Paclobutrazol Distribution following Application to Two Media as Determined by Bioassay

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Abstract. A broccoli (Brassica oleracea var. botrytis L.) seedling bioassay was used to measure paclobutrazol activity and distribution in two growing media following drench or subirrigation applications. The bioassay exhibited a saturation-type response curve for paclobutrazol concentrations up to 1000 μg·L⁻¹ in solution and 100 μg·L⁻¹ in the media. The concentration of paclobutrazol required to achieve one-half of the maximum observed bioassay activity was 3-fold as high in bark-based commercial potting medium as in a peat-based medium. Less than 2% of applied paclobutrazol leached out during the drench application despite the collection of up to 50 mL of leachate per 120 mL of the solution (1000 μg·L⁻¹) that was applied per 15-cm pot. Immediately following drench application, paclobutrazol concentrations in both media were highest in the uppermost 2.5 cm and decreased downward. By 3 weeks after treatment, drench-applied paclobutrazol had moved into lower depths. Distribution of paclobutrazol was limited to the bottom 2.5 cm of media when applied as a subirrigation soak. Chemical name used: (±)-(R*,R*)-β-[4-(3-chlorophenyl)methyl]-α-[1,1-dimethyl]-1H-1,2,4-triazole-1-ethanol (paclobutrazol).

Paclobutrazol is a growth retardant used in commercial floriculture to control plant size and quality. Although commonly applied as a foliar spray, it also can be applied as a medium drench. This increases its activity and results in fewer problems with excessive stunting or delayed flowering.

Factors affecting paclobutrazol activity and movement in soil and soilless media have received very little attention compared to the multitude of studies on its distribution in plants (Davis et al., 1988). This inattention may be attributed in part to the exacting laboratory procedures required for paclobutrazol measurement (Reed, 1988). The use of bioassays for quantifying the activity of growth retardants has proved useful in a limited number of studies. Hunter and Proctor (1990) described a grape (Vitis sp.) auxillary shoot bioassay for paclobutrazol. They found specific leaf mass was log-linearly correlated with soil paclobutrazol concentration from 1 to 1000 μg·L⁻¹ of soil. McArthur and Eaton (1987) used a bioassay with rape (Brassica napus L.) and observed residual paclobutrazol activity in the soil 11 weeks after a foliar application to strawberry (Fragaria ×ananassa Duchesne).

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Frequent movement in soil and soilless media have received very little attention compared to the multitude of studies on its distribution in plants (Davis et al., 1988). This inattention may be attributed in part to the exacting laboratory procedures required for paclobutrazol measurement (Reed, 1988). The use of bioassays for quantifying the activity of growth retardants has proved useful in a limited number of studies. Hunter and Proctor (1990) described a grape (Vitis sp.) auxillary shoot bioassay for paclobutrazol. They found specific leaf mass was log-linearly correlated with soil paclobutrazol concentration from 1 to 1000 μg·L⁻¹ of soil. McArthur and Eaton (1987) used a bioassay with rape (Brassica napus L.) and observed residual paclobutrazol activity in the soil 11 weeks after a foliar application to strawberry (Fragaria ×ananassa Duchesne).

Materials and Methods

Bioassay procedure and standard curves. The two media tested were: a) Metro Mix 300 (MM300) (O.M. Scotts, Marysville, Ohio) composed (volume basis) of 20% sphagnum peat, 30% vermiculite, 40% composted pine bark, 10% processed pine bark ash, and b) Vergro Klay Mix A (VKMA) (Verlite, Tampa, Fl.) composed of 50% sphagnum peat, 20% vermiculite, 20% perlite, and 10% calcined clay. Broccoli was chosen as a bioassay crop because preliminary trials indicated that it was sensitive to the growth retardants in the range of concentrations typically found with media applications. To establish a standard response curve 100-mL solutions of paclobutrazol (0, 10, 25, 50, 75, 100, 250, 500, 750, 1000, and 2000 μg·L⁻¹) were mixed with 1-L batches of media premoistened with tap water. Each batch of medium was mixed uniformly by shaking and massaging in a polyethylene bag and was placed into twelve 4 × 4 cm “cell-pack” cells each 5 cm deep. Six broccoli seeds were planted in each cell, and the cell-packs were placed in a greenhouse on capillary matting. Seedlings were thinned to two per cell 5 d after sowing. Average daily minimum and maximum temperatures were 19 and 30°C, respectively. The cells were subirrigated continuously with fertilizer solution (20N–4.4P–16.6K with micro-nutrients; O.M. Scotts Co., Marysville, Ohio), with N at 150 mg·L⁻¹. A bioassay procedure for estimating paclobutrazol concentration in solutions was accomplished by pipetting 2 mL of standard solution into each of 12 vermiculite-filled cells. Nine standard concentrations ranging from 0 to 1000 μg·L⁻¹ were included to establish a standard response curve. Broccoli was planted in the cells and the test carried out as described above.

