes showed evidence of synergy between *Xylella* and drought stress. Cell wall modification genes (expansins, pectin esterases, pectate lyases, polygalacturonases, etc.) were among the major class of water stress repressed genes in *Arabidopsis* (Bray, 2004). In the current study, these genes were induced by the pathogen, providing a possible counterpoint to the argument in favor of *Xylella*-induced drought stress.

**CONCLUSIONS**

In summary, a wide array of genes are up regulated (or in some cases down regulated) in grapes in response to *Xylella* infection. We found limited correlation between the nature of genes induced by moderate drought stress and the genes induced by pathogen infection. Interestingly, however, the results suggest a synergistic effect of drought stress on *Xylella*-induced gene expression. We have also identified numerous genes where induction was specific to the pathogen, and not synergistic with drought. This later class of genes included pathogenesis related protein genes and genes involved in plant cell wall metabolism. Ongoing experiments are using Real Time PCR to validate and extend the Affymetrix GeneChip results (data not shown) and to determine the spatial pattern of gene expression for the various classes of transcriptional response, as shown by example for gene 8946 in Figure 2.

Our earlier work with a small set of pathogen-induced genes has permitted us to characterize the kinetics and specificity of the host response to *Xylella*, and to isolate and begin the characterization of *Xylella*-responsive gene promoters. The recent results, reported above, provide a large suite of new genes and predicted biochemical pathways for investigation. We suggest that these results are a first step toward a comprehensive understanding of host responses to PD, and the relationship of disease to whole plant physiology including water relations, photosynthesis and defense responses. Our continuing work will explore in detail the relationship between gene expression in resistant and susceptible plants, and to begin more precise analysis of the spatial relationship between gene expression and pathogen localization. Moreover, we anticipate providing many additional and potentially useful gene promoters to Dave Gilchirst's project to develop a pathogen-inducible transgene system. How will these technologies help in solving PD? In the short term they will:

1. Provide gene-promoters for effective genetic engineering in grapes.
2. Inform us about the nature of host responses to *Xylella* infection.
3. Allow pathogen detection based on Real Time PCR using a "biomarker" strategy.
4. In the long term, transcriptional profiling will identify candidate genes and gene pathways that may confer resistance to the pathogen (Xf).

Other strategies, such as reverse genetics and analysis of natural genetic variation, will be needed to establish a causal role for candidate genes.

**REFERENCES**


Analysis of Microarray (SAM) Data (Aubert et al., 2004). Differential regulation was assessed by comparison to uninoculated control plants grown under identical conditions.

In total, 238 genes were identified as being differentially expressed (T-test \( a < 0.05, \geq 2\)-fold induction) in response to \( Xf \) treatment or drought stress (Figure 1). There are 2 primary conclusions from this study: First, we have identified several genes where expression is induced strongly in diseased tissue and where drought stress does not appear to impact this transcriptional response. The majority of such genes have predicted roles in defense and cell wall metabolism. Second, a large fraction of the \( Xylella \)-induced transcriptome is synergistically modified in plants that are doubly-treated by pathogen infection and moderate drought stress. These genes fall into two categories: synergistically upregulated are primarily from the flavonoid biosynthesis pathway, while synergistically down regulated are primarily from the photosynthesis pathway. These results are consistent with the existence of two distinct classes of transcriptional response in grapes to \( Xylella \). One response is sensitive to plant water status and results in redirection of flavonoid synthesis and photosynthesis genes, and one response is independent of plant water status leading to the activation of defense-related transcripts. Although we observed limited overlap in the genes induced in response to moderate drought stress and the genes induced in diseased tissue, we cannot rule out the possibility that a more severe drought stress may lead to an increase in the coincidence of PD and drought-associated gene expression.

As shown in Table 1, physiological measurements of the plants used for microarray analysis also suggest an additive interaction between water stress and PD. We note that the level of water stress imposed in these experiments induced an acclimation response in treated plants, as evidenced by measurements of stomatal conductance, internal \( CO_2 \) concentrations and transpiration rates. However, reductions to pre-dawn water potential and net assimilation rates document a clear water stress response. By contrast, pathogen infection had a strong influence on virtually all of these parameters. Moreover, drought stress combined with pathogen infection tended to increase the magnitude of change in all parameters assayed. These results suggest a reduced capacity for acclimation to water stress in infected plants and they agree well with the results of gene expression, described below.

A 2-Dimensional hierarchical cluster generated with the DChip software (Li and Wong, 2001) was used to depict the expression 238 genes that were responsive to one or more of the treatments. The most striking aspect of this particular analysis is the massive transcriptional response that occurs in infected and symptomatic plants. Major categories and/or expression patterns of genes identified so far are described briefly below.

I. Disease related gene expression.
Seventeen transcripts were annotated as disease related genes, including many pathogenesis related or PR protein genes. On average these genes were up regulated 7-fold in response to pathogen infection. Expression of these genes was not influenced by drought either in healthy or diseased plants. The sole exception are two PR protein genes that were down regulated 2.5-fold in response to drought stress, but up regulated >10-fold in response to the pathogen. These results suggest the occurrence of a pathogen-specific defense response in susceptible \( V. \ vinifera \).

II. Photosynthetic gene expression.
One of the most common responses of plants to drought stress is a down regulation of photosynthesis. Consistent with physiological measurements, 11 photosynthesis-related transcripts were significantly down regulated in \( Xylella \)-infected plants. While moderate water stress had little or no effect on expression of these genes, the combination of pathogen infection and water stress resulted in an even greater reduction in gene expression compared with either treatment alone. \( Xylella \) causes a decrease in photosynthetic gene expression that is accentuated by reduced water availability.

III. Flavonoid pathway gene expression.
The largest transcriptional effect of \( Xylella \) infection was a massive re-direction of enzymes and regulatory proteins for flavonoid biosynthesis. In total, 27 genes were 4-fold upregulated in \( Xylella \) infected plants, compared to healthy control plants. Approximately 50% of these transcripts were induced an additional 2.5-fold when drought stress and \( Xylella \) infection were combined. The transcription of flavonoid pathway genes was not significantly affected by drought stress alone. \( Xylella \) causes an increase in flavonoid gene expression that is accentuated by reduced water availability.

IV. Genes induced uniquely in the interaction between disease and drought.
Twelve genes were unaffected by either drought or \( Xylella \) infection, but were significantly induced in plants that were challenged with both \( Xylella \) and water stress simultaneously. On average, these genes were induced 3.5-fold in double-treated plants. Annotations for these genes do not suggest function in a common pathway.
provide the basis for new lines of experimental inquiry focused on testing the efficacy of specific host genes for PD resistance. It should be possible, for example, to determine the extent to which resistance responses in grapes are related to well-characterized defense responses in other plant species (e.g., Maleck et al., 2002; Tao et al., 2003; de Torres et al., 2003).

Three co-lateral benefits from the identification of pathogen-induced genes are: (1) the promoters for such genes are candidates to control the expression of transgenes for resistance to PD, (2) the protein products of induced genes may have roles in disease resistance, and (3) knowledge of host gene expression can be used to develop improved diagnostic assays for disease. In a related project, we are currently characterizing pathogen-responsive promoters, which will facilitate testing of candidate genes for resistance phenotypes.

OBJECTIVES
1. Identify genes and gene pathways in susceptible *V. vinifera* correlated with *Xf* infection: (a) identify *Xylella*-responsive genes in *V. vinifera*, (b) distinguish early from late gene expression, and (c) determine the correlation between drought stress and PD.
2. Determine host genotype affects on gene expression in response to *Xylella* infection: (a) susceptible *V. vinifera* compared to resistant genotypes of *Vitis* and *Muscadinia* species, and (b) comparison of pathogen-induced gene expression with gene expression triggered by salicylic acid and ethylene.
3. Detailed analysis of candidate genes: (a) Real Time PCR to validate candidate genes identified in objectives 1 and 2, (b) Real Time PCR to study kinetics and specificity of the host response in susceptible and resistant genotypes, and (c) *in situ* hybridization to establish precise location of plant gene expression relative to bacterial infection.

