Methyl Jasmonate-Induced Ethylene Production Is Responsible for Conifer Phloem Defense Responses and Reprogramming of Stem Cambial Zone for Traumatic Resin Duct Formation

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Conifer stem pest resistance includes constitutive defenses that discourage invasion and inducible defenses, including phenolic and terpenoid resin synthesis. Recently, methyl jasmonate (MJ) was shown to induce conifer resin and phenolic defenses; however, it is not known if MJ is the direct effector or if there is a downstream signal. Exogenous applications of MJ, methyl salicylate, and ethylene were used to assess inducible defense signaling mechanisms in conifer stems. MJ and ethylene but not methyl salicylate caused enhanced phenolic synthesis in polyphenolic parenchyma cells, early sclereid lignification, and reprogramming of the cambial zone to form traumatic resin ducts in *Pseudotsuga menziesii* and *Sequoiadendron giganteum*. Similar responses in internodes above and below treated internodes indicate transport of a signal giving a systemic response. Studies focusing on *P. menziesii* showed MJ induced ethylene production earlier and 77-fold higher than wounding. Ethylene production was also induced in internodes above the MJ-treated internode. Pretreatment of *P. menziesii* stems with the ethylene response inhibitor 1-methylcyclopropene inhibited MJ and wounding responses. Wounding increased 1-aminocyclopropane-1-carboxylic acid (ACC) oxidase protein, but MJ treatment produced a higher and more rapid ACC oxidase increase. ACC oxidase was most abundant in ray parenchyma cells, followed by cambial zone cells and resin duct epithelia. The data show these MJ-induced defense responses are mediated by ethylene. The cambial zone xylem mother cells are reprogrammed to differentiate into resin-secreting epithelial cells by an MJ-induced ethylene burst, whereas polyphenolic parenchyma cells are activated to increase polyphenol production. The results also indicate a central role of ray parenchyma in ethylene-induced defense.

Resistance in conifer stems to invasion by bark beetles, wood borers, and fungal pathogens includes constitutive defenses that deter initial invasion and inducible responses that may include mass oleoresin secretion and increased phenolic synthesis surrounding the wound zone following invasion. The bark (periderm and secondary phloem) is the first line of defense against stem-invading organisms and in many conifers contains basal levels of compartmentalized phenolic and terpenoid compounds. However, little is known about the activity of cells in the secondary phloem with respect to defense response mechanisms.

Polyphenolic parenchyma (PP) cells are a common component of the secondary phloem of all conifers and are active in the constitutive synthesis, storage, and modification of various phenolic compounds (Franceschi et al., 1998; Krekling et al., 2000). Following abiotic or biotic damage, PP cells give rise to the wound periderm and are activated to accumulate and release phenolics around the wound site, but these cells can also be induced to accumulate phenolics 10 to 30 cm away from damaged tissue (Franceschi et al., 1998, 2000; Hudgins et al., 2003a; Krekling et al., 2004).

A number of studies have revealed qualitative alterations in phenolic compounds following fungal inoculations, including increased activity of the flavonoid pathway (Brignolas et al., 1995; Lieutier et al., 1996; Bois and Lieutier, 1997; Bonello et al., 2003), and accumulation of phenolics is considered a significant part of induced defense responses in the bark (Nicholson and Hammerschmidt, 1992; Schultz et al., 1992; Viiri et al., 2001).

