Alterations in Anatomy and Ultrastructure of Pecan Leaves Treated with Propiconazole during Shoot Expansion

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Abstract. Propiconazole, a triazole fungicide, has been reported to inhibit leaf expansion in pecan [Carya illinoensis (Wangenh.) K. Koch] trees when applied under field conditions. This study was conducted to determine the effect of propiconazole on pecan leaf morphology and structure using light and transmission electron microscopy. Mature pecan trees were sprayed once or three times per week from budbreak to pollen maturity. Fungicide sprays resulted in significantly reduced leaf area. Compared to controls, leaves from propiconazole-treated shoots had alterations in cell arrangement, and cortical tissue (Wang and Dunlap, 1994). Paclobutrazol [3RS,3RS]-1-(4-chlorophenyl)-4-propyl-1,3-dioxolan-2-yl (methyl)-1H-1,2,4-triazole (propiconazole).

Proper timing and good coverage of fungicide applications are very important for successful pecan ([Carya illinoensis] production (Ellis et al., 1997; Sparks, 1995). Current spray programs are designed to assure that a protective fungicide shield is consistently present, particularly during the early season to protect rapidly developing young foliage (Ellis et al., 1997; Graves and Diehl, 1991). Recently, climatically-based fungicide applications have been proposed (Sparks, 1995) and are used frequently. However, calendar-based procedures are still in use, with sprays scheduled every 10 to 14 d from budbreak through pollination (Ellis et al., 1997). Under extremely wet weather conditions, intervals between sprays may be shorter.

Propiconazole belongs to the group of triazole derivatives related to sterol biosynthesis inhibitors (Steffens, 1988), and is one of the primary early season fungicides used in pecan (Ellis et al., 1997; Reilly and Wood, 1996). These compounds bind to the cytochrome P450 component of the C-14 demethylase and inhibit 14 α-methylation, and thus function as fungicides (Coolbaugh and Hamilton, 1976; Coolbaugh et al., 1978; Worthington, 1989) and/or plant growth regulators (Buchaner and Röhner, 1981; Burden et al., 1987; Köller, 1987) primarily by reducing gibberellin biosynthesis.

Uniconazole [(E)-1-(p-chlorophenyl)-4,4-dimethyl-2-(1,2,4-triazol-1-yl)-1-penten-3-ol] and paclobutrazol [(2RS,3RS)-1-(4-chlorophenyl)-4,4-dimethyl-2-(1H-1,2,4-triazol-1-yl)pentan-3-ol] are triazole compounds that are used widely in horticulture as growth retardants. The morphological and cytological effects of these growth inhibitors have been documented in a number of crops and include suppressing shoot elongation and limiting the rate of leaf production (Wang and Gregg, 1989, 1994; Wang et al. 1992). Uniconazole applications in hibiscus (Hibiscus rosa-sinensis L.) resulted in shorter pedicels with larger pith, vascular, and cortical tissue (Wang and Dunlap, 1994). Paclobutrazol application resulted in thicker leaves and reduced stem diameter in ‘Lillian Hoek’ chrysanthemum [Dendranthema ×grandiflora (Ramat.) Kitam. (syn. Chrysanthemum ×morifolium Ramat.)] (Burrows et al., 1992), and decreased leaf length in wheat (Triticum aestivum L.) as a result of cell length reduction (Tonkinson et al. 1995).

Applications of triazole fungicides have exhibited adverse effects on various higher plants. In apple [Malus sylvestris (L.) Mill var. domestica (Borkh.) Mansf. ‘Cox’s Orange Pippin’], decreased tree growth (Church et al., 1984; Steffens, 1988), decreased ovule longevity, and reduced ovary size (Williams et al., 1987) were observed. Reilly and Wood (1996) found that propiconazole sprays suppressed leaf expansion in young pecan seedlings, and reported a progressive decline in leaf area as dosage increased. In our previous studies with pecan (He and Wetzstein, 1994), propiconazole applied during shoot development limited shoot growth, leaf expansion, catkin elongation, and pollen development. Nonetheless, the cytological events associated with the inhibitory effects of propiconazole on leaf growth have not been ascertained. In the present study, we describe differences in the anatomy and ultrastructure of pecan leaves following propiconazole applications under field conditions.

Materials and Methods

Plant material and field applications and measurements. Mature, ‘Desirable’ pecan trees, located at the University of Georgia Horticulture Farm near Watkinsville, Ga., were used for this investigation. Terminal shoots used in the study were selected at budbreak for uniformity (i.e., from the outer portion of the canopy, with a caliper of ≈1 cm, and a length of 25 to 35 cm). Five shoots were randomly assigned per treatment. Leaves and shoots were sprayed with propiconazole, in the form of Orbit (Ciba-Geigy, Greensboro, N.C.) at the recommended rate of 0.3 mL·L⁻¹ in water (0.125 mg·L⁻¹ of propiconazole), either once or three times weekly. Control shoots were sprayed with water. Sprays were applied with a hand-held spray bottle until runoff. Applications were from budbreak to pollen maturity, a period of ≈1

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The experiment was repeated a second year on different trees in the same orchard to confirm anatomical observations.

**LEAF AREA AND SHOOT LENGTH MEASUREMENTS.** Current season’s shoots were harvested after leaf expansion had ceased. Leaf area measurements were taken using a leaf area meter (LI-3000; LI-COR, Lincoln, Nebr.).

**PREPARATION OF LEAF SAMPLES FOR MICROSCOPY.** For anatomical and ultrastructural evaluations, tissues were sampled from leaves located midway along the shoot, after leaf expansion had ceased. Leaves in pecan are pinnately compound. A leaflet from the midportion of the leaf was selected, and tissue samples were obtained from the central region of the blade adjacent to the main veins of the leaflet. Tissues were prepared for light (LM) and transmission electron microscopy (TEM) as described previously for pecan leaves (Wetzstein and Sparks, 1983). Briefly, samples were fixed in 2% glutaraldehyde in 0.1 M cacodylate buffer (pH 7.2), postfixed with 1% osmium tetroxide in the same buffer, and dehydrated in a graded ethanol series. The samples were infiltrated and embedded in Spurr’s (1969) low viscosity resin (EM Sciences, Fort Washington, Pa.). For LM, 1-µm-thick sections were cut with an ultramicrotome (MT6000-XL; RMC, Inc., Tucson, Ariz.) and collected on glass microscope slides coated with gelatin and chrome-alum. Sections were stained with 1% pergamon solution (Fisher Scientific, Inc., Springfield, N.J.), and mounted in a medium containing glycerol and Tris-HCl (He et al., 1995). The slides were examined with a microscope (Axioscop; Carl Zeiss, Thornwood, N.Y.) equipped for differential interference contrast. Leaf thickness, heights of palisade and spongy mesophyll regions, and number of palisade cells along the leaf sections were measured. Ten different leaf samples were evaluated per treatment, and mean values per leaf were obtained from measurements of five different microscopic fields.

For TEM, thin sections (∼80 nm) were obtained also using an ultramicrotome, and collected on Formvar-coated, gilded copper slot grids (Ted Pella, Redding, Calif.) and placed on Formvar-coated bridges to dry. The sections were poststained with 4% (w/v) aqueous uranyl acetate and Reynolds (1963) lead citrate. Sections were examined with a transmission electron microscope (Zeiss EM 902A; LEO Electron Microscopy, Thornwood, N.Y.) at 80 kV.

**STATISTICAL ANALYSES.** Numerical data were subjected to analysis of variance procedures and means separated by Duncan’s multiple range test using general linear model procedures of SAS for personal computers (SAS Inst., Inc., 1990). For leaf area, data were transformed with natural logarithms.