RESULTS
Testing the effect of plant water status on PD
Two lines of evidence suggest that plant water status may have a significant impact on the development of PD symptoms. First, it is frequently observed that well-watered plants develop reduced symptoms relative to water-stressed plants. Thus, one might expect to see an enhanced transcriptional response in plants that are both water-stressed and infected by the pathogen. Second, it has been proposed that *Xylella* infection of xylem elements obstructs water flow, leading to whole-plant water stress and consequently to symptom development. Despite the logic of this reasoning, a causal relationship between *Xylella* infection and water stress has not been established. It is noteworthy, that the “water stress” hypothesis does not explain the absence of symptoms early in the season, even though high pathogen titers can be observed at this phase of disease, and it does not explain the absence of symptoms in tolerant genotypes of grapes, which can be heavily infected by the pathogen but without disease.

The experimental design described below permits a comparison of (1) pre-symptomatic and post-symptomatic host responses, (2) drought stressed versus diseased individuals, and (3) the interaction between drought stress and pathogen infection. In total, fourteen different transcriptional states that were compared to address these issues.

The experimental design involved 42 three-year-old vines of Cabernet Sauvignon clone 8 grafted to Freedom rootstock. In the spring of 2004, potted vines were moved from greenhouse to growth chamber prior to budbreak. Subsequent to a 3 to 4 week acclimation period, vines were pruned to produce a uniform shoot architecture consisting of two shoots per plant and ten leaves per shoot. Plants were grown in a block design of 3 rows with all treatments randomized in each row. Water use was calculated by watering 5 plants to field capacity and using a mini-lysimeter to establish water usage over a 24-hour period. The resulting average value was used to define 100% estimated water use. Plants were watered either at 100% water usage, 50% water usage (mild stress), or 25% water usage (moderate stress) throughout the remainder of the experiment. For each plant, measurements were made on the second leaf opposite to cluster to infer the level of drought stress pre- and post-veraison. Gas exchange and stomatal conductance values were obtained with a Licor 6400 gas exchange analyzer. C13:C12 ratios were measured on the same leaf samples used for transcriptional profiling to estimate long term effects of treatments on stomatal conductance, gas exchange and water use efficiency. On April 12, corresponding to full bloom, plants were either inoculated with a suspension of *Xf* or mock inoculated with water. Four weeks following inoculation, the third and fourth leaves were harvested from three plants of each treatment type. At 8 weeks following inoculation, when symptoms were evident on infected individuals, the remaining plants (3 from each treatment) were harvested. On the day of harvest for arrays the 5th leaf from each plant was destructively sampled to measure “pre-dawn” water potential. Symptom development was recorded using a visual scale.

RNA was extracted from tissue using protocols that we have optimized for quality and yield of RNA from grape (Iandolino et al., 2004). cRNA synthesis was carried out according to procedures described in the Affymetrix technical manual. Hybridization and data collection were performed using standard Affymetrix protocols, with the aid of the University of California, Davis microarray facility in the University of California, Davis Genome Center. Technical and biological replicates demonstrated highly consistent results within and between similarly treated samples. Quality control analyses were conducted using GCOS 1.2 (Affymetrix), Dechip (Li and Wong, 2001), and the Affy R package. Robust Multichip Average or RMA (Irizarre et al., 2003) was used to estimate differentially expressed genes by two different strategies: a) application of t-test and fold change filters (Sottosanto et al., 2004); and b) false discovery rate determinations using Significance
FUNCTIONAL GENOMICS OF THE GRAPE-XYLELLA INTERACTION: TOWARDS THE IDENTIFICATION OF HOST RESISTANCE DETERMINANTS

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Reporting Period: The results reported here are from work conducted October 2004 to October 2005.