Many, but not all, conifers also contain resin-producing structures in the secondary phloem or xylem as a constitutive defense (Langenheim, 2003). Some Pinaceae species (e.g., *Pinus, Picea*) constitutively produce xylem resin ducts (Wu and Hu, 1997) but can be induced to form additional xylem traumatic resin ducts (TD) following herbivory, pathogen invasion, or wounding (Reid et al., 1967; Álvaro, 1995; Franceschi et al., 1998, 2000; Tomlin et al., 1998; Nagy et al., 2000; Hudgins et al., 2003a; McKay et al., 2003), while others (*Albes, Cedrus*) only develop xylem resin ducts in response to injury (Bannan, 1936; Fahn et al., 1979). The resin formed from inducible TDs may be more toxic or fungistatic because of the addition of phenolics and compositional changes of terpene content (Nagy et al., 2000; Fälldt et al., 2003; Krokene et al., 2003). Copious resin production provides an effective defense system that functions in flushing the wound, discouraging herbivory and sealing the injury through...
Pinaceae (Franceschi et al., 2002; Martin et al., 2002, 2003; Fälldt et al., 2003; Hudgins et al., 2003a) and Taxodiaceae (Hudgins et al., 2004). While there is little information on natural jasmonates in conifers, jasmonic acid has been shown to be induced in conifer cell culture (Taxus baccata) following elicitation with fungal cell wall fragments (Mueller et al., 1993). By contrast, jasmonates are known to modulate many activities in angiosperms, including defense responses (see Creelman and Mullet, 1997; Cheong and Choi, 2003; Farmer et al., 2003). A few investigations have also shown that MJ is capable of inducing terpenoid biosynthetic enzymes and their associated genes in Norway spruce (Picea abies; Martin et al., 2002, 2003; Fälldt et al., 2003) and chalcone synthase, a key enzyme in the phenylpropanoid pathway, in white spruce (Picea glauca; Richard et al., 2000). Thus, MJ appears to be involved in signal transduction pathways for both the phenylpropanoid and terpenoid defense responses in conifer stems. A question that remains is whether MJ directly induces TD development and PP cell activation or if it activates other factors that ultimately induce these responses.

A potential clue to the factors directly involved in TD formation may be inferred from the developmental features of these structures. TD formation occurs as a result of reprogramming of cambial cell derivatives that would normally develop into tracheids (Weker and Fahn, 1969; Nagy et al., 2000; Krekling et al., 2004), but mechanisms controlling this process are unknown. The phytohormone ethylene is involved in a diverse array of plant growth and developmental processes (Abeles et al., 1992; Kende, 1993; Bleecker and Kende, 1997) and is a possible candidate effector. In conifers, a number of studies have indicated that ethylene evolution is positively correlated with cambial zone activity, primarily with tracheid production and compression wood formation (Barker, 1979; Telewski and Jaffe, 1986; Eklund, 1990, 1991; Ingemarsson et al., 1991; Eklund and Little, 1995, 1996, 1998; Little and Pharis, 1995; Dolan, 1997; Little and Eklund, 1999; Klintborg et al., 2002; Andersson-Gunneras et al., 2003; Du and Yamamoto, 2003). Reports have also indicated that monoterpane synthesis and ethylene production are correlated with stem damage following fungal inoculations and wounding in several species of Pinus (Peters, 1977; Popp et al., 1995). Katoh and Croteau (1997) found that application of Ethrel, in addition to wounding, enhanced monoterpane synthase activity over wounding alone in Abies grandis (Pinaceae) saplings. Exogenous application of Ethrel has also been shown to induce enhanced resin exudation in a member of the Cupressaceae (Kusumoto and Suzuki, 2001), and an increase in the number (linear density) of resin ducts in seedlings of loblolly pine (Pinus taeda) and Pinus densiflora (Telewski et al., 1983; Yamamoto and Kozlowski, 1987). However, the direct effect of ethylene on resin production in conifers is not clear since many studies have included wounding as part of the application method, or continued application over many days.

A link between jasmonates and ethylene has been shown in a number of systems (Farmer et al., 2003). MJ has been shown to induce ethylene production in such diverse organs as apple and tomato fruits and olive leaves (Saniewski, 1997; Fan et al., 1998). In Arabidopsis, responses to different pathogens have been shown to include a synergistic effect of jasmonate and ethylene for induction of defense-related genes (Xu et al., 1994; Penningckx et al., 1998). For conifers, information on jasmonates has only recently become available but is limited, and a correlation between jasmonate and ethylene has not been established.

Another phytohormone that needs to be considered with respect to PP cell activation and TD formation in conifers is salicylic acid (SA), a known mediator of the expression of various defense-related genes (Ryals et al., 1996; Sticher et al., 1997). In conifers, SA has been shown to accumulate in response to pathogen challenge (Franich et al., 1986; Kozlowski and Metraux, 1998; Kozlowski et al., 1999), and Davis et al. (2002) showed that following either pathogen challenge or salicylic acid treatments, Pinus species were induced to express three chitinase homologs. Kozlowski et al. (1999) found that MJ induces the accumulation of salicylic acid in Norway spruce seedlings, establishing a link between the MJ and SA pathways in conifers. Although evidence indicates SA to be an important activator of genes encoding pathogenesis-related proteins (Durner et al., 1997), various proteins induced in the host plant in response to pathogens (van Loon et al., 1994), there is no information for its role in induction of other defense responses important to conifers, such as terpene and phenolic synthesis.

