Eliminating Thrips From In Vitro Shoot Cultures of Apple with Insecticides

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Abstract. The insecticides acephate, dichlorvos, and imidacloprid were assayed, using in vitro shoot cultures of apple (Malus ×domestica Borkh.), to determine their phytotoxicity at several concentrations and their effectiveness for eradication of the Western Flower Thrips (Frankliniella occidentalis, Pergande) from infested apple shoot cultures. Commercial formulations of acephate (Orthene), dichlorvos (Vaportape II), and imidacloprid (Admire) and a technical grade of imidacloprid were used in the experiments. For acephate and imidacloprid, concentrations of 1 to 80 mg·L⁻¹ a.i. in shoot culture medium were used, while for dichlorvos, a fumigant, particles of the formulated product containing concentrations of 0.7 to 6.4 mg a.i. were suspended in the head space of the 500-mL glass culture jar. Acephate, dichlorvos, and the technical grade of imidacloprid did not cause phytotoxicity and growth of shoot cultures was unaffected at all treatment concentrations tested for a 6-week treatment period. Imidacloprid (20 mg·L⁻¹ of the commercial formulation) caused chlorosis at the end of the 6-week treatment period. None of the treatments tested resulted in the death of shoots. Thrips were eradicated by acephate or imidacloprid treatments of 5 mg·L⁻¹ and by dichlorvos treatment of 0.7 mg per 500-mL culture jar. Shoot cultures grew normally after the treatment period. Chemical names used: O3,S-dimethyl acetylphosphoramidothioate (acephate), 2,2-dichlorovinyl dimethyl phosphate (dichlorvos), 1-(6-chloro-3-pyridylmethyl)-N-nitroimidazolidin-2-ylidineamine (imidacloprid).

Contamination is the most important cause for losses of in vitro cultures at commercial micropropagation facilities (Cassells, 1986; Boxus and Terzi 1987; Leifert and Waite, 1990; Leifert and Woodward, 1998). Cultures can be contaminated with viruses, bacteria, fungi, yeast, mites and thrips (Knauss and Miller, 1978; Leifert et al., 1994). Thrips and mites often spread microorganisms that contaminate plant tissue cultures. Bacteria are by far the most common and most serious contaminants.

Contamination by thrips is less common but a few invading thrips can develop to high population levels that completely destroy cultures and may often go unnoticed because of their small size. Additionally, thrips may introduce into sterile cultures other contaminants such as bacteria and fungi (Deberch and Maene, 1984; Klocke and Myers, 1984; Blake, 1988). Little information is available concerning control of thrips in in vitro cultures (Klocke and Myers, 1984).

Thrips are common pests of greenhouse and field crops including apple and sweet cherry. They usually feed on actively-growing tissues such as young leaves and terminal buds or flowers and developing fruit. Eggs are laid just below the surfaces of tissues. Thrips’ affinity for tight places, like flower buds, and the ability to deposit eggs in plant tissues may prevent decontamination during surface sterilization when in vitro cultures are initiated and makes eradication more difficult if a contact/respiratory insecticide is used. Thrips frequently gain entry to in vitro cultures through loose-fitting closures of culture vessels when they migrate from nearby infested plants.

Damage done by thrips is not always severe, but in a survey done by Blake (1988), 15% of tissue culture labs had experienced problems with thrips infestation, sometimes quite frequently and with severe consequences. Besides damage done by feeding, thrips may carry fungal spores and bacteria in and on their bodies, inoculating cultures and medium causing serious contamination problems. The presence of thrips in cultures can go undetected for several years, causing losses and reducing the efficiency of commercial labs (Blake, 1988).

The objective of this study was to evaluate the effectiveness of some insecticides with systemic and contact/respiratory modes of action for the control of thrips on apple shoot cultures and develop a protocol for their use to eliminate thrips from infested cultures without harming the cultures.