After 12 d, seedlings from both bioassay procedures were cut at the medium surface and the hypocotyl length from the base to the cotyledonal node was measured to the nearest mm. Bioassay activity was expressed as inhibition relative to the control:

\[ I_1 = \frac{(H_0 - H_1)}{H_0} \]

where \( I_1 \) = inhibition at concentration x, \( H_0 \) = hypocotyl length at concentration x, and \( H_1 \) = mean hypocotyl length of the controls. A derivative-free (DUD) nonlinear regression procedure (PROC NLIN) (SAS Institute, 1987) was used to fit a Michaelis–Menten equation (Causton, 1983) to the standard curve data for each medium.

Media effects on drench distribution (Expt. 1). To determine the distribution of paclobutrazol following drench treatments, 15-cm (1400-mL) pots were filled 11 cm deep with either VKMA or MM500. The filled pots were watered with tap water until the media were saturated. After settling, medium depth was ≈10 cm. All pots were allowed to stand for 24 h, at which time 120 mL of paclobutrazol at 0 or 1000 μg·L⁻¹ were applied to the surface of each pot (0 or 120 μg·pot) (eight pots per treatment). Leachates from the drench application were collected from each pot and their volumes measured.

Paclobutrazol in the leachates was bioassayed immediately by pipetting 2 mL of leachate from each pot onto each of 12 vermiculite-filled cells. This procedure was carried out at the same time and in the same
manner as described previously for the standards. Paclobutrazol activity in the leachate was expressed as inhibition using the hypocotyl length of the 0 µg·L⁻¹ treatment as H₀. The Michaelis-Menten regression equation fitted to the standard curve data was subsequently used to estimate paclobutrazol concentration in the leachates.

Treated medium from each pot was divided into four depths: 0–2.5, 2.5–5, 5–7.5, and 7.5–10 cm. For the bioassay, each sample was mixed and placed into six cells. This bioassay was conducted at the same time and under the same conditions as the standard curve series described previously. For each medium, inhibition was calculated separately for each sample depth using the control hypocotyl length of the 0 µg·L⁻¹ rate for a given sample depth as H₀. Analysis of variance (ANOVA) was used to evaluate treatment effects on hypocotyl length.

A completely randomized design was used for analyzing leachate data, while a split-plot design was used to analyze media data. Drench concentration was the main plot factor and layer depth was the subplot factor. Because inhibition was calculated using the 0 µg·L⁻¹ treatment results, only media and sample depth were included as treatment effects in the ANOVA for this variable. Treatment means for inhibition were used to estimate paclobutrazol concentration according to bioassay standard curve regression equations.

**Effect of application method on paclobutrazol distribution** (Expt. 2). Three rooted cuttings of chrysanthemum [Dendranthema ×grandiflora (Ramat.) cv. Nob Hill] were planted per 15-cm (1400-mL) pot filled with VKMA. Plants were grown in a greenhouse with light levels between 800 and 1000 µmol·m⁻²·s⁻¹, and average minimum and maximum temperatures of 19 °C and 28 °C, respectively. Plants were fertilized at every watering with a nutrient solution (20N–4.4P–16.6K with minor elements; O.M. Scotts, Marysville, Ohio) containing N at 300 mg·L⁻¹. Paclobutrazol concentrations of 0 or 500 µg·L⁻¹ were applied as either a drench or through a subirrigation soak in a randomized block design with four blocks and four pots per block. All pots were treated 10 d after transplanting when roots had reached the edges of the pot. Subirrigation treatments were accomplished by placing preweighed pots in trays maintained 2-cm deep with either 0 or 500 µg·L⁻¹ of paclobutrazol solution. After 20 min, pots were removed and weighed to determine the volume of solution applied by subirrigation. Equivalent volumes were then applied to the surface for drench treatments. Two pots per block were sampled immediately after treatment and the distribution of paclobutrazol among the four sample depths was determined by the bioassay procedure (Expt. 1).