The purpose of this study was to determine if MJ directly induces PP cell activation and TD formation, as well as their systemic induction in untreated regions, or if it is an intermediate signaling compound. We used non-wounding exogenous applications of MJ, methyl salicylate (MS), and ethylene to assess stem responses in two diverse conifers: Douglas fir (Pseudotsuga menziesii Mirbel Franco), which has constitutive resin ducts, and giant redwood (Sequoia giganteum Lindl. Buchholz), which only produces TDs. Our results demonstrate that a major part of the MJ-induced response is due to increased ethylene production, and the sites of ethylene production are identified. The results are relevant to mechanisms of induced resistance in conifers, as previous studies have demonstrated that wounding or MJ pretreatment of mature Norway spruce stems induces these same
responses and also confers local resistance to the bark beetle-vectored fungal pathogen, *Ceratocystis polonica* (Franceschi et al., 2002; Krokene et al., 2003).

**RESULTS**

**Methyl Jasmonate and Ethylene Induce Resin Accumulation and Morphogenic Changes**

Following application of MJ or ethylene to Douglas fir and giant redwood, resin accumulated on the bark of the treated internode of all the saplings within 21 d but never appeared on the bark of MS or control saplings (data not shown; see Hudgins et al., 2004). Observations of stem surfaces indicated that resin exudation was considerably greater with 100 mM MJ than lower MJ concentrations and ethylene, which although not quantified correlates with the cross-sectional area of TDs formed at the treated internode at each concentration (see Figs. 3 and 4 for quantitative values).

Normal anatomy of Douglas fir secondary phloem, as seen in cross-section, consists of sieve cells and associated albuminous cells, PP cells, radial ray parenchyma, and scattered radial resin ducts. In control tissue (Tween 20 and water), PP cells generally occurred as semicomplete tangential rows with 5 to 10 sieve cells between rows, though individual PP cells were occasionally scattered among the sieve cells (Fig. 1A). All PP cells contained small phenolic bodies. Twenty-eight days after treatment with MJ or ethylene, PP cells appeared in an activated stage (Franceschi et al., 2000), which included swelling and massive accumulation of phenolics (Fig. 1, B and C; quantified in Fig. 3). PP cells showed about a 2-fold increase in size after 100 mM MJ and ethylene treatments versus Tween 20, MS, and water treatments (138 ± 17 μm² 0.1% Tween 20 versus 220 ± 37 μm² 100 mM MJ). MJ and ethylene also induced early lignification of stone cells (sclereids) in the secondary phloem (data not shown). PP cell and stone cell changes were not observed following Tween 20 (Fig. 1A), MS (Fig. 1D), or water treatments (Fig. 3).

Following MJ treatments to Douglas fir saplings, a tangential row of large axial xylem TDs developed proximal to the cambium, and radial resin ducts in secondary phloem and xylem were activated to accu-

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**Figure 1.** Douglas fir stem cross-sections 28 d after application of 0.1% Tween 20 (A), 100 mM MJ (B), ethylene (C), and 100 mM MS (D). Figures include similar region from each treatment with secondary phloem, cambium, and xylem for comparison. Bars = 200 μm. C, Cambial zone; PP, PP cell; PD, phloem resin duct; R, ray parenchyma; S, sieve cells; X, xylem. A, Normal anatomy of Douglas fir (Tween 20 control) with rows of PP cells, normal cambium, and the absence of constitutive axial xylem ducts. B, MJ-induced PP cell swelling, phenolic accumulation, and formation of radial phloem ducts and large axial xylem TDs. C, Ethylene treatment-induced PP cell swelling and phenolic deposition and a single row of TDs in the xylem. D, MS treatment did not induce anatomical changes.
mulate resin as indicated by expansion and increase in lumen size (Fig. 1B). The radial ducts of the secondary phloem were often seen to be connected with the induced xylem TDs (Fig. 1B). These same responses were found at all four internodes from which sections were collected, although the response was reduced in magnitude at both internodes above the treatment site (Fig. 4). Ethylene also induced xylem TDs, but the ducts were smaller than observed with MJ treatments (Figs. 1C and 3). Copper acetate staining indicated that induced axial and radial TDs contained resin acids (data not shown). Control treatments (Figs. 1A and 3) or MS (Figs. 1D and 3) did not induce axial or radial TDs.