### Results

Effects of propiconazole on shoot growth were apparent within 2 weeks after initial sprays, and persisted in affected shoots. Shoots treated with propiconazole were substantially shorter (data not presented), and leaf expansion was severely reduced (Table 1). The mean area of leaves sprayed weekly with propiconazole during expansion was only 54% of water controls. The inhibition symptoms were more severe in shoots treated more frequently. Leaves sprayed three times per week had a leaf area that was only 48% of control leaves.

![Fig. 1. Light micrographs of transverse sections of leaves from control and propiconazole-treated shoots of pecan. (A) Leaf from water-sprayed control shoot, (B) leaf from shoot treated once a week with propiconazole, and (C) leaf from shoot treated three times a week with propiconazole. Scale bar in (C) = 50 µm and applies to all figures.](image-url)
frequently had peltate, glandular trichomes that were more numerous on the abaxial or lower surface. The mesophyll was differentiated into a palisade and spongy parenchyma with dominant intercellular spaces, especially in the spongy region. Leaves treated with propiconazole exhibited marked differences which were more extensive with the higher spray frequency (Fig. 1B and C). The mesophyll of leaves treated with propiconazole was composed of more closely packed cells and had fewer intercellular spaces, and the number of palisade cells per 100-µm cross-sectional length was significantly greater (Table 1). Leaves sprayed at the higher frequency had 1.3× more palisade cells than controls. Neither total leaf thickness nor thickness of spongy and palisade cell layers were significantly affected by sprays.

Electron micrographs of mesophyll tissues from control (Fig. 2A) and propiconazole-sprayed leaves (Fig. 2B and C) illustrate differences in palisade cell organization. In control leaves, the palisade cells were aligned primarily in a single layer with cells of variable lengths interspersed with intercellular spaces. Cells had large vacuoles. More internal cells (from the adaxial surface) were elongated, but loosely packed. Palisade cells from leaves sprayed with propiconazole were more uniformly columnar in shape and length, and were more closely packed (Fig. 2B and C). Leaves sprayed at the higher frequency had two distinct, compactly arranged palisade layers with few intercellular spaces (Fig. 2C). Palisade cells had less vacular space, that was frequently composed of numerous vacuoles.

Control leaves had well defined membrane structures (Fig. 3A and B). The double membrane structure of the nuclear envelope was clearly evident. Chloroplasts generally had well defined thylakoid membranes organized into granal stacks of varying heights. In leaves treated weekly with propiconazole, starch grains were more numerous within the chloroplasts (Fig. 3C and D). Although some chloroplasts exhibited structural characteristics similar to the control, there was a tendency for internal membranes to be less defined. Thylakoids showed less stacking and the stroma often stained more densely than controls. Leaves treated three times per week with propiconazole had a greater incidence of membrane perturbations (Fig. 3E and F). Thylakoid membranes were less organized and indistinct. Starch grains were observed frequently. In some cases, the chloroplast stroma was amorphous with few membrane profiles observed. Occasionally, nuclear membranes were more sinuous and irregularly spaced.

Discussion

Although the inhibitory effects of triazoles on leaf expansion have been documented, their possible anatomical manifestations have received little attention. Burrows et al. (1992) evaluated leaf structure in chrysanthemum leaves treated with a paclobutrazol soil drench. Leaves had an additional layer of palisade mesophyll, although the individual palisade cells were shorter, and more tightly packed. Benton and Cobb (1995) reported epoxiconazole (a triazole used as a fungicide) caused elongated palisade and spongy mesophyll, as well as upper epidermal cells in cleavers (Galium aparine L.). The triazoles, triadimefon and S-3307, applied as seed treatments increased epicuticular wax and reduced the length but increased the width and thickness of wheat leaves (Gao et al., 1988). In the current study, the palisade cells of the leaves from the propiconazole-treated shoots were similarly more densely packed with less intercellular spaces than the controls. The palisade mesophyll was composed of more cell layers, with cells more elongated and better defined.