ABSTRACT
In silico mining of EST data, Real Time PCR, and Affymetrix GeneChip technology was used to characterize the transcriptional response of *Vitis vinifera* to the Pierce’s disease (PD) pathogen *Xylella fastidiosa* (*Xf*). We have determined that susceptible *V. vinifera* responds to *Xylella* infection with a massive re-direction of gene transcription. This transcriptional response includes the up regulation of transcripts for phenylpropanoid and flavonoid biosynthesis, ethylene production, adaptation to oxidative stress, and homologs of pathogenesis related (PR) proteins. In addition to highlighting potential metabolic and biochemical changes that are correlated with disease, the results suggest that susceptible genotypes respond to *Xylella* infection by induction of limited defense response.

A long-standing hypothesis states that PD results from pathogen-induced drought stress, with the consequent development of disease symptoms. To test this hypothesis, we compared the transcriptional and physiological response of plants treated by pathogen infection, low or moderate water deficit, or a combination of pathogen infection and water deficit. We determined that the transcriptional response of plants to *Xylella* infection is not the same as the response of healthy plants to moderate water stress. However, there is an apparent synergistic interaction between water stress and disease, such that water stressed plants exhibit a stronger physiological and transcriptional response to the pathogen. Qualitative and quantitative estimates of gene expression derived from the Affymetrix gene chip were confirmed by a combination of Real Time PCR and *in situ* hybridization analysis with ~20 candidate marker genes.

Real Time PCR analysis involving six marker genes was used to survey the specificity of *Xylella*-induced gene expression under field conditions. The results demonstrate that the marker genes are up-regulated in response to *Xylella* infection but not in response to the other pathogens assayed, including common viral, nematode and fungal pathogens, or by *Phylloxera* infestation or herbicide damage. Similarly, moderate drought stress did not result in increased transcript levels for these marker genes. By contrast, each of the marker genes was strongly induced in non-infected leaves where the vascular system was compromised by biotic or abiotic factors, including girdling by insect damage and severe drought stress leading to death. We hypothesize that an aspect of xylem dysfunction, but not drought stress per se, is one trigger for *Xylella*-induced gene expression.

INTRODUCTION
All organisms adapt to external stressors by activating the expression of genes that confer adaptation to the particular stress. In the case of Pierce’s disease (PD), such genes are likely to include those coding for resistance or susceptibility to *Xylella fastidiosa* (*Xf*).

Genomics technology offers an opportunity to monitor gene expression changes on a massive scale (so-called "transcriptional profiling"), with the parallel analysis of thousands of host genes conducted in a single experiment. In the case of PD of grapes, the resulting data can reveal aspects of the host response that are inaccessible by other experimental strategies. In May of 2004, the first Affymetrix gene chip was made available for public use, with ~15,700 *Vitis* genes represented. This gene chip has been developed based primarily on a collaboration between the Cook laboratory and researchers at the University of Nevada-Reno (Goes da Silva et al., 2005). With the arrival of the Affymetrix gene chip, we are poised to make a quantum leap in the identification of host gene expression in response to *Xf*.

In addition to enumerating differences between susceptible and resistant genotypes of *Vitis*, this research is testing a long-standing but largely untested hypothesis that pathogen-induced drought stress is one of the fundamental triggers of PD symptom development. The utility of this type of data will be to inform the PD research community about the genes and corresponding protein products that are produced in susceptible, tolerant and resistant interactions. Differences in the transcriptional profiles between these situations are expected to include host resistance and susceptibility genes, and thus
examined, a previous survey of 16 species in riparian areas in Napa showed that BGSSs were more common on California blackberry, California grapevine, elderberry, and Himalayan blackberry, than on periwinkle from April to July (Purcell 1976). From September to March, BGSSs were more common on periwinkle (Purcell 1976). Differences in our results may be due to differences in locations, study years, or sampling methods.

Successful long-term management of PD may require removal of certain reservoir hosts, given that insecticides do not significantly reduce the spread of the disease (Purcell 1979) and that resistant winegrape varieties are not available. Wistrom and Purcell (2005) ranked the most important reservoir hosts, in terms of vector acquisition, as those that are feeding hosts of the BGSS, are frequently infected after transmission events, are systemic hosts of $X_f$, and support high pathogen populations. Revegetation of a riparian area adjacent to a diseased vineyard offers the potential to reduce the pathogen reservoir outside the vineyard, but it may be of limited efficacy in controlling the disease when infected grapevines remain in the vineyard. Grapevines satisfy all of Wistrom and Purcell’s (2005) criteria of important reservoir hosts and, thus, serve as a source of the pathogen for acquisition by BGSSs, even if riparian hosts are removed from an adjacent riparian area. Furthermore, removal of reservoir hosts may not diminish the ability of a riparian area to support BGSSs. Riparian areas are considered to be a habitat of the BGSS; they harbor many feeding and reproductive hosts (Freitag and Frazier 1954; Purcell 1975, 1976), in addition to plants that provide shelter for the overwintering adults.

REFERENCES

FUNDING AGENCIES
Funding for this project was provided by the USDA Agricultural Research Service, and the CDFA Pierce’s Disease and Glassy-winged Sharpshooter Board.
CONCLUSIONS
We measured abundance of the BGSS on five species (California blackberry, California grapevine, elderberry, Himalayan blackberry, and periwinkle) of naturally-established plants in riparian areas adjacent to vineyards on the North Coast of California. We assessed the ability of the same species to support Xf, based on results from controlled inoculations of potted plants kept in screenhouses in the field. None of the species were characterized by both an abundance of BGSSs and a high frequency of Xf detection. California grapevine and periwinkle may be more important pathogen reservoirs than California blackberry, elderberry, and Himalayan blackberry. Despite a significantly lower frequency of Xf detection in California grapevine, 19%, this species supported eight times as many nymphs and three times as many adults as periwinkle, suggesting that more frequent visitations by the BGSS may result in a high probability of acquisition of Xf from California grapevine. While periwinkle supported the fewest BGSSs, 71% of tested plants were Xf-positive, suggesting that a high percentage of transmission events result in systemic infection and that infrequent visitation by the vector may, nonetheless, result in a high acquisition rate. California blackberry, elderberry, and Himalayan blackberry are likely less important pathogen reservoirs because Xf was infrequently detected in their tissues and BGSSs were rare on these species.

Our finding that abundance of nymphs, but not that of adults, differed significantly among the riparian hosts we examined are consistent with those of Purcell (1976) who found that nymphs utilize fewer species than do adults. We might expect that nymph BGSSs have more restricted host ranges than adults based on different feeding requirements, as has been demonstrated for Homalodisca coagulata (Say) (glassy-winged sharpshooter) (Brodbeck et al. 1995), an introduced vector of PD in southern California (Blua et al. 1999). The significance of nymph BGSSs in the spread of PD is not known. Although nymphs lose infectivity after molting (Purcell and Finlay 1979), this may not preclude their importance in the epidemiology of PD relative to that of adults, which are infective for life once they acquire Xf (Purcell and Finlay 1979). The low mobility of nymphs, due to their flightlessness and small size, likely results in more transmission of Xf within an infected host than between hosts. Consequently, nymphs may spread Xf to new tissues within an infected host faster than the pathogen can move systemically. Systemic hosts of Xf on which nymphs are abundant, such as California grapevine, may support infections in more tissues than hosts on which nymphs are rare and, therefore, may serve as important sources of Xf for acquisition by adults.

It is possible that Xf infection of the species we examined through controlled inoculations are different in naturally-established plants of the same species. Natural levels of infection are related to a host’s ability to support Xf and its attractiveness to the BGSS. California grapevine, for example, may have higher levels of infection in the field than we measured in our inoculated plants, based on the high number of BGSSs we found on this species. There are few published surveys of Xf in naturally-established plants (Raju et al. 1983; Raju et al. 1980). In one such study of 28 native and non-native species in riparian areas in Napa County, Xf was detected in only four species: Himalayan blackberry, periwinkle, Fragaria vesca L. (wood strawberry), and Claytonia perfoliata Willd. (miner’s lettuce) (Raju et al. 1983). Although they surveyed California grapevine and elderberry, hosts that we also examined, their study was designed with the objective of identifying reservoir hosts, as opposed to comparing natural levels of infection among species.

Throughout the growing season, BGSSs occur in both riparian areas and vineyards (Freitag and Frazier 1954; Purcell 1975, 1976). Their whereabouts and behavior outside the growing season, when the population consists of adults (Purcell 1975; Severin 1949), are not well understood, mainly because cold temperatures limit BGSS flight activity (Feil et al. 2000) and, thus, hamper monitoring efforts. We measured BGSS abundance from spring to early summer, as this time of the year is characterized by BGSS flight activity, the presence of both adults and nymphs, and active growth of the five riparian host species we examined. Although we detected no significant differences in abundance of BGSSs on the five riparian hosts we

![Figure 3. Correlation of abundance of adult BGSSs with mean abundance of nymph BGSSs](image)
inoculation technique ensured that all plants were challenged by the pathogen; (ii) by sampling tissues distal to the inoculation site, \(Xf\)-positive identifications were known to represent systemic infections; and (iii) plants were inoculated once and, therefore, the presence of \(Xf\) was known to result from a single infection. In other words, our approach did not rely on natural infection by the BGSS, which likely reflects not only the hosts’ abilities to maintain \(Xf\) infections, but also BGSS feeding behavior.

**RESULTS**

Abundance of nymphs varied significantly among species \((P<0.0001)\). Nymphs were significantly more abundant on California grapevine which had a mean of 3.1 nymphs per sample, compared to all other hosts, but especially compared to periwinkle and elderberry, which had means of 0.4 nymphs per sample and 0.7 nymphs per sample, respectively (Figure 1). Abundance of adults was not significantly different among species \((P=0.0676)\). California grapevine, the species with the most nymphs, also had the most adults, 2.4 per sample (Figure 1). In contrast, periwinkle, the species with the fewest nymphs, also had the fewest adults, 0.9 per sample (Figure 1).

Frequency of detection of \(Xf\) varied significantly among species \((P<0.0001)\). Periwinkle had the highest frequency of detection with 70.8% of all tested plants, averaged across three sampling periods, found to be \(Xf\)-positive (Figure 2). Frequency of detection of \(Xf\) did not vary significantly between the two detection methods, colony counts in culture and real-time PCR \((P=0.09)\). Results from both detection methods showed the same relative differences among species; the interaction of species x detection method was not significant \((P=0.3582)\). For example, periwinkle had the highest percentage of plants that were found to be \(Xf\)-positive by culture (113 out of 160 total samples tested, summed across sampling periods) and by real-time PCR (140 out of 160 total samples tested, summed across sampling periods). In contrast, none of the 202 culture attempts from elderberry samples yielded \(Xf\) colonies, and real-time PCR analyses of the same tissues resulted in only six \(Xf\)-positive samples.

Despite the lack of statistical significance for differences in abundance of adults among riparian hosts from ANOVA \((P=0.07)\), there was a significant positive correlation between abundance of adults and nymphs \((r=0.96, P=0.01)\). Samples with many nymphs also had many adults (Figure 3). There were no correlations between detection frequency of \(Xf\) and abundance of adults \((r=-0.44, P=0.45)\) or nymphs \((r=-0.34, P=0.58)\).

**Figure 1.** Abundance of BGSSs on naturally-established riparian hosts adjacent to vineyards in northern California. A sample consisted of 25 sweeps per plant; \(n=13\) to 95 samples per species per year. Each column is the sum of the mean number of adults and nymphs per sample per species, averaged over years. Columns within each life stage with different letters are significantly different at \(P<0.05\) (Tukey’s test).

**Figure 2.** Frequency of detection of \(Xf\) from riparian hosts. Plants were inoculated in the greenhouse. Infected plants were placed in the field and subsequently tested at 3, 11, & 13 mos., by culture and real-time PCR; \(n=45\) to 76 plants per species per sampling period. Each column is the mean percentage of plants that were \(Xf\)-positive, averaged over sampling periods and detection methods. Columns with different letters are significantly different at \(P<0.05\) (Tukey’s test).