Stems of giant redwood have a considerably different anatomy than Douglas fir and do not normally produce resin ducts, which is why we chose it to compare responses to ethylene. The secondary phloem of giant redwood consist of a highly organized structure with the following tangential pattern: fiber cells followed by sieve cells, followed by PP cells, then sieve cells, and then another layer fiber cell to repeat the pattern (Fig. 2A). Tangential layers of mature (thickened) fibers occur in the secondary phloem layers older than 2 years, while the fiber walls appear weakly lignified in the first two layers near the cambium. MJ treatment strongly induced PP cell activation as seen by both swelling of PP cells and the accumulation of phenolics (Figs. 2B and 3). Early lignification of young fiber cells also occurred in response to MJ treatment (Fig. 2B). Ethylene treatment also strongly induced PP cell activation and lignification of fiber cells (Figs. 2C and 3), while MS had no apparent effect on PP cells or fiber lignification (Fig. 2D). Differences were significant at $P < 0.05$.

Some species of Pinaceae but no species of Taxodiaceae that have been studied produce constitutive axial xylem resin ducts (Hudgins et al., 2004); however, MJ and ethylene both induced TD formation in giant redwood, a member of the Taxodiaceae (Fig. 2, B and C). The TDs that developed in giant redwood in response to 100 mM MJ were approximately 2.5-fold larger (in cross-sectional area) than those formed in response to exogenous ethylene treatments (Fig. 3C). Copper acetate stained the MJ- and ethylene-induced TD lumens blue, indicating the TDs produced resin acids and were functional following both treatments.

![Figure 2](image_url). Giant redwood stem cross-sections 28 d after application of 0.1% Tween 20 (A), 100 mM MJ (B), ethylene (C), and 100 mM MS (D). Figures include similar region from each treatment with secondary phloem, cambium, and xylem for comparison. Bars = 200 μm. C, Cambial zone; F, fiber; PP, PP cell; R, ray parenchyma; S, sieve cells; X, xylem. A, Tween 20 control has normal anatomy of giant redwood, with regular secondary phloem pattern of a row of PP cells followed by sieve cells, fibers, and another row of sieve cells, and the absence of constitutive axial xylem ducts. B, MJ-induced PP cell swelling, phenolic accumulation, lignification of young fibers, and formation of axial xylem TDs. C, Ethylene treatment induced PP cell swelling and phenolic deposition and a row of TDs in the xylem. D, MS treatment did not induce anatomical changes.
Dose Response to Methyl Jasmonate

A differential dose response was observed with respect to TD size and numbers, and the amount of MJ applied to giant redwood and Douglas fir (Fig. 4). In both species, TD size generally increased with increasing MJ concentration in the treated internode as well as internodes immediately above and below the treatment site. For a given MJ concentration, the size of the TDs decreased moving from the treated internode to the internode immediately above, and then again to the second internode above. TD size also decreased in the internode below the treated internode. Previous studies showed that gaseous MJ alone was not able to penetrate the bark at a level that gave a response (Hudgins et al., 2004), and, thus, the effects in distant internodes are due to internal transport of a signal.

A statistical analysis was done on the number (linear density in mm$^{-1}$) and size (area in $\mu m^2$) of MJ-induced TDs at the internode below the treated internode (internode A), the treated internode (B), and both internodes above the treatment site (C and D; 10 and 20 cm above, respectively). In giant redwood, significant differences were found only between the number of induced TDs at internodes A, B, C, and D within 10 and 50 mM MJ treatments ($P < 0.0001$; Fig. 4A). Significant differences with 25 mM MJ were found at internodes A and C ($P = 0.0037$) and internodes C and D ($P = 0.0037$; Fig. 4A). For 100 mM MJ, the number of TDs was significant between internodes B and A ($P = 0.0007$), C ($P = 0.0168$), and D ($P < 0.0001$). Within each MJ concentration in giant redwood, TD lumen sizes also showed significant differences ($P < 0.0001$; Fig. 4B). In Douglas fir, significant differences occurred between the number of induced TDs at all internode locations except A and C with 10 mM MJ treatment ($P = 0.3339$), while with 25 mM MJ differences were found between internodes A and C ($P = 0.0017$), B and C ($P < 0.0001$), B and D ($P = 0.006$), and C and D ($P = 0.0198$; Fig. 4C). The number of induced TDs were not significantly different within 50 and 100 mM MJ treatments at internodes A, C, and D ($P > 0.06$ in all cases); however, internode B was significantly different from A ($P = 0.011$) at 50 mM MJ. Lumen sizes of the induced TDs in Douglas fir at 10 mM MJ treatment were significant between internodes A and B, B and D, and C and D (all $P = 0.0372$; Fig. 4D). For 25 mM MJ, significant differences were found between A and B ($P = 0.0372$), A and D, B and D, and C and D at $P < 0.0001$. With 50 mM MJ, significant differences were found between internodes A and B ($P = 0.0002$), C and D ($P = 0.0025$), B and C ($P < 0.0001$), B and D ($P = 0.0001$), and B and D ($P < 0.0001$). For 100 mM MJ treatments, significant differences ($P < 0.0001$) were found between all internodes except A and C ($P = 0.81$).

Phenolic accumulation (data not shown) and TD formation (Fig. 4) were also found at the internodes above and below treatment sites for both MJ and ethylene treatments, although the magnitude of the remote response was reduced compared with the response at the application site. TD induction was not observed with MS (Fig. 2D) or control treatments (Fig. 2A).

Figure 3. Analysis of change in secondary phloem PP cell phenolics and xylem resin ducts 28 d after application of 0.1% Tween 20 and water (controls), and 100 mM MS, 100 mM MJ, and ethylene in giant redwood and Douglas fir. Shown are total mean polyphenolic body area (A), mean xylem resin duct cross sectional area (B), and mean number of resin ducts per millimeter of xylem (C). Measurements ($\mu m^2$ mm$^{-1}$) represent means from cross-sections from three individual replicate saplings. Means are of three replicates ± se. Forty PP cells and 10 xylem resin ducts were analyzed per section. Different letters indicate significant differences ($P < 0.05$) for each treatment within a species. Significant differences were found between MJ and ethylene, and MJ and ethylene and Tween 20, water, and MS treatments. Significant differences were not found among Tween 20, water, and MS treatments.
relationship between TD size and number is purely due to physical restraints.

Methyl Jasmonate and Wounding Induce Ethylene Production

Exogenous MJ treatment on Douglas fir stems induced ethylene production at both the treated internode and the untreated internode above (Fig. 5). The response was rapid, with a 77-fold increase of ethylene at 6 h over wound-induced ethylene at the same time point (significant at $P < 0.0001$). In the treated internode, there was a decrease in ethylene production from 6 to 24 to 48 h after treatment. The ethylene production by the untreated internode peaked at 24 h, indicating a lag phase probably associated with transport of the signal from the treated internode. Although significantly lower, ethylene above control levels was detected at wounded internodes after 6 h and increased at 24 h, and remained at this level up to 48 h (Fig. 5). An ethylene signal was observed at the internode above the wound; however, it remained below the quantifiable limits of the calibration curve. Statistical analysis revealed MJ treatments were significantly higher ($P < 0.0001$) over wounding at all sample time points, but differences between the treated and untreated internodes at 24 ($P = 0.131$) and 48 h ($P = 0.2211$) with MJ treatments were not significantly different. Ethylene was not detected in internodes of the Tween 20-treated control trees.

The Ethylene Antagonist, 1-Methylcyclopropene, Reduces the MJ and Wound Response

We sought to clarify and differentiate the role of MJ and ethylene by using 1-methylcyclopropene (1-MCP) to inhibit the ethylene response in Douglas fir. As found previously, in the absence of 1-MCP, MJ-treated stems developed activated PP cells and a tangential
band of xylem TD at the treated internode (Fig. 6A) and the internode directly above the treatment site (data not shown). In the presence of 1-MCP, only a few small scattered xylem TD were formed at the treated internode (Fig. 6B), and resin ducts were not formed at the internode above the treatment site. Quantitative analysis showed that 1-MCP reduced MJ-induced TD area to only 16% of that formed by MJ treatment without 1-MCP (908 ± 74 versus 5,850 ± 408 mm²/mm) in the treated internode, and no TDs were formed in the internode above the treated internode when 1-MCP was used. The TDs in Figure 6B are shown as an example of the anatomy of the few TDs formed in the presence of 1-MCP, and it must be noted that this picture represents a rarity, as in most of the stem no TDs could be found. However, PP cells did appear to be activated by MJ in the presence of 1-MCP (Fig. 6B), though it was not as extensive as in the absence of 1-MCP. Following wounding with a cork borer, a band of 10 to 15 phenolic filled cells and undifferentiated TDs formed in the xylem in a circumferential band some distance from the wound (Fig. 6, C and D). In saplings treated with 1-MCP and wounding, the same response only occurred directly below the wound and not in adjacent tissue (Fig. 6, E and F). Use of 1-MCP did not produce observable negative effects on growth of control saplings.

Expression and Localization of ACC Oxidase

To determine which cells may be involved in the MJ-induced ethylene production, we used immunohistochemical techniques and an antibody to 1-aminoacyclopropane-carboxylic acid (ACC) oxidase. Western immunoblots demonstrate that the antibody recognized ACC oxidase from Douglas fir, as indicated by a single band of the correct molecular mass of approximately 40 kD (Fig. 7). This enzyme is shown here to be constitutively expressed, as it is present in control tissue. The enzyme is also inducible; both wounding and MJ treatment results in an increase in the amount of protein (Fig. 7). A clear increase can be seen 96 h after wounding, whereas in MJ-treated plants, the increase can be seen at 6 and 48 h, with a subsequent decrease at 96 h after treatment (Fig. 7). These data are consistent with the data on ethylene production under similar treatments (Fig. 5).

Immunocytochemical localization demonstrated that ACC oxidase was most highly expressed within the cytoplasm of ray parenchyma cells of Douglas fir followed by cambial and PP cells (Fig. 8, A–E). Label was also abundant in secretory epithelial cells of resin ducts, including secondary phloem radial resin ducts (Fig. 8F), cortical axial resin ducts (Fig. 8, G and H), and TD (data not shown). Some label was associated with the nucleus (Fig. 8, A and D–G). The enzyme was present in control (Fig. 8E), wound (Fig. 8G), and MJ-treated (Fig. 8D) tissue, but the greatest amount of labeling occurred with wounding and MJ treatments. Labeling for ACC oxidase was associated with the cytoplasm and nucleus but not with the cell wall, vacuole, or other organelles. Nonimmune serum controls did not give any appreciable labeling of any cell type (Fig. 8C).

DISCUSSION

Constitutive phenolic and terpenoid compounds are important barriers to pests attacking the stems of conifers, but once compromised, additional defenses, some of which require extensive anatomical changes (Christiansen et al., 1999; Franceschi et al., 2000; Nagy et al., 2000), are elicited. The signals involved in eliciting these defenses are not understood. Here, it
is demonstrated that exogenous applications of ethylene can elicit similar defense responses as MJ treatment in stems of two diverse conifers, Douglas fir and giant redwood. The results indicate that ethylene is a downstream signaling agent from MJ and likely the ultimate elicitor of the extensive responses seen involving phenolics and especially terpenoids (Fig. 9). These MJ/ethylene-induced responses are relevant to induced resistance of conifer trees, as both wounding and MJ pretreatments can make mature Norway spruce completely resistant to the bark beetle-vectored fungal pathogen *C. polonica* (Franceschi et al., 2002;
Krokene et al., 2003). MS was ruled out as a signaling agent for activation of phenol and terpenoid synthesis and the alteration of cambial cell programming required for TD development. However, there are many other defense responses we have not yet explored, such as pathogenesis-related protein induction, where MS may play a role.

A major site for phenolic-based defenses in conifer stems is found in the PP cells of the secondary phloem (Franceschi et al., 1998, 2000; Krekling et al., 2000; Nagy et al., 2000; Kusumoto and Suzuki, 2003). Previous studies comparing MJ treatments and wounding found MJ is capable of activating PP cells similar to what was previously shown for Norway spruce in response to wounding or bark beetle attack (Franceschi et al., 2000), in all conifer lineages tested (Franceschi et al., 1998, 2000; Krekling et al., 2000; Nagy et al., 2000; Hudgins et al., 2004) and that appear to exist as a discrete symplasmic domain as indicated by synchronous TD formation (Krekling et al., 2004). Given our results that ethylene alone can induce TDs and the ethylene response inhibitor can inhibit TD induction by MJ and by wounding, it appears that ethylene is directly inducing reprogramming of cambial zone xylem mother cell differentiation from tracheid (Lachaud et al., 1999; Savidge, 2001) into secretory epithelial cells. While basal levels of ethylene in concert with other growth hormones (indole-3-acetic acid; Little and Pharis, 1995; Eklund and Little, 1998) control normal tracheid formation, a high-level ethylene burst induced by upstream signaling agents dramatically alters cell programming. Because the ethylene burst is dissipated in a day or so, as shown by our ethylene measurements, the subsequently produced derivatives from the cambial initial cells resume their regular developmental programming under a normal ethylene level and differentiate into tracheids, resulting in the TDs being embedded in xylem.

In a study of Norway spruce cuttings, significant increases in ethylene occurred 6 h after wounding (Ingemarsson and Bollmark, 1997), consistent with our results, and this included a turnover of ACC, the substrate for ACC oxidase (Kende, 1993; Bleecker and Kende, 2000). In conifers, ACC oxidase activity has also been shown to affect lignification, another process involving the phenylpropanoid pathway. Lignification appears to exist as a discrete symplasmic domain as indicated by synchronous TD formation (Krekling et al., 2004). Given our results that ethylene alone can induce TDs and the ethylene response inhibitor can inhibit TD induction by MJ and by wounding, it appears that ethylene is directly inducing reprogramming of cambial zone xylem mother cell differentiation from tracheid (Lachaud et al., 1999; Savidge, 2001) into secretory epithelial cells. While basal levels of ethylene in concert with other growth hormones (indole-3-acetic acid; Little and Pharis, 1995; Eklund and Little, 1998) control normal tracheid formation, a high-level ethylene burst induced by upstream signaling agents dramatically alters cell programming. Because the ethylene burst is dissipated in a day or so, as shown by our ethylene measurements, the subsequently produced derivatives from the cambial initial cells resume their regular developmental programming under a normal ethylene level and differentiate into tracheids, resulting in the TDs being embedded in xylem.

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been characterized in stems (Plomion et al., 2000) and needles (Ievinsh and Tillberg, 1995) of a couple of species. Here, it is shown that ACC oxidase is constitutively expressed in conifer stems but that wounding and, especially, exogenous MJ induce a large increase in levels of this enzyme. This increase is rapid and is correlated with the time course of MJ-induced enhancement of ethylene production in stems. Immunolabeling shows the enzyme is highest in the ray parenchyma and cambial zone cells. In addition, epithelial cells of radial resin ducts and axial cortical resin ducts have abundant labeling for the enzyme, which could help explain the rapid induction of resin flow by wounding, shown by Ruel et al. (1998), which occurs too quickly to be accounted for by TD formation. The ray parenchyma cells may be particularly important for ethylene signaling since they are involved in radial flow between secondary phloem and cambium/xylem (Fig. 9), and the cambium is considered to be supplied primarily by the rays (Levy-Yadun and Aloni, 1995). They are also linked anatomically to a tangential pathway in the secondary phloem.
of the stem via the rows of PP cells to which they are connected by abundant plasmodesmata in the interconnecting radial walls (Krekling et al., 2000).

There is conflicting information about the subcellular location of ACC oxidase. Various reports place it in the cell wall/apoplast (Rombaldi et al., 1994; Gomez-Jimenez et al., 2001), the external face of the plasma membrane (Ramassamy et al., 1998), and the cytosol (Chung et al., 2002). Our results show that the enzyme is present in the cytoplasm and the nucleus but not in other compartments. Nuclear localization in plants and animal cells has been shown for a number of different types of enzymes, including some in the glycolytic and gluconeogenic pathways (Yanez et al., 2003), Man-6-P reductase (Everard et al., 1993), plant hemoglobin (Seregelyes et al., 2000), apyrase (Shibata et al., 2002), lipoxygenase (Brock et al., 2001), and glutathione peroxidase (Orbea et al., 2000). Ethylene-responsive transcription factors are present in the plant nucleus (Johnson and Ecker, 1998; Ohta et al.,...
closely associated in slash pine (Pinus elliottii) production and the monoterpene concentrations were

increased production of ethylene is also often associated with plant-microbial interactions (Boller, 1991). Popp et al. (1995) demonstrated that following inoculations with pathogenic fungi, the rate of ethylene production and the monoterpene concentrations were closely associated in slash pine (Pinus elