Effects of Chitin and Chitosan on the Incidence and Severity of Fusarium Yellows of Celery

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ABSTRACT

The effects of chitin and chitosan on disease incidence and severity of Fusarium yellows of celery and on populations of Fusarium oxysporum were investigated between 1994 and 1996. Field experiments were conducted at two locations with a history of severe Fusarium yellows. Disease incidence and severity were significantly reduced by pre-plant chitin amendments to soil. Chitosan applied as a root dip alone did not reduce disease incidence but significantly (P < 0.05) reduced disease severity when used with a tolerant celery cultivar. Standard soil dilution methods were used to enumerate populations of soil microflora. Chitin increased bacterial and actinomycete populations in soil in 2 of the 3 years of study. The effects of potential biocontrol agents recovered from chitin-treated plots in 1995 were studied in 1996; enriching the transplant medium with isolates of bacteria and actinomycetes 4 weeks and 1 week prior to transplanting did not alter the established equilibrium in the field, and no biocontrol effect was observed. Chitin amendments to soil or chitosan treatment of transplants did not reduce soil populations of F. oxysporum. Whether these treatments affected the F. oxysporum f. sp. apii subpopulation within the F. oxysporum population could not be determined.

Additional keywords: nonchemical control, sustainable agriculture

Fusarium yellows, incited by Fusarium oxysporum f. sp. apii (Nelson & Scherb), is a vascular disease of celery (Apium graveolens L. var. dulce) that causes losses in both the quality and quantity of yield. Symptoms include brown or reddish-brown vascular discoloration and necrosis in roots and crowns. Vascular discoloration of the petiole and yellowing of the foliage occurs in plants with advanced infection.

From the late 1930s, Fusarium yellows was managed in California by the use of tolerant cultivars. In 1978, a new race, designated race 2, was reported in four coastal California counties (18). Initially, race 2 rendered all commercially available green cultivars susceptible, but cultivars later bred for resistance to race 2 were successful in managing the disease. However, since 1991, higher-than-expected levels of yellows-related discoloration occurred in tolerant celery lines. No new races of Fusarium yellows have been detected in California, but higher yellows incidence in tolerant cultivars may have been caused by increased inoculum density in the soil owing to continued celery culture.

Alternative methods of disease control to reduce the inoculum in soil or suppress the disease in celery cultivars have had limited success. Attempts to suppress the pathogen by fertilization with potassium, chloride, and nitrate ions (12,30), non-pathogenic strains of the pathogen (2,23,29), other rhizobacteria (4,14,37), and crop rotation (13) have been either ineffective in production fields or have not been adopted by growers.

Chitin (2) and chitosan amendments (6,11) may effectively reduce soilborne diseases. Addition of small quantities of chitin to soil resulted in a marked reduction in the severity of root rot of beans caused by F. solani f. sp. phaseoli and vascular wilt of radish caused by F. oxysporum f. sp. conglutinans (25). Chitin (β-1,4-linked-glucosamine polymer), a deacetylated derivative of chitin, has demonstrated fungal activity against several fungi (6,11,16,33). Chitosan is also known to be a potential elicitor of many plant defense responses, including stimulation of plant chitinases (6,11), pisatin induction (20), occlusion of the pit fields in xylem vessels, and formation of wall appositions (6). Application of chitosan may enhance the vitality of plant cells and the plant’s ability to degrade the walls of fungi upon entry (6). Moreover, the ready availability of chitin and chitosan from crustacean shell wastes has kindled interest in their potential agricultural applications.

The objectives of this research were to determine the effects of chitin and chitosan on the incidence of Fusarium yellows in celery cultivars with varying levels of yellows resistance, and on populations of F. oxysporum f. sp. apii, and to determine the potential mechanisms involved in disease suppression.

MATERIALS AND METHODS
Effects of chitosan concentration on mycelial growth of F. oxysporum f. sp. apii. To determine the optimal concentration of chitosan for treating celery transplants and to determine the uniformity of its effects on populations of F. oxysporum f. sp. apii, effects of chitosan were evaluated on five isolates of F. oxysporum f. sp. apii. Purified chitosan was prepared by dissolving chitosan (Sigma Chemical Co., St. Louis) in 0.25 N HCl by stirring for 8 h at 45°C. Undissolved particles were removed by centrifugation (10,000 × g, 15 min). Chitosan was precipitated with 2 N NaOH and washed three times in deionized water to remove salts. The purified chitosan was then air-dried and stored at room temperature until required. For incorporation into media and treatment of plant roots, purified chitosan was dissolved in 0.25 N HCl, then adjusted to pH 5.6 with 2 N NaOH. Chitosan was incorporated into potato dextrose agar (PDA; Difco Laboratories, Detroit, MI) at concentrations of 0, 1, 2, 3, 4, and 6 mg ml⁻¹. The chitosan solution and PDA were autoclaved separately and combined after autoclaving. Equal volumes of acid were used for all concentrations of chitosan, adjusted to pH 5.6 to keep the salt concentration constant. Five F. oxysporum f. sp. apii isolates were tested for inhibition by chitosan. A 4-mm diameter plug from the advancing margins of colonies of each isolate on PDA was seeded centrally onto 4 plates of each chitosan concentration. Cultures were incubated at 25°C. The diameter of all colonies was measured daily until the leading edge of the fastest-growing colony had reached the edge of the plate. The experiment was
Each season, the celery crop was grown using grower practices recommended for each location. The crop was furrow-irrigated in Santa Maria and sprinkler-irrigated in Oxnard. In 1994 and 1995, Fusarium yellows incidence in each plot was recorded 1 week prior to harvest by counting the total number of plants and the number showing characteristic symptoms. In 1996, the severity of Fusarium yellows in each treatment was recorded at harvest by scoring 10 randomly-chosen plants from the center of each plot for vascular discoloration of root and crown on a 0 to 5 scale, where 0 = no disease, 1 = slight discoloration of root tissue, 2 = extensive discoloration of root tissue, 3 = crown discolored, 4 = crown extensively discolored, and 5 = plant mortality. To determine whether treatments affected the marketable yield of the celery crop, fresh weight measurements of 15 plants in 1995 and 10 plants in 1996, selected at random from each plot, were recorded.

Soil samples in each season were collected prior to the introduction of treatments, and at monthly intervals until harvest, to determine the treatment influence on F. oxysporum populations. Samples from three random spots in each bed were collected using a trowel to a depth of 20 cm, and bulked. The samples were air-dried on greenhouse benches and the number of F. oxysporum propagules were determined by dilution plating on modified Komada’s medium (21). Data from each year were analyzed separately because the cultivars were different each year. Analysis of variance was conducted on the incidence and severity data, plant fresh weights, and on log-transformed F. oxysporum CFU’s. A least significant difference (LSD) test was used to compare treatment means.

Effects of chitin and chitosan on microbial populations in soil. Because chitin is known to significantly increase microorganisms in soil (15,17,25,28) that, in turn, may reduce populations of F. oxysporum f. sp. apii, soil samples collected during 1995 and 1996 were assayed for total actinomycetes, bacteria, and fungi. Soil samples collected as described above were transported to the laboratory in a cooler and processed within 24 h. To assess soil microflora, 10 g of each soil sample was suspended in 90 ml sterile water, stirred for 10 min, and serially diluted. For isolation of fungi, final dilutions of 10^5 to 10^0 were plated on PDA amended with 50 µg ml^-1 rose Bengal and 100 µg ml^-1 rifampicin. Plates were incubated at 25°C for 5 to 7 days before recording the numbers and types of fungi. For isolation of bacteria, final dilutions of 10^5 to 10^0 were placed on 0.5% tryptic soy broth (Difco Laboratories) and incubated at 37°C for 48 h before recording the numbers of bacteria. For isolation of actinomycetes, soil samples were air-dried at room temperature for 10 days before isolation. Final dilutions of 10^2 to 10^0 were placed on actinomycete isolation agar (Difco Laboratories) and incubated at 25°C for 5 to 7 days before recording numbers of actinomycetes. Representative colonies of fungi and actinomycetes were purified and stored on PDA slants at 4°C. Bacteria were stored at –70°C in vials with tryptic soy broth (Difco Laboratories) amended with 15% glycerol.

Soil moisture in each sample was determined gravimetrically. Populations of soil microorganisms from each treatment were adjusted for soil moisture and transformed to log CFU g^-1 soil and analyzed to determine treatment effects.

Repeated measures analysis of variance was used (GLM, SAS 6.11, SAS Institute, Inc., Cary, NC) to evaluate the effects of treatment, cultivar, date of sampling, and interaction using appropriate error terms. A LSD test (P < 0.05) was used to compare treatment means.

Evaluating the biocontrol potential of the microflora. Results from the soil assays in 1995 revealed both quantitative and qualitative differences in the microflora recovered from chitin-amended plots compared with unamended plots. Additionally, active inhibition of F. oxysporum colonies by some actinomycetes and bacteria on assay plates with soil from chitin-amended plots was observed. To evaluate the potential of these microorganisms to control Fusarium yellows in celery, two additional treatments, planting transplants of each celery cultivar treated with potential biocontrol agents in plots with or without chitin, were included in 1996.

Twenty isolates each of bacteria and actinomycetes that inhibited F. oxysporum in vitro were grown in 100 ml potato dextrose broth on a shaker for 2 and 5 days, respectively. Broth containing the isolates was mixed in a Waring blender for 8 s and combined. Approximately 1.5 ml was applied to each cell (2.5 by 4 cm) in the transplant tray at 4 weeks and 1 week prior to transplanting in the field. Treated transplants were transferred to chitin-amended and unamended plots at the same time as the establishment of other treatments. Fusarium yellows severity was recorded at harvest as described above. Soil samples collected at transplanting and afterwards at monthly intervals were processed for mi-
croflora, including F. oxysporum, as described above. The data were analyzed using the repeated measures analysis of variance described above.

RESULTS

Effects of chitosan concentration on F. oxysporum f. sp. apii in vitro. All isolates of F. oxysporum f. sp. apii were progressively and uniformly inhibited by concentrations of chitosan up to 3 mg ml⁻¹ (Fig. 1). Concentrations above 3 mg ml⁻¹ did not result in significantly greater inhibition of growth (P = 0.05). The small but insignificant growth observed above 3 mg ml⁻¹ was probably due to clumping of the chitosan that occurred above 3 mg ml⁻¹ (Fig. 2).

Effects of chitin and chitosan on Fusarium yellows. In 1994, incidence of yellows in treatments with chitin plus chitosan was significantly lower than in the unamended plots (Table 1) 60 and 90 days after transplanting (P = 0.05). At 60 days, disease incidence in chitin and chitin-plus-chitosan treatments was significantly less than incidence in chitosan treatments. Similarly, the incidence of Fusarium yellows was significantly reduced by chitin and chitin-plus-chitosan treatments relative to the unamended control (P = 0.05) on cultivar VTR1331 at Oxnard in 1995 (Table 1). At this location, disease incidence in plants treated with chitin plus chitosan was lower than disease incidence in plants treated with either chitin or chitosan alone. However, disease incidence in Conquistador in 1995 at Santa Maria was significantly reduced only in the chitosan treatment (P = 0.05); chitin-plus-chitosan treatments were not significantly different from either the control or chitosan or chitin alone (Table 1).

Disease severity was significantly different between cultivars (P = 0.0001) and between treatments (P = 0.04) in 1996. As expected, disease severity was least in Promise, intermediate in Conquistador, and highest in TU52-70R. Disease severity was reduced by all treatments in cultivar Promise (P = 0.05). In Conquistador, chitin and chitin plus chitosan reduced disease. There were no significant differences in disease severity between treatments in cultivar TU52-70R (Table 2).

There were no significant differences in plant fresh weight between treatments in 1995 or 1996. In 1996, plant fresh weight between cultivars was significantly different (data not shown).

Effects of chitin and chitosan on soil propagules. Overall, the chitin-plus-chitosan treatment significantly reduced the number of F. oxysporum propagules in soil (2.06 × 10⁴ vs. 2.75 × 10¹ CFU g⁻¹ dry soil in control; P = 0.02) in 1994, but within sampling dates significant reduction was observed only at 60 days. The F. oxysporum CFUs were nearly identical on all sampling days. Mean F. oxysporum population prior to treatment application was 2.06 × 10⁴ CFU g⁻¹ dry soil.

In 1995 in Oxnard, populations of F. oxysporum in soil planted with VTR 1331 were not significantly different between treatments or sampling dates. Mean F. oxysporum population prior to treatment application was 3.6 × 10⁴ CFU g⁻¹ dry soil.

In 1996, populations of F. oxysporum propagules were not affected by treatment or cultivar until 90 days after transplanting, when populations were significantly (P = 0.03) lower in the chitin-plus-chitosan and biocontrol-plus-chitin than the biocontrol treatment (data not shown). Mean F. oxysporum population prior to treatment application was 2.16 × 10⁴ CFU g⁻¹ dry soil.

Effects of chitin and chitosan on microbial populations in the soil. Bacteria. In 1995 at Oxnard, populations of bacteria were greater in the chitin and chitin-plus-chitosan treatments than in the unamended plots at 30 and 60 days after transplanting (Fig. 3A). At the end of the season, populations of bacteria in the chitin treatment remained significantly higher than in the control. In Santa Maria, populations of bacteria increased significantly 30 days

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Table 1. Effects of chitin and chitosan on the incidence of Fusarium yellows in 1994 and 1995 at Oxnard and Santa Maria, California

<table>
<thead>
<tr>
<th></th>
<th>TU 52-70R (S)</th>
<th>VTR1331 (M)</th>
<th>Conquistador (M)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treatment</td>
<td>60 days × 90 days</td>
<td>90 days</td>
<td>90 days</td>
</tr>
<tr>
<td>Control</td>
<td>90.5 a</td>
<td>91.7 a</td>
<td>37.7 a</td>
</tr>
<tr>
<td>Chitin</td>
<td>53.3 c</td>
<td>74.8 b</td>
<td>24.3 b</td>
</tr>
<tr>
<td>Chitosan</td>
<td>68.9 b</td>
<td>75.3 b</td>
<td>29.0 ab</td>
</tr>
<tr>
<td>Chitin + chitosan</td>
<td>52.3 c</td>
<td>70.9 b</td>
<td>8.7 c</td>
</tr>
</tbody>
</table>

\( ^\text{S} \) = susceptible cultivar, \( ^\text{M} \) = moderately tolerant cultivar.
\(^\text{a} \) Number of days after transplanting. Values followed by the same letter within columns are not significantly different (P = 0.05) according to a least significant difference test.

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Table 2. Effects of chitin and chitosan on the severity of Fusarium yellows at harvest in 1996 at Santa Maria, California

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Promise (T)</th>
<th>Conquistador (M)</th>
<th>TU52-70R (S)</th>
<th>Mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>0.80 a</td>
<td>3.53 a</td>
<td>4.28 a</td>
<td>2.87 a</td>
</tr>
<tr>
<td>Chitin</td>
<td>0.33 c</td>
<td>2.73 c</td>
<td>4.23 a</td>
<td>2.43 b</td>
</tr>
<tr>
<td>Chitosan</td>
<td>0.30 c</td>
<td>3.46 a</td>
<td>4.40 a</td>
<td>2.72 ab</td>
</tr>
<tr>
<td>Chitin + chitosan</td>
<td>0.43 bc</td>
<td>3.00 bc</td>
<td>4.02 a</td>
<td>2.48 b</td>
</tr>
<tr>
<td>Biocontrol</td>
<td>0.40 bc</td>
<td>3.38 a</td>
<td>4.13 a</td>
<td>2.64 ab</td>
</tr>
<tr>
<td>Biocontrol + chitin</td>
<td>0.57 b</td>
<td>3.25 b</td>
<td>3.97 a</td>
<td>2.59 ab</td>
</tr>
<tr>
<td>Mean</td>
<td>0.47 a</td>
<td>3.22 b</td>
<td>4.17 c</td>
<td></td>
</tr>
</tbody>
</table>

\( ^\text{a} \) = no disease, \( ^\text{b} \) = blight discoloration of root tissue, \( ^\text{c} \) = extensive discoloration of root tissue, \( ^\text{d} \) = discoloration in crown, \( ^\text{e} \) = crown extensively discolored, and \( ^\text{f} \) = dead.
\(^\text{T} \) = tolerant cultivar, \( ^\text{M} \) = moderately resistant cultivar, and \( ^\text{S} \) = susceptible cultivar. Values followed by the same letter within columns are not significantly different (P = 0.05) according to a least significant difference test.
after transplanting in both chitin treatments, and remained significantly higher than the unamended control throughout the season (Fig. 3B). At the end of the season, populations of bacteria in chitosan-alone treatments were not significantly different from those with chitin amendments. Populations of bacteria varied significantly with date of sampling \((P = 0.0001)\) and with treatment \((P = 0.03)\) in 1996. Cultivars did not influence the populations of bacteria in soil \((P = 0.63)\); Fig. 4).

**Fungi.** In 1995 in Oxnard, there were significantly fewer fungi in chitosan-alone and chitin-plus-chitosan treatments than in the unamended control at 30 days. Similarly, in the chitosan-alone and chitin-alone treatments at 60 days, there were fewer fungi in the unamended control (Fig. 3A). *Fusarium* spp. and *Penicillium* spp. were the most common fungi isolated, and there were no apparent differences in the species of fungi from different treatments. In Santa Maria in 1995, there were fewer fungi in the chitin-plus-chitosan plots before treatments were applied and at the end of the experiment (Fig. 3A).

In 1996, populations of fungi were not influenced by either the treatment or cultivar, but time of sampling influenced the type of species recovered \((P = 0.0001)\). The observed differences in populations of fungi between treatments most likely resulted from an unequal distribution in the plots (Fig. 4). Treatment effects for individual cultivars were not consistent. The most frequently isolated fungi were again *Penicillium* spp. and *Fusarium* spp., regardless of the treatment or cultivar.

**Actinomycetes.** In 1995, numbers of actinomycetes were not affected by treatment until 60 days after transplanting (Fig. 3A and B). Subsequently, a significant increase in actinomycetes in the chitin and chitin-plus-chitosan treatments were observed in Oxnard (Fig. 3A). At 60 and 90 days after transplanting in Santa Maria, numbers of actinomycetes in treatments with chitin, chitosan, and chitin plus chitosan were greater than numbers in the unamended plots (Fig. 3B).

Actinomycetes in the 1996 experiment varied significantly with the time of sampling \((P = 0.0001)\) and with treatment \((P = 0.003)\). Cultivars did not affect actinomycete populations, and effects were consistent across all treatments. There were no differences between treatments at any time with the cultivar Promise (Fig. 4).

**Biocontrol potential of the microflora on Fusarium yellows.** Disease severity was significantly different between cultivars \((P = 0.0001)\) and between treatments \((P = 0.04)\) in 1996 (Table 2). Severity of Fusarium yellows was lowest in Promise, intermediate in Conquistador, and highest in TU52-70R. Biocontrol treatments did not reduce either the disease severity (Table 2) or increase plant fresh weight over that provided by chitin alone. No significant reductions in Fusarium propagules were observed in the biocontrol treatment, nor were the differences in populations of microflora consistent over time or cultivar (Fig. 5).

**DISCUSSION**

The higher incidence of Fusarium yellows in recent years in California, possibly caused by increased inoculum density in the soil, prompted this investigation of alternative methods of disease control using chitin and chitosan. The experiments confirmed previous studies (25), which suggested that chitinous materials reduce the severity of some root diseases caused by phytopathogenic fungi. Although some researchers (3,33) reported chitosan to be more effective than chitin in vitro, we found chitin to be the more effective compound in the field. Other advantages of chitin include easier handling and application, lower costs of application and labor, and more consistent beneficial effects.

In 1994 and 1995, yellows incidence was significantly reduced by soil amendments with chitin. In 1996, using cultivars ranging from susceptible to tolerant, chitin reduced severity of yellows in the moderately resistant cultivar Conquistador and tolerant cultivar Promise, but was ineffective in reducing the disease in the susceptible cultivar TU 52-70R. The disease pressure may have been too extreme for this method of control in 1996.

The apparent reduction in disease severity with chitin treatments may have been due to direct inhibition of the pathogen or to a fertilizer effect of the chitin. Other studies have shown disease reduction with

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Fig. 3. Effects of chitin and chitosan on populations of bacteria, fungi, and actinomycetes (log CFU g⁻¹ dry soil) during 1995 (A) at Oxnard with cultivar VTR1331 and (B) at Santa Maria with cultivar Conquistador.
Chitin amendments (8, 25, 32, 34), although some studies (10, 27) reported conflicting results. Chitin decomposition releases volatile compounds such as ammonia which suppress activity of some fungi (19, 32). Specific chitinolytic microflora increase in soils with chitin amendments (10, 15, 24). Chitin decomposition products such as ammonia may also promote growth of celery. While rapid greening of transplants in plots amended with chitin was observed in this study, fresh weight of marketable celery stalks was not significantly different between treatments. Size of individual plants does not appear to increase in chitin-amended plots, but yield is increased by the reduction in yellows incidence.

Our study showed increases in bacteria and actinomycete populations in 1995 and erratic increases in 1996 with chitin treatments. Fungal populations were not consistently affected, suggesting that, in the loam soils of our study, their role as chitin degraders may be less important than bacteria or actinomycetes. The treatment of transplants with isolates of actinomycetes and bacteria with demonstrated biocontrol activity on *F. oxysporum* f. sp. *apii* in vitro did not influence populations of microflora in the soil, indicating that they did not alter the established population equilibrium. Bacteria and actinomycetes, particularly streptomycetes, are known to be the most abundant degraders of chitin in soil (31, 35, 36), and increases in their populations in chitin treatments is indirect confirmation of their role in chitin degradation. Identification of select strains using fatty acid analysis showed that a majority of the actinomycetes were *Streptomyces* spp. (K. V. Subbarao, unpublished data).

By increasing populations of chitinolytic microflora, we hypothesized that the number of pathogenic fungi would decrease as a direct result of feeding activity on the chitinous hyphal walls of *F. oxysporum*. Thus, the number of propagules in the soil was expected to decrease. However, there were no effects of chitin or chitosan on total *F. oxysporum* populations. The assay used to determine the number of *F. oxysporum* propagules in soil provided an estimate of the total population, not of *F. oxysporum* f. sp. *apii* specifically. With techniques currently available, it is not feasible to distinguish *F. oxysporum* f. sp. *apii* within a large soil population of *F. oxysporum*. The availability of color mutants that are similar in growth and pathogenicity to their wild-type counterparts (22) would make this task far simpler, but none are available for this pathogen. Introducing mutant strains would also carry a certain risk, discouraging the use of this technique in commercial production fields. Plating *F. oxysporum* f. sp. *apii* on sorbose medium (9) has been reported to distinguish between pathogenic and non-pathogenic strains of this pathogen. This technique, however, is successful only with a few isolates, and is thus not universally applicable (K. V. Subbarao and J. C. Hubbard, unpublished data). Therefore, even if the treatments reduced *F. oxysporum* f. sp. *apii* populations, we were unable to detect it and cannot indicate that a reduction in pathogenic propagules occurred as a result of treatment.

Suppression of *F. oxysporum* by competition between pathogenic and nonpathogenic species for nutrients (1) or infection sites (29), or by enhanced resistance induced by nonpathogenic species (7, 23, 26), has also been reported to result in decreased disease incidence. Measurement of entire *F. oxysporum* populations did not

Fig. 4. Effects of cultivar (Tall Utah 52-70R, Conquistador, and Promise), chitin, and chitosan treatments on populations of bacteria, fungi, and actinomycetes (log CFU g⁻¹ dry soil) at Santa Maria during 1996.
allow us to determine whether our treatments caused any of these effects. We demonstrated that five test *F. oxysporum* isolates were inhibited by chitosan in vitro. The role of chitosan as both an elicitor of plant defenses (17) and as an active inhibitor of fungal growth is well-documented (3,32). Application of chitosan to tomato plants increases resistance to *F. oxysporum* f. sp. *radicis-lycopersici* (5,6), but results from our field study indicate that dipping transplants in chitosan gave less protection against Fusarium yellows than soil amendment with chitin. Benhamou and Theriault (6) showed that an application of chitosan to tomato roots prior to inoculation with *F. oxysporum* delayed onset of lesion development. In our study, chitosan was generally ineffective in reducing disease. It is possible that chitosan protects the plant in the early stages of growth by triggering plant defense mechanisms (6) but that this protection may be transitory, and under high disease pressure the pathogen may overcome the conferred resistance. Interestingly, disease severity was reduced by chitosan treatment in a tolerant cultivar (Promise), but in the moderately resistant (Conquistador) and the susceptible (TU52-70R) cultivars, chitosan was ineffective.

Chitin significantly reduced the incidence of Fusarium yellows and increased populations of potentially chitinolytic actinomycetes and bacteria in the soil, which in turn may reduce pathogenic populations of *F. oxysporum* f. sp. *apii*. Furthermore, reduction in the incidence of Fusarium yellows may also directly contribute to the reduction in pathogenic populations of *F. oxysporum* f. sp. *apii*. More studies are needed to understand fully the mechanisms involved. Although chitin is readily available from crustacean shell wastes, at current market prices it is not economically viable to use chitin as a soil amendment for Fusarium yellows management.

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LITERATURE CITED