Reports on the effects of triazoles on leaf thickness have been mixed. Burrows et al. (1992) demonstrated that paclobutrazol increased leaf thickness of ‘Lillian Hoek’ chrysanthemum. Benton and Cobb (1995) showed that epoxiconazole caused increase of leaflet thickness of cleavers. In contrast, Thetford et al. (1995) reported that uniconazole treatments of ‘Spectabilis’ forsythia
(Forsythia × intermedia Zab.) had no effect on leaf thickness. Likewise, propiconazole did not affect pecan leaf thickness in the current study. Wang and Gregg (1989) demonstrated uniconazole restricted cell division and differentiation in treated hibiscus. Differences in response may reflect the time of application and/or genetic differences in patterns of cell elongation and division in leaves. For example, spray applications made after completion of marginal meristem activity would be anticipated to have less impact on the number of cell layers comprising the palisade and spongy mesophyll than earlier sprays.

It is reported that leaves from triazole-treated plants contain higher chlorophyll concentrations or an alteration of chlorophyll synthesis (Benton and Cobb, 1995; Thetford et al., 1995; Wang and Gregg, 1989), thus resulting in a greening effect. In the current study, no apparent increase in the number of chloroplasts in individual mesophyll cells was evident with propiconazole treatments. However, the number of mesophyll cells per unit leaf area increased due mainly to a closer cell arrangement. Benton and Cobb (1995) also found that epoxiconazole-treated leaflets of cleavers had more palisade and spongy cells per unit area with fewer air spaces. They proposed that the tightly packed cells might be due to inhibition of cell separation caused by epoxiconazole. Thus, relatively few extracellular spaces and vacuolar areas may contribute to the observed greener leaves.

Triazoles may restrict plant growth by inhibition of phytosterol biosynthesis (Burden et al., 1987; Köller, 1987). Buchenauer and Röhrner (1981) suggested that the growth-retarding activity of triadimefon and triadimenol, two triazole fungicides, was the result of inhibition of both gibberellin and sterol biosynthesis. By simultaneous application of gibberellic acid (GA₃) with epoxiconazole on cleavers, Benton and Cobb (1995) indicated that the inhibitory effect of epoxiconazole on stem elongation could be counteracted by GA₃. Such inhibition of both gibberellin and sterol biosynthesis can result in accumulation of sterol precursors, possibly altering membrane integrity and functioning (Steffens, 1988). In the present study, chloroplast in leaves sprayed with propiconazole had thylakoid membranes with less stacking and internal membranes that were less defined. Modifications in membrane structure have likewise been reported with other triazole compounds. Paclobutrazol treatments in maize (Zea mays L.) seedlings increased the abundance of stroma lamellae and reduced the number of grana stacks in mesophyll chloroplasts (Sopher et al., 1999). Pring (1986) found that triadimefon applied to wheat seedlings caused fragmentation of the tonoplast and plasmalemma, and disintegration of chloroplasts. However, effects were localized and limited to cells adjacent to stomata which they attribute to accumulation and movement of the chemical in the transpi-
ration stream. In studies with corn and soybean [Glycine max (L.) Merrill] seedlings (Barnes et al., 1989), paclobutrazol and uniconazol had no visible effect on chloroplast ultrastructure, in contrast to amitro-t, which caused serious disruption of chloroplast membrane structure as well as a reduction of grana and stroma thylakoids.

Triazole fungicides are used widely in many agricultural and horticultural crops. While these fungicides have contributed greatly to control a variety of plant diseases, considerations should be made on their affects on plant development. In the present study, a triazole compound, propiconazole, applied to pecan leaves inhibited leaf expansion, altered mesophyll cell plast membrane structure as well as a reduction of grana and opaque stain in electron microscopy. J. Cell. Biol. 17:108–212.

Virtually all fungi produce sterols, which are precursors to ergosterol, the sterol of choice for the fungus and which is necessary for normal mycelial growth. A number of very potent fungicides, including benzimidazoles and triazoles, which are known as fungicidal growth regulators, interfere with sterol biosynthesis. Ellis Horwood, Chichester, United Kingdom.

**Literature Cited**