For the drench treatments only, the remaining two pots per block were sampled three weeks after treatment and the paclobutrazol bioassay was repeated. Separate standard curves were included for both sampling dates. We excluded the subirrigation treatments for the 3-week sampling because the use of overhead watering prevented our obtaining any useful information.

Statistical analysis was identical to Expt. 1 except that the main plot factor for the first sampling date was application method instead of medium.

**Results and Discussion**

**Bioassay and standard curves.** A saturation-type response was observed for both media (Fig. 1) and leachate (Fig. 2) bioassay standards. A Michaelis-Menten equation (Causton, 1983) was selected to fit the standard curve data:

\[ y = \frac{ax}{b + x} \]

where \( y \) = inhibition, \( x \) = media paclobutrazol concentration (µg·L⁻¹), and \( a \) and \( b \) are parameters. This equation was chosen because the parameters \( a \) and \( b \) have useful interpretations. Parameter \( a \) is the horizontal asymptote, which represents the maximum inhibition observed, while \( b \) represents the concentration at which half the maximum inhibition was observed. Therefore, the parameters could be used to compare paclobutrazol activity in MM500 vs. VKMA. The 95% confidence intervals for parameter \( a \) estimates were 0.874 ± 0.034 for VKMA and 0.87 ± 0.03 for MM500. This indicates that predicted maximum inhibition for the two media did not differ. However, parameter estimates for \( b \) were 27.4 ± 6.4 for MM500 and 9.5 ± 1.5 for VKMA, indicating that the concentration of paclobutrazol required to achieve one-half maximum observed inhibition was nearly 3x as high in MM500 as in VKMA.
Reduced efficacy of paclobutrazol (Million et al., 1998) and ancymidol (Bonaminio and Larsen, 1978; Tschabold et al., 1975) in pine bark–based media has been reported previously. Because MM500 and VKMA differ relative to several components, the reduced activity in MM500 cannot be directly attributed to the bark in this study. However, the procedure for establishing the standard curve was conducted under nonleaching conditions, and the reduced efficacy in MM500 cannot be explained by increased leaching losses caused by the bark component, as proposed by Bhat and Tayama (1990) with ancymidol.

Media effects on drench distribution (Expt. 1). Paclobutrazol concentration was lower in leachates from MM500 than in those from VKMA (Table 1). Considering the leachate volumes collected, total estimated leaching losses of paclobutrazol were 1 and 1.9 µg for MM500 and VKMA, respectively. These totals represent <2% of the 120 µg of paclobutrazol applied, which indicates that leaching during application should not cause a significant loss in chemical efficacy.

Paclobutrazol activity was not uniformly distributed in either media immediately following drench applications (Table 2). Bioassay results on the different sample layers indicated that inhibition (paclobutrazol activity) was highest in the uppermost 2.5 cm of both media and decreased with depth. This decrease with depth was more pronounced in MM500 than in VKMA (Table 2). Because activity was lower in MM500 than in VKMA for a given concentration of paclobutrazol (Fig. 1), the estimated concentrations of paclobutrazol at each sample depth for both media did not differ significantly (Fig. 3).

The estimated concentration of paclobutrazol was 3 to 4x as high in the uppermost 2.5-cm layer than in the lower depths for both media (Fig. 3), indicating that paclobutrazol was being retained by the upper layers of media during the drench application. The type of growing medium affected paclobutrazol activity more than the initial distribution of the chemical during the drench application.

**Effect of application method on paclobutrazol distribution (Expt. 2).** The standard curve equations used to estimate media concentration of paclobutrazol were:

\[
y = 0.853x/(9.5 + x); \quad r^2 = 0.89
\]

for the first sample date bioassay, and

\[
y = 0.80x/(9.5 + x); \quad r^2 = 0.86
\]

for the second. These standard curves were similar to that observed in Expt. 1 (Fig. 1).

The drench application method resulted in more uniform distribution of paclobutrazol activity than did the subirrigation method (Table 3). For the drench application, bioassay activity was distributed mostly in the upper 5 cm of medium with highest activity observed in the uppermost 2.5 cm. The average application volume was 158 mL/pot, which is 30% more than the 120 mL recommended by the product label. A similar pattern of paclobutrazol distribution was observed in Expt. 1.

For the subirrigation soak method of application, the procedure for establishing the standard curve equations used to estimate media concentration of paclobutrazol were:

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cation, very little paclobutrazol was detected above the bottom 2.5 cm of the medium when the medium was sampled immediately after treatment. Upward capillary movement of solution during subirrigation apparently resulted in more effective adsorption by the medium than was evident for the drench application method.

For the 3-week posttreatment period, during which pots were surface-irrigated, paclobutrazol activity in the medium remained relatively constant (Fig. 4). However, estimated concentrations decreased in the top 2.5-cm layer of medium and increased in the lower three layers, indicating that paclobutrazol was being redistributed to the lower portions of the medium.

The bioassay procedure described in this report should be useful in future studies with growth retardants. The bioassay results are obtained within 2 weeks, as broccoli seedlings emerge quickly and are relatively nondemanding in their growth requirement. Furthermore, many seedlings can be grown in a small area, which reduces the variability inherent in bioassay procedures.

### Literature Cited


### Table 3

Broccoli seedling bioassay of Vergro Klay Mix A, a peat-based medium containing no bark, sampled immediately after applying paclobutrazol by drench or subirrigation methods to chrysanthemums in 15-cm pots (Expt. 2).

<table>
<thead>
<tr>
<th>Application method</th>
<th>Paclobutrazol concn (µg·L⁻¹)</th>
<th>Sample depth (cm)</th>
<th>Broccoli hypocotyl length (cm)</th>
<th>Inhibition¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drench</td>
<td>0</td>
<td>0–2.5</td>
<td>2.3</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>2.5–5</td>
<td>2.2</td>
<td></td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>5–7.5</td>
<td>2.4</td>
<td></td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>7.5–10</td>
<td>2.4</td>
<td></td>
<td>0.75</td>
</tr>
<tr>
<td></td>
<td>1000</td>
<td>0.6</td>
<td>1.0</td>
<td>0.55</td>
</tr>
<tr>
<td></td>
<td>2.5–5</td>
<td>2.1</td>
<td>0.13</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>5–7.5</td>
<td>2.2</td>
<td>0.10</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>7.5–10</td>
<td>2.3</td>
<td>0.01</td>
<td>---</td>
</tr>
<tr>
<td>Subsurface</td>
<td>0</td>
<td>0–2.5</td>
<td>2.3</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>2.5–5</td>
<td>2.3</td>
<td></td>
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</tr>
<tr>
<td></td>
<td>5–7.5</td>
<td>2.3</td>
<td></td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>7.5–10</td>
<td>2.4</td>
<td></td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>1000</td>
<td>0.9</td>
<td>0.63</td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>0–2.5</td>
<td>0.19</td>
<td></td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>2.5–5</td>
<td>0.07</td>
<td></td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>5–7.5</td>
<td>0.08</td>
<td></td>
<td>---</td>
</tr>
<tr>
<td></td>
<td>7.5–10</td>
<td>0.63</td>
<td></td>
<td>---</td>
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<tr>
<td></td>
<td></td>
<td>HSD 0.05 (compare mean concn within method)</td>
<td>0.22</td>
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<tr>
<td></td>
<td></td>
<td>HSD 0.05 (compare all means)</td>
<td>0.10</td>
<td></td>
</tr>
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</table>

¹An average of 158 mL was applied via subirrigation; equivalent drench volumes were applied.

Inhibition was calculated as the reduction in broccoli seedling hypocotyl length relative to the 0 µg·L⁻¹ rate for a given method and sample depth.

### Fig. 4

Distribution of paclobutrazol in Vergro Klay Mix A, a peat-based medium, sampled immediately (●) or 3 weeks after (■) paclobutrazol drench application of 120 mL at 500 µg·L⁻¹ (Expt. 2).