A Soil Inoculant Inhibits *Armillaria mellea* In Vitro and Improves Productivity of Grapevines with Root Disease

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ABSTRACT


A soil inoculant, Vesta (Biologically Integrated Organics, Inc., Sonoma, CA), was tested for its ability to inhibit *Armillaria mellea*, causal agent of Armillaria root disease of grapevine (*Vitis vinifera*). Colony diameter of *A. mellea* was significantly inhibited by undiluted inoculant (*P* < 0.0001) and by bacterial isolates cultured from the inoculant (*Bacillus subtilis*, *B. lentimorbus*, *Comamonas testosteroni*, *Pseudomonas aeruginosa*, *P. mendocina*; *P* < 0.0001) relative to diameter of the nontreated control. Efficacy of the inoculant for postinfection control of Armillaria root disease of grapevine was examined in an *A. mellea*-infested vineyard in northern California. Inoculant was applied via drip-irrigation to vine rows in replicate blocks in 2003 and 2004. Yield, growth, mineral nutrition, and juice quality parameters of healthy and symptomatic vines were measured in treated and nontreated vine rows. Significantly decreased petiole P and K concentrations and significantly lower soluble solids content in fruit from symptomatic vines demonstrated that Armillaria root disease negatively affects vine mineral nutritional status and fruit quality, findings that have not been previously reported for an agronomic host of *A. mellea*. The inoculant significantly increased cluster weights of symptomatic vines (109.63 g/cluster), relative to those of symptomatic-nontreated vines (92.05 g/cluster), to levels comparable to those of healthy vines (122.09 g/cluster). However, the inoculant did not decrease the rate of symptom development or mortality of treated vines from 2002 to 2004. The results of our field experiment suggest that the inoculant may not prevent Armillaria root disease, but can provide therapeutic benefit by improving productivity of infected vines.

Additional keywords: biological control

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Armillaria root disease of grapevine (*Vitis vinifera* L.) is caused by *Armillaria mellea* (Vahl:Fr.). P. Kumm., a fungal pathogen that also infects and decays the woody roots of many common forest trees in California, such as *Quercus kelloggii* Newb. (California black oak) (5). Where forests are converted to vineyards, mycelium of *A. mellea* can survive in partially decayed tree roots and infect grapevines up to several years after planting (6).

Most research on the control of Armillaria root disease on cultivated plants has focused on preplant use of soil fumigants to kill inoculum of the pathogen in partially decayed tree roots (1,8,20,21). Methyl bromide and carbon disulfide can kill mycelium in partially decayed tree roots, but not to the depths in soil necessary to reach all of the infested roots (8), especially on sites that were cleared of mature forest trees. Soil fumigants typically provide control for several years (17). However, replants eventually become infected when their roots contact inoculum that escapes the effects of soil fumigation. In this way, reliance on soil fumigation for control of Armillaria root disease leads grape-growers into an endless cycle of removing dead vines, fumigating, and replanting. Nonetheless, soil fumigation is one of few available control treatments. Given that methyl bromide will eventually be banned from use in California vineyards, an alternative to soil fumigation is needed.

Another preplant treatment that has been examined is the use of antagonistic fungi for biological control of *A. mellea* mycelium in buried tree roots. This approach was pursued based on findings that fumigants either kill or weaken *A. mellea* mycelium in buried wood segments (16,19,20,22), which predisposes the pathogen to attack by antagonistic soilborne fungi such as *Trichoderma viride* Pers.:Fr., a hypothesis that was first proposed by Bliss (8). Although some tested *Trichoderma* isolates show excellent in vitro and in vivo (25) inhibition of *A. mellea* growth, it is not clear that necessary concentrations of antagonistic fungi can be achieved in field soil (26).