Materials and Methods

Plant material and culture conditions. Shoot cultures of apple cv. Golden Delicious were maintained on a shoot culture medium of Murashige and Skoog (1962) minerals and vitamins (MS), supplemented with 3.0% benzyladenine (BA), 87.6 mm sucrose and 0.6% Agar (Sigma Chemical Co., St. Louis). Cultures were maintained in 500-mL jars (100 mL medium/jar) closed with plastic lids fitted with 1.0 cm diameter open cell foam plugs to allow for exchange of gases. Additionally, the junction between the lid and the jar was sealed with a plastic wrap. Cultures were incubated at 25 ± 2 °C with a 16-h light/8-h dark photoperiod and light intensity of 18 µmol·m⁻²·s⁻¹ provided by cool-white fluorescent tubes.

Insecticides. The insecticides used in this investigation were acephate, dichlorvos, and imidacloprid. Acephate and imidacloprid are systemic insecticides while dichlorvos has a contact/respiratory mode of action. The insecticidal preparations used were commercial formulations of acephate, Orthene 75% SP (Chevron Chemical Co., Richmond, Calif.) and of dichlorvos, Vaportape II (a.i. = 10%, Hercen Environmental Co., Emigsville, Pa.) and two formulations of imidacloprid, a technical grade (a.i. = 98.3%) and a commercial formulation, Admire (a.i. = 21.4%, Bayer Corp., Kansas City, Mo.).

Phytotoxicity bioassay. Shoot tip explants from well-established shoot cultures (3 weeks after subculture) were transferred to treated medium to determine the phytotoxicity of the insecticides. The technical grade and commercial formulation of imidacloprid and commercial formulation of acephate, were dissolved in dimethyl sulfoxide (Sigma Chemical Co., St. Louis) and added to autoclaved shoot culture medium cooled to 55 °C, to give final concentrations of 1.5, 10, 20, 40 and 80 mg·L⁻¹ a.i. One hundred milliliters of medium was dispensed into 500-mL culture jars. For the dichlorvos bioassay, pieces of Vaportape II measuring 4, 9, 16, 25, or 36 mm² were used. This corresponded to 0.7, 1.6, 2.8, 4.4 and 6.4 mg a.i. of dichlorvos, respectively. One piece of Vaportape II, suspended from the foam plug in the lid, was used per jar. It was sterilized by dipping in 70% alcohol for 20 s. Control cultures were grown on medium without insecticide added.

Shoot tip explants of apple cv. Golden Delicious were inoculated into the various treatments and allowed to grow 6 weeks. The fresh weights of the shoot tip explants were recorded at the start of the incubation period and the fresh weights of shoot clusters that developed from these shoot tip explants were recorded at the end of the 6-week treatment period.

Insecticidal bioassay. Because all shoots remained alive in the phytotoxicity bioassays, experiments were done to determine the insecticidal efficacy of the various compounds using identical treatments to those described above for the phytotoxicity bioassay. Shoot tip explants were cultured in the various treatments for 2 weeks before inocu-
lation with thrips. Inoculation with thrips was accomplished by placing one infested shoot in each culture jar. The number of thrips per infested inoculum shoot averaged 50. This was determined in a separate, preliminary experiment by placing 20 infested shoots in individual sealed plastic bags for 3 weeks. This allowed the shoot to remain healthy but was sufficient time for thrips eggs on the inoculum to hatch. The number of thrips was counted after 3 weeks by washing the thrips off the shoots and bag with ethanol, collecting the ethanol in a petri dish and counting the insects using a stereo microscope. From this experiment, it was determined that an average of 50 adult thrips can develop over a 3-week period. The exact number of thrips inoculated to each culture jar replicate was not crucial since by the end of the treatment period the treatments resulted either in control with no multiplication nor thrip damage symptoms or severe infestations.

Shoot culture medium and incubation conditions were identical to those described above for maintaining the stock cultures. Cultures were observed for thrip damage at 4-week intervals, when subcultured, for a total time period of 12 weeks. This experiment was duplicated. After 12 weeks, shoot clusters of all treatments were removed from the culture jars, placed in Petri dishes which were sealed with Parafilm and observed with a stereo microscope for the presence of live thrips.

Data collection and statistical analysis. Shoot tip explant fresh weights were recorded in the phytotoxicity bioassay experiments at the start and shoot cluster fresh weights at the end of the experiments. The total weight of the five shoot tip explants/shoot clusters in each of the replicate jars was recorded. Also, the lengths of the main shoots in the shoot clusters and the number of shoots >1 cm in each cluster were recorded. The overall condition of the cultures were assessed visually to note untoward detrimental effects on the quality of the cultures caused by the treatments. Visual assessments were used to record reduction in culture quality resulting from chlorosis, tip burn, condition of the stems and leaves, and vigor of the cultures. For the insecticide efficacy tests, shoot damage was assessed visually by estimating the percentage of total shoot surface mottled by feeding. Data were analyzed using analysis of variance (ANOVA), with pairwise comparisons of the means using Bonferroni’s method (Neter et al., 1990).

Results and Discussion

Phytotoxicity bioassay. Treatments with the technical grade of imidacloprid did not significantly affect the means of shoot cluster fresh weights, number of shoots per cluster, or length of the main shoots. Visual assessment of the cultures suggested the treatments did not cause obvious debilitating effects on culture quality (Table 1).

With the commercial formulation of imidacloprid (Admire), the means of shoot cluster fresh weights of treatments at 1 and 80 mg L⁻¹ differed significantly from the means of all other treatments (Table 2). The 20 to 80 mg L⁻¹ treatments induced chlorosis and reduced shoot proliferation, as measured by the number of shoots at the end of the treatment period (Table 2). This adverse effect of Admire on the shoot cultures was attributed to unidentified components other than the a.i. in the commercial formulation since it was not observed in cultures treated with the technical grade. However, even at 80 mg L⁻¹ all shoots survived and grew reasonably well and recovered when transferred to insecticide-free medium. Callus formed on the stems in the 80 mg L⁻¹ treatment.

The means of fresh weights of shoot clusters treated with acepethate (Orthene) was not affected by the treatments except the 10 mg L⁻¹ treatment mean that was significantly higher (Table 3). At the end of the 6-week treatment period, cultures treated with 10 mg L⁻¹ had mean fresh weights 45% greater than that of the control. The 40 and 80 mg L⁻¹ treatments resulted in reduced shoot proliferation but no treatments caused chlorosis (Table 3).

Insecticidal bioassay. Four weeks after inoculation with thrips, shoots treated with all of the insecticidal treatments lacked symptoms

| Table 1. Effect of imidacloprid (technical formulation) on shoot cultures of apple after a 6-week treatment period. |
|---------------------------------------------------------------|
| Treatment (mg L⁻¹) | Shoot quality | Length of main shoots in cluster (cm²) | No. of shoots in cluster | Shoot cluster fresh wt (g) | Estimated shoot damage (%)* |
|--------------------|---------------|----------------------------------------|--------------------------|---------------------------|-----------------------------|
| 0                  | normal        | 4.0 ± 0.5 a                           | 6.0 ± 2.4 a               | 8.5 ± a                   | 100                         |
| 5                  | normal        | 4.2 ± 0.6 a                           | 5.8 ± 1.5 a               | 7.1 ± a                   | 50                          |
| 10                 | normal        | 4.4 ± 0.5 a                           | 5.6 ± 1.6 a               | 6.5 ± a                   | 0                           |
| 20                 | normal        | 4.3 ± 0.6 a                           | 5.6 ± 1.4 a               | 6.8 ± a                   | 0                           |
| 40                 | normal        | 4.4 ± 0.6 a                           | 6.4 ± 1.3 a               | 7.3 ± a                   | 0                           |
| 80                 | normal        | 5.1 ± 0.5 a                           | 7.5 ± 1.0 a               | 7.3 ± a                   | 0                           |

*Visual assessment suggested a reduction in quality for all treatments (Table 4). Shoot proliferation was significantly reduced with the 4.4 and 6.4 mg/500 mL culture jar treatments.

Table 2. Effect of imidacloprid (commercial formulation, Admire) on shoot cultures of apple after a 6-week treatment period.

| Treatment (mg L⁻¹) | Shoot quality | Length of main shoots in cluster (cm²) | No. of shoots in cluster | Shoot cluster fresh wt (g) | Estimated shoot damage (%)* |
|--------------------|---------------|----------------------------------------|--------------------------|---------------------------|-----------------------------|
| 0                  | normal        | 4.0 ± 0.7 a                           | 7.4 ± 1.8 a               | 8.5 ± 3.1 a               | 100                         |
| 1                  | normal        | 4.0 ± 0.7 a                           | 5.2 ± 1.3 a               | 5.2 ± 1.5 b               | 70                          |
| 5                  | normal        | 4.1 ± 0.7 a                           | 5.4 ± 1.0 a               | 6.6 ± 0.9 a               | 0                           |
| 10                 | normal        | 13.4 ± 0.3 a                          | 5.8 ± 1.3 a               | 6.3 ± 1.2 a               | 0                           |
| 2                  | chlorotic     | 3.4 ± 0.4 a                           | 3.6 ± 0.5 b               | 6.0 ± 0.9 a               | 0                           |
| 40                 | chlorotic     | 3.2 ± 0.4 a                           | 3.8 ± 1.5 b               | 5.7 ± 1.4 a               | 0                           |
| 80                 | chlorotic with callus | 3.1 ± 0.5 b | 3.2 ± 1.5 b | 4.5 ± 0.7 b | 0 |

*Means of length of main shoots in clusters, number of shoots in clusters >1 cm, and shoot cluster fresh weight.

Table 3. Effect of acepethate (commercial formulation, Orthene) on shoot cultures of apple after a 6-week treatment period.

| Treatment (mg L⁻¹) | Shoot quality | Length of main shoots in cluster (cm²) | No. of shoots in cluster | Shoot cluster fresh wt (g) | Estimated shoot damage (%)* |
|--------------------|---------------|----------------------------------------|--------------------------|---------------------------|-----------------------------|
| 0                  | normal        | 4.5 ± 0.6 a                           | 7.2 ± 1.2 a               | 4.9 ± 0.6 a               | 100                         |
| 1                  | normal        | 4.4 ± 0.7 a                           | 6.8 ± 1.3 a               | 6.5 ± 0.8 a               | 5                           |
| 5                  | normal        | 4.4 ± 0.6 a                           | 6.4 ± 1.0 a               | 6.7 ± 1.4 a               | 0                           |
| 10                 | normal        | 3.8 ± 0.2 a                           | 5.8 ± 0.7 a               | 7.1 ± 2.0 b               | 0                           |
| 20                 | normal        | 3.8 ± 0.2 a                           | 5.2 ± 1.3 a               | 5.6 ± 0.9 a               | 0                           |
| 40                 | normal        | 3.9 ± 0.6 a                           | 4.4 ± 1.0 b               | 5.2 ± 1.1 a               | 0                           |
| 80                 | normal        | 3.8 ± 0.3 a                           | 5.6 ± 1.2 b               | 5.6 ± 1.2 a               | 0                           |

*Means of length of main shoots in clusters, number of shoots in clusters >1 cm and shoot cluster fresh weight.

Visual estimate of shoot surfaces mottled by feeding.

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of thrip damage. In contrast, control cultures with no insecticide treatment had 100% shoot damage with death, indicated by browning and loss of cell turgor of most tissues not protected by submersion in the medium (Fig. 1). Eight weeks after inoculation, some of the treated shoots had developed thrip damage. Five percent of shoots cultured in the 1 mg L⁻¹ treatments of both acephate and technical grade of imidacloprid, developed symptoms of thrip damage, while 30% to 40% damage occurred to shoots cultured in the 1 mg L⁻¹ commercial formulation of imidacloprid (Admire). None of the other treatments showed any symptoms of thrip damage.

After 12 weeks, 50% damage had occurred in the 1 mg L⁻¹ treatments with the technical grade and 70% damage with the commercial formulation of imidacloprid, but damage remained at 5% in the 1 mg L⁻¹ acephate treatment. This suggested that with the 1 mg L⁻¹ acephate treatment, shoot damage was caused initially by the thrips inoculated into the culture but thrips multiplication was prevented. No symptoms of thrip damage were seen in any of the treatments at concentrations of 5 mg L⁻¹ or greater. When examined under the microscope, live thrips were observed only on shoots treated with 1 mg L⁻¹ of the technical grade and commercial formulation of imidacloprid. No live thrips were observed on shoots treated with 1 mg L⁻¹ acephate or on shoots of any of the other treatment. Acephate and imidacloprid are systemic insecticides with low mammalian toxicity and low phytotoxicity (Bath Press, 1994). This, combined with the results presented above, indicate they are good candidates for use in tissue culture to control or eliminate thrip infestations. They were effective at the low concentrations used in these experiments and had no apparent deleterious effects on apple cultures even when cultures were treated for three subculture periods (12 weeks). One milligram per liter of both the technical grade and commercial formulation of imidacloprid were not effective in eradicating the thrips. However, eradication was achieved at 5 mg L⁻¹ and above.