Given the demonstrated futility of preplant soil fumigation and the difficulty involved with inoculating soils with sufficient populations of antagonistic fungi, efforts may be better spent on postinfection treatments that improve an infected grapevine’s tolerance of Armillaria root disease. We might expect postinfection treatments to be a successful approach for two important reasons: (i) *A. mellea* is a slow-growing fungus (colony growth on 1% malt extract agar incubated at room temperature is approximately 5 mm/week), and (ii) grapevines are relatively tolerant of infection, based on the fact that it typically takes several years for an infected grapevine to die from Armillaria root disease (6). In addition, a recent study demonstrated the efficacy of root collar excavation for postinfection treatment of grapevines infected with *A. mellea* (4). The practice significantly mitigates one of the disease symptoms, decreased fruit productivity (4), by causing mycelial fans to recede from the root collar, thereby improving the function of vascular tissue at the base of the trunk. Therefore, control measures designed to enhance the yield of infected grapevines may prove to be more cost-effective than replanting, given that replants are not immediately productive and may eventually become infected by *A. mellea*.

The objective of this research was to examine the efficacy of a commercially available soil inoculant, Vesta (Biologically Integrated Organics, Inc., Sonoma, CA), for postinfection control of Armillaria root disease of grapevine. The inoculant is produced by a proprietary compost fermentation process, the product of which contains viable populations of bacteria that may serve as antagonists of *A. mellea*. Microbial inoculants and other biologically based products are of interest to grape-growers who want to reduce their reliance on pesticides.

A series of laboratory experiments was used to examine inhibition of *A. mellea* by the inoculant, and to isolate and identify bacteria associated with inhibition. A field experiment was conducted in an *A. mellea*-infested vineyard to determine if applications of the inoculant to symptomatic vines could improve the following performance parameters: yield, growth, mineral nutrition, and juice quality. Given that yield and growth of symptomatic vines are known to be significantly lower
than those of healthy vines (4), we viewed amelioration of these impacts as measures of the efficacy of the inoculant. Examining naturally infected vines in the field is preferable to inoculating potted vines with *A. mellea* in the greenhouse, in that the latter is neither a reliable nor timely means of obtaining infected vines (23,24,27). The factorial design of the field experiment necessitated the inclusion of healthy vines, allowing for quantification of effects of the disease on vine mineral nutrition and juice quality, a logical complement to existing knowledge of the negative effects of the disease on vine yield and growth.

**MATERIALS AND METHODS**

**Inhibition assays.** The inoculant, Vesta (Biologically Integrated Organics, Inc., Sonoma, CA), is produced by fermentation of a proprietary blend of composted materials, the product of which contains viable populations of bacteria that are monitored throughout the production process using colony counts of functional groups of microorganisms (heterotrophic bacteria, aerobic bacteria, Pseudomonads, nitrogen-fixing bacteria). After fermentation, the resulting product is amended with humic acid and other organic acids. Vesta is a brown, opaque liquid that is applied to vine rows by injection into the drip-irrigation system, with agitation, but can also be applied to individual vines as a soil drench. Recommended application rates are 75 to 112 liters/ha (8 to 12 gallons/acre) per season.

The soil inoculant was used to challenge *A. mellea* in inhibition assays. Assays were conducted on yeast extract malt agar (YMA) on 100-mm-diameter petri plates with an *A. mellea* isolate cultured from symptomatic vines in the experimental vineyard. A sterile, 6-mm-diameter paper disk was saturated with 20 µl of undiluted inoculant and placed 5 mm from an *A. mellea* agar plug on each of four YMA plates. Disks saturated with sterile water served as nontreated controls. Diameters of *A. mellea* colonies were recorded after incubation at 25°C for 21 days. Inhibition assays were repeated three times, to give a total of 12 plates for the inoculant and 12 plates for the nontreated control (water). Bacteria were isolated from expanding colonies surrounding the inoculating disks from plates on which *A. mellea* growth had been inhibited the most. These isolates were used in the remaining laboratory tests.

Bacterial isolates cultured from the inoculant were used to challenge *A. mellea* in a second set of inhibition assays. Assays were conducted as described above with four plates per bacterial isolate. A sterile paper disk saturated with 20 µl of a turbid suspension of vegetative cells (approximately 10⁶ cells/ml collected from a 24-h-old culture on YMA) in sterile water of each bacterial isolate was placed 5 mm from each *A. mellea* plug. Sterile disks with sterile water served as nontreated controls. Diameters of *A. mellea* colonies were evaluated as described above. Inhibition assays were repeated twice, to give a total of eight plates for each bacterial isolate and eight plates for the nontreated control (water). Bacterial isolates that limited *A. mellea* colonies to the same diameter as did the inoculant were identified by fatty acid methyl ester (FAME) analysis. Fatty acids were extracted and analyzed by gas chromatographic analysis (BBC Laboratories, Tempe, AZ) using the technique described by Kloeper et al. (18). Identifications were based on similarity index values of each isolate, which represent the similarity between the FAME profile of the unknown isolate and a bacterial species from a database of FAME profiles of known isolates, on a scale of 0.001 (least similar) to 0.999 (most similar).

**Field trials.** A field experiment was conducted in a commercial vineyard in Napa County, CA, during the 2003 and 2004 growing seasons. *A. mellea* had been discovered in 2000 on dead and dying vines in the vineyard. The vineyard was planted in 1997 with dormant bench grafts of *V. vinifera* cv. Cabernet Sauvignon (clone 337) on 110R rootstock (*V. berlandieri* Planch. × *V. rupestris* Scheele). The 3.4-ha site was previously an oak woodland. Vine spacing was 1.8 m within rows and 2.3 m between rows, with east-west row orientation. Vines were trained as unilateral cordons to a vertical shoot positioning trellis system. The vineyard was drip-irrigated (43 kl/ha/week, July-October) with one drip emitter per vine positioned 0.3 m from each vine trunk.

In July 2002, the status of every vine was categorized, based on approximate shoot length, as symptomless (≥1 m), moderately symptomatic (0.3 to 1 m), or severely symptomatic (≤0.3 m), using the procedure described previously (4). Severely symptomatic vines were excluded from this study because they often die before the fruit can mature (4). Vine status was documented at harvest on the following dates: 19 September 2002, 18 October 2003, and 23 September 2004. In November 2002, the presence of *A. mellea* infection on moderately symptomatic vines was verified by clearing soil away from the root collars to a depth of approximately 0.3 m and removing a small piece of bark, ca. 2 cm² area, from the base of the trunk and from each main root to expose mycelial fans. The same method was used to verify absence of *A. mellea* on symptomless vines. Hereafter, symptomless vines with no mycelial fans at their root collars will be referred to as “healthy”, with the understanding that some might actually have been infected, but were not yet symptomatic. Moderately symptomatic vines with mycelial fans at their root collars will be referred to as “symptomatic”.

The vineyard included three replicate blocks (0.44 ha/block), with a nontreated buffer row between blocks. Experimental treatments were arranged in a split plot design with inoculant treatment (treated or nontreated) as the main plot and vine status (healthy or symptomatic) as the subplot. An experimental block consisted of 10 vineyard rows: four adjacent treated rows plus five adjacent nontreated rows, separated by a nontreated buffer row. Healthy and symptomatic vines, the subplot treatments, were randomly selected from within the treated and nontreated rows, the main plot treatments, which were randomly assigned to a half of each block.

Healthy vines were included for comparison with symptomatic vines, in order to verify the negative effects of Armillaria root disease on the symptomatic vines, evidence without which inoculant effects may be considered somewhat specious. In 2003 and 2004, the inoculant was injected into the irrigation system of treated rows at the following rates: budbreak (46.77 liters/ha), full bloom (46.77 liters/ha), 15% veraison (onset of fruit ripening) (18.71 liters/ha), and 85% veraison (18.71 liters/ha). All three phenological stages are visible in the field and are, therefore, typically used by grape-growers to schedule fungicide and fertilizer applications. Nontreated rows received equal rates of irrigation water. In treated and nontreated sections of each block, 12 healthy and 12 symptomatic data vines were randomly chosen for measurement of yield, growth, mineral nutrition, and juice quality parameters, for a total of four experimental treatments.

Fruit clusters from sampled vines were collected, harvested, and weighed on 18 October 2003 and 23 September 2004, resulting in a total of three yield parameters for each vine: cluster number, yield (kg/vine), and average cluster weight (yield [g]/cluster number). At harvest, one cluster/vine from five vines per treatment per block were subsampled and analyzed for three juice quality parameters: total soluble solids (°Brix), measured with a table-top refractometer (Carl Zeiss Inc., Thornwood, NY); pH (Accumet pH meter, Fisher Scientific, Fair Lawn, NJ); and titratable acidity (g tartaric acid/liter of juice; [TA]), using standard methods (9). TA was estimated based on the volume of 0.1 N NaOH required to bring about a color change in a mixture of 5 ml juice from each five-cluster sample, 100 ml distilled water, and 2 drops phenolphthalein (indicator). Total soluble solids, pH, and TA are routinely measured by grape-growers and winemakers for the purpose of timing harvest because all three parameters affect wine quality. From the same vines from which yields were measured, dormant canes were pruned and weighed on 21 January 2004 and 14 December 2004. Average shoot weight was obtained
RESULTS

Inhibition assays. In the presence of the inoculant, *A. mellea* colonies grew to diameters of 5 to 12 mm (including the 4-mm-diameter agar plug) on all 12 plates, compared to 18 to 22 mm on control plates. This differential reflected a significant inhibitory effect of the inoculant on the pathogen (*P* < 0.0001). Twelve bacteria, isolated from plates on which *A. mellea* colonies were limited to <8 mm, were used in a second set of inhibition assays. Seven of these isolates limited expansion of *A. mellea* colonies to 5 to 8 mm. When compared to the control plates, where *A. mellea* colonies ranged from 18 to 22 mm, inhibition of the pathogen by the seven isolates was significant (*P* < 0.0001). Five species were identified from the seven isolates that inhibited *A. mellea* in vitro to the same extent as the inoculant. These were (with FAME similarity index values following in parentheses) two isolates of *Bacillus subtilis* (Ehrenberg 1835) Cohn 1872 (0.741 and 0.728), two isolates of *B. lentimorbus* Dutky 1940 (0.707 and 0.671), one isolate of *Comamonas testosteroni* (Marcus and Talalay 1956) Tamaoka et al. 1987 (0.944), one isolate of *Pseudomonas aeruginosa* (Schroeter 1872) Migula 1900 (0.932), and one isolate of *P. mendocina* Palleroni 1970 (0.661).

Field trials. In both years, the inoculant had significant effects on yields and cluster weights of symptomatic vines, as detected by ANOVA, but no effects on healthy vines; hence, the significant status × treatment interaction effects on yield and cluster weight (Table 1). Although there was a significant effect of the status × treatment interaction on yield, means comparisons with Tukey’s tests following ANOVA showed no significant differences between yields of symptomatic-treated and symptomatic-nontreated vines (Fig. 1A). In contrast, cluster weights of symptomatic vines treated with the inoculant were 17.58 g more than those of symptomatic-nontreated vines, and these differences were significant (Fig. 1B). In fact, cluster weights of symptomatic-treated vines were as high as those of healthy vines.

Among the three factors examined by ANOVA (year, vine status, inoculant treatment), inoculant treatment was the only factor that significantly affected soil

Table 1. Analyses of variance for combined 2003 and 2004 cluster number, yield, and cluster weight of healthy and symptomatic grapevines, with or without inoculant treatment, in a vineyard infested with *Armillaria mellea*

<table>
<thead>
<tr>
<th>Source/interaction</th>
<th>Cluster number</th>
<th>Den df&lt;sup&gt;a&lt;/sup&gt;</th>
<th>F value</th>
<th>Yield</th>
<th>Den df&lt;sup&gt;a&lt;/sup&gt;</th>
<th>F value</th>
<th>Cluster weight</th>
<th>Den df&lt;sup&gt;a&lt;/sup&gt;</th>
<th>F value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Year</td>
<td></td>
<td></td>
<td>3.69</td>
<td></td>
<td></td>
<td>3.98</td>
<td></td>
<td>3.72</td>
<td></td>
</tr>
<tr>
<td>Status</td>
<td>236</td>
<td>165.11***</td>
<td>237</td>
<td>150.79***</td>
<td>237</td>
<td>6.13*</td>
<td>239</td>
<td>29.58***</td>
<td></td>
</tr>
<tr>
<td>Year × status</td>
<td>236</td>
<td>7.21**</td>
<td>237</td>
<td>0.52</td>
<td>237</td>
<td>3.88*</td>
<td>239</td>
<td>6.81**</td>
<td></td>
</tr>
<tr>
<td>Treatment</td>
<td>2</td>
<td>0.61</td>
<td>2</td>
<td>0.78</td>
<td>4</td>
<td>2.81</td>
<td></td>
<td>0.53</td>
<td></td>
</tr>
<tr>
<td>Year × treatment</td>
<td>237</td>
<td>0.01</td>
<td>237</td>
<td>0.52</td>
<td>237</td>
<td>3.88*</td>
<td>239</td>
<td>6.81**</td>
<td></td>
</tr>
<tr>
<td>Status × treatment</td>
<td>237</td>
<td>0.46</td>
<td>237</td>
<td>0.83</td>
<td>237</td>
<td>0.83</td>
<td>239</td>
<td>0.53</td>
<td></td>
</tr>
<tr>
<td>Year × status × treatment</td>
<td>237</td>
<td>0.87</td>
<td>237</td>
<td>0.83</td>
<td>237</td>
<td>0.83</td>
<td>239</td>
<td>0.53</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup> Source of variation: 2003 or 2004 (year), healthy or symptomatic (status), nontreated or treated (inoculant treatment).

<sup>b</sup> Denominator degrees of freedom. Numerator degrees of freedom are 1 for all main and interaction effects; *, **, and *** indicate significance at *P* ≤ 0.05, 0.01, and 0.0001, respectively.
composition, specifically soil C ($P = 0.0351$). Soil from treated vines had significantly higher soil C than that of nontreated vines ($34.45$ and $28.80$ mg/g dry soil of total C, respectively). There were no significant differences in vine growth, petiole nutritional status, or juice quality between treated and nontreated vines (data not shown). In comparisons of counts of healthy, symptomatic, and dead vines between nontreated and treated rows (Table 2), chi-square analyses indicated that there were no significant differences in the proportions of healthy vines that developed symptoms or died from 2002 to 2003 or from 2003 to 2004 (Table 2). Similarly, the inoculant had no effect on the proportion of symptomatic vines that died in 2003 or 2004.

Symptomatic vines had significantly fewer clusters, lower yields, and lower cluster weights than healthy vines in both study years (Table 1), despite a significant decrease among healthy vines in cluster number per vine ($26.52$ versus $31.73$; $P = 0.0125$) and yield per vine ($3.23$ versus $3.88$; $P = 0.0111$) in 2004, compared to 2003. Unlike healthy vines, symptomatic vines did not differ in cluster number or yield from 2003 to 2004 ($P = 0.9997$ and $P = 0.9912$, respectively); hence the significant year × status interaction effects on cluster number and yield (Table 1). Compared to healthy vines, symptomatic vines had far fewer clusters, resulting in much lower yields in both years ($17.01$ clusters/vine and $1.79$ kg/vine, averaged across 2003 and 2004, respectively).

Among the three factors examined in ANOVA (year, vine status, inoculant treatment), vine status was the only factor that significantly affected juice quality, vine growth, and petiole nutrition (Table 3). In addition to having fewer clusters, lower yields, and lower cluster weights, symptomatic vines produced clusters with significantly lower, by $1.38$ °Brix, concentrations of total soluble solids than healthy vines in both years ($P = 0.0170$). Symptomatic vines had significantly fewer shoots ($P < 0.0001$), lower pruning weights ($P < 0.0001$), and lower shoot weights ($P < 0.0001$) than healthy vines. In fact, pruning weights of symptomatic vines were $58\%$ lower than those of healthy vines. Symptomatic vines also had significantly lower petiole concentrations of P and K than healthy vines ($P = 0.0021$ and $P = 0.0108$, respectively). There were no significant differences in soil nutrition between healthy and symptomatic vines (data not shown). Averaged across healthy and symptomatic vines, the soil contained $1.95$ mg/g dry soil of total N, $49.68$ µg/g dry soil of NO$_3$-N, $51.00$ µg/g dry soil of Olsen P, and $430.89$ µg/g dry soil of exchangeable K, with pH $6.2$, and $241.72$ µmol/g dry soil of cation exchange capacity.

**DISCUSSION**

Using inhibition assays in the lab and repeated measurements of vine yield, growth, mineral nutrition, and juice quality in the field, we examined the potential of a microbial soil inoculant for postinfection management of Armillaria root disease of grapevine. The inoculant inhibited A. mellea, based on measures of colony diameter in culture. Our findings that symptomatic vines treated with inoculant had significantly higher cluster weights in both years and in all three replicate blocks of the vineyard indicate that applications of the inoculant may mitigate some of the effects of Armillaria root disease. Furthermore, there was no stimulatory effect of the inoculant on performance parameters of healthy vines, suggesting that the efficacy of the inoculant in improving cluster weights of symptomatic vines is associated with its effects on the pathogen.

Given that bacteria isolated from the inoculant (B. lentimorbus, B. subtilis, C. testosteroni, P. aeruginosa, P. mendocina) were individually found to inhibit A. mellea to the same extent as did the inoculant, it is possible these species are involved with inhibition. Reports that some of the bacteria we identified are antagonists of other pathogens, specifically B. subtilis (7,28), P. aeruginosa (2,3,14), and P. mendocina (12), suggest that these species may inhibit A. mellea in infected vine roots. For example, B. subtilis has been shown to inhibit Phytophthora cactorum (Leb. & Cohn) Schroeter (1886) in vitro and to reduce root rot incidence among infected apple seedlings (28). Dumas (15) and Delong et al. (13) demonstrated that Bacillus spp. and fluorescent pseudomonads isolated from forest soils inhibit the conifer pathogen A. ostoyae (Romagn.) Herink in vitro, suggesting that indigenous populations of antagonists may have a natural role in regulating Armillaria root disease in conifer forests. Even though we examined a different Armillaria species on a different host, results of research on antagonists of A. ostoyae have relevance to A. mellea inhibition by bacteria in the inoculant, given that A. mellea and A. ostoyae share a similar biology and infection cycle (29).

Bacteria isolated from the inoculant inhibited growth of A. mellea in vitro, and the inoculant compensated for compromised productivity of symptomatic vines in the field, results that infer direct or indi-

**Table 2.** Effects of inoculant on status of healthy and Armillaria-affected vines, based on chi-square tests* of independence between inoculant treatment and changes in health status

<table>
<thead>
<tr>
<th>Change in health status</th>
<th>Years</th>
<th>Percentage of vines that changed health status</th>
<th>$\chi^2$</th>
<th>$P^*$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Healthy to dead</td>
<td>2002 to 2003 $^b$</td>
<td>0.45</td>
<td>0.57</td>
<td>0.111</td>
</tr>
<tr>
<td></td>
<td>2003 to 2004 $^b$</td>
<td>1.39</td>
<td>2.07</td>
<td>1.069</td>
</tr>
<tr>
<td>Healthy to symptomatic</td>
<td>2002 to 2003</td>
<td>1.47</td>
<td>2.57</td>
<td>2.475</td>
</tr>
<tr>
<td></td>
<td>2003 to 2004</td>
<td>9.94</td>
<td>9.91</td>
<td>0.000</td>
</tr>
<tr>
<td>Symptomatic to dead</td>
<td>2002 to 2003</td>
<td>12.00</td>
<td>8.60</td>
<td>0.630</td>
</tr>
<tr>
<td></td>
<td>2003 to 2004</td>
<td>3.19</td>
<td>3.45</td>
<td>0.009</td>
</tr>
</tbody>
</table>

*Chi-square tests are based on comparisons between the number of vines in each category, summed across three replicate blocks of the experiment, in inoculant-treated and nontreated rows. $H_0$: there is no relation between treatment with the inoculant and change in health status; degrees of freedom = 1.

$^b$ Probability that the differences between nontreated and treated rows in observed counts of vines that changed status are due to chance alone; differences are significant at $P \leq 0.05$.

$^v$ Vine status was assessed on 19 September 2002, 18 October 2003.

$^a$ Vine status was assessed on 18 October 2003, 23 September 2004.

**Table 3.** Effects of Armillaria root disease on combined 2003 and 2004 growth, nutrition, and juice quality parameters of symptomatic vines, relative to those of healthy vines, in a vineyard infested with Armillaria mellea

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Healthy</th>
<th>Symptomatic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Juice quality</td>
<td>6.81 a</td>
<td>6.34 a</td>
</tr>
<tr>
<td>Total soluble solids (°Brix)</td>
<td>26.19 a</td>
<td>24.81 b</td>
</tr>
<tr>
<td>pH</td>
<td>3.39 a</td>
<td>3.46 a</td>
</tr>
<tr>
<td>Growth</td>
<td>22.04 a</td>
<td>19.66 b</td>
</tr>
<tr>
<td>Shoot number</td>
<td>0.60 a</td>
<td>0.25 b</td>
</tr>
<tr>
<td>Petiole nutrition</td>
<td>27.77 a</td>
<td>12.93 b</td>
</tr>
<tr>
<td>Total N (mg/g dry tissue)</td>
<td>9.32 a</td>
<td>10.07 a</td>
</tr>
<tr>
<td>Total P (mg/g dry tissue)</td>
<td>2.73 a</td>
<td>2.18 b</td>
</tr>
<tr>
<td>K (mg/g dry tissue)</td>
<td>30.05 a</td>
<td>26.51 b</td>
</tr>
</tbody>
</table>

$a$ Written, sums over the growing seasons of 2003 and 2004, for means within each set of parameters are as follows: juice quality, $n = 12$; growth, $n = 184$; petiole nutrition, $n = 12$. Means in the same row with different letters are significantly different at $P \leq 0.05$, Tukey’s test.
irect effects of the bacteria on *A. mellea*. However, the inoculant did not decrease rates of symptom development or mortality. Without knowing the mechanism(s) by which bacteria in the inoculant inhibit *A. mellea*, it is difficult to explain these somewhat contradictory findings. One possible explanation is that the bacteria replace *A. mellea* in infected roots, a situation that Bliss (8) found with *T. viride* on *Citrus* root segments infected with *A. mellea*, demonstrating that the pathogen may not be capable of indefinitely defending infected roots from soil microbes. Replacement of *A. mellea* in infected roots by the bacteria in the inoculant would have no effect on vascular tissue already destroyed by *A. mellea*, but it would reduce the pathogen’s food base and, thus, inhibit the colonization rate of adjacent root tissue. By inhibiting the spread of existing *A. mellea* infections, the bacteria in the inoculant may afford some level of root function recovery in symptomatic vines, thereby increasing cluster weights. Whatever the mechanism(s) of inhibition, the effects of the inoculant seem to be insufficient to prevent the onset of symptoms or eventual death of an infected vine.

While cluster weight increases were detected among symptomatic-treated vines in all three replicate blocks of the vineyard in both study years, our experiment included only one vineyard. Nonetheless, differences in yield and growth between healthy and symptomatic vines that we observed are similar to those reported for two other *A. mellea*-infested Napa vineyards (4). Given that the progression of Armillaria root disease in the examined vineyard does not appear to be unique in comparison with other Northern California vineyards where similar measurements were made, our results should be applicable to other vineyards.

Our findings of significantly increased soil C among treated vines, including both healthy vines and symptomatic vines, in both study years, and in all replicate blocks of the vineyard, are puzzling. The inoculant clearly contains bacteria that, after application to the soil, may establish populations in the grapevine rhizosphere and contribute to soil C. However, it seems unlikely that the bacteria could bring about such a considerable change in soil C, 5.7 mg/g dry soil of total C, in such a brief time. Research on sustainable cropping systems in northern California, in which one of the main objectives was to increase soil C, showed 10-fold lower increases in soil C than we observed in our study, and this was after 4 years of adding large quantities of compost and manure to the soil (11). If it was possible to raise soil C only by 0.5 mg/g dry soil of total C with 4 years of compost and manure amendments (11), it is surprising that we detected an increase in soil C of 5.7 mg/g dry soil of total C among treated vines in our experiment with 2 years of inoculant applications. Future investigations of the C sources and their concentrations in the inoculant might clarify this issue.

Although Armillaria root disease is known to decrease yield and growth of grapevines (4), negative effects of the disease on vine mineral nutrition and juice quality as documented in this study have not previously been reported. Above-ground symptoms of the disease result from destruction of vascular tissue at the root collar and on primary roots by *A. mellea* mycelium (6). Thus, an infected vine is unable to absorb an adequate supply of mineral nutrients. The lower yield and growth, reduced tissue P and K, and lower soluble solids in fruit from symptomatic vines measured in this study are consistent with this expectation. Destruction of vascular tissue in the woody roots and associated poor nutrient uptake reduce a symptomatic vine’s capacity; it has smaller shoots, and the buds on the shoots are less fruitful. In turn, deficiencies in P and K reduce photosynthetic capability, limiting yield and ripening. Growers could remedy this by reducing the crop load on symptomatic vines, in order to ensure proper ripening. While the petiole P and K concentrations that we measured among symptomatic vines were not technically deficient (adequate petiole concentrations at bloom are >1.5 mg P/g dry tissue and >15 mg K/g dry tissue; 10), they were statistically lower than those of healthy vines in all replicate blocks of the vineyard and in both study years. Therefore, it may be useful to evaluate foliar applications of P and K fertilizers to determine if they can compensate for lower root absorption of these macronutrients and, thus, help symptomatic vines tolerate Armillaria root disease.

The fact that significant cluster weight increases were measured among symptomatic-treated vines only 5 months after the first application are a clear indication of the inoculant’s therapeutic benefit. Focusing control efforts on improving the productivity of infected vines that still produce normal clusters is more cost-effective than replanting, which requires the expenditure of new plant material, the labor costs for planting, and the lack of profit while waiting at least 2 years for replants to bear fruit. Inoculant applications require only a standard drip irrigation system already found in most California vineyards. The grower incurs a relatively small cost for purchase of the inoculant. The high value of Napa County grapes, $4,400/1,000 kg at the vineyard gate, examined, means that yield increases on symptomatic vines may offset the cost of the inoculant.

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**LITERATURE CITED**