No symptoms of thrip damage were seen at any treatment level in cultures treated with dichlorvos (Vaportape II). However, visual assessment of cultures suggested a progressive decline in quality towards the end of the experiment at the two highest treatment concentrations. Shoots tips in the 6.4 mg treatment died and associated brown tissue developed (Fig. 1). Dichlorvos has a respiratory, contact and stomach mode of action. It has high volatility and is known to rapidly kill insects, but it also has high mammalian toxicity and may also be phytotoxic to some plants (Bath Press, 1994). Dichlorvos was highly phytotoxic to sweet cherry cultures in some preliminary experiments (results not shown). With apple cultures, 0.7 mg dichlorvos/500-mL culture jar controlled thrips with 100% effectiveness, with little negative effect on the growth of shoot cultures or their quality assessed visually, but prolonged exposure (>6 weeks) to high concentrations was deleterious. The low dose of 0.7 mg/jar was found to give 100% control of thrips in our studies and may or may not be a health hazard depending on the number of jars treated, ventilation and other factors.

The thrips used as inoculums in this study appeared to be free of bacteria and fungal spores since no cultures became contaminated with bacteria or fungi. Shoot cultures with thrip damage symptoms were always infested with large numbers of thrips that had multiplied from the original inoculum. By the end of the third subculture (12 weeks), shoot cultures were either infected because the treatment was not effective or were thrip-free and did not become infested when transferred to insecticide-free medium indicating that the insects introduced with the inoculum plus insects developing from eggs on the inoculum had been killed.

To achieve complete eradication of thrips from in vitro cultures with insecticides used in this way, the duration of treatment should be long enough so that insects emerging from eggs are killed and do not reinitiate the infestation.

### Table 4. Effect of dichlorvos (commercial formulation, Vaportape II) on shoot cultures of apple after a 6-week treatment period.

| Treatment (mg L⁻¹) | Shoot quality | Length of main shoots in cluster (cm) | No. of shoots in cluster | Shoot cluster fresh wt (g) | Estimated shoot damage (%) |
|-------------------|---------------|---------------------------------------|--------------------------|----------------------------|-----------------------------|
| 0.7               | small leaves  | 3.9 ± 0.7 ab                          | 4.0 ± 0.9 a              | 2.8 ± 0.6 a                | 0                           |
| 1.6               | small leaves  | 2.8 ± 0.5 c                           | 3.6 ± 0.8 a              | 3.0 ± 0.6 a                | 0                           |
| 2.8               | small leaves  | 3.0 ± 0.6 c                           | 4.0 ± 0.9 a              | 2.7 ± 0.4 a                | 0                           |
| 4.4               | small leaves  | 3.0 ± 0.4 c                           | 3.2 ± 1.2 b              | 2.9 ± 0.3 a                | 0                           |
| 6.4               | small leaves  | 2.8 ± 0.5 c                           | 3.2 ± 2.2 b              | 2.6 ± 0.1 a                | 0                           |

*Means of length of main shoot in cluster, number of shoots in cluster >1 cm and shoot cluster fresh weight.

*Mean separation within columns by Bonferroni’s pairwise comparison at 0.05. Means in each column followed by the same letters are not significantly different.

*Visual estimate of shoot surfaces mottled by feeding.
Insecticidal treatment of our apple cultures for two subculture periods (6 weeks) resulted in eradication of the thrips and cultures returned to optimum growth and quality in the subculture period following cessation of treatment. The chemicals used in these experiments are, to our knowledge, not registered for use with in vitro plant cultures.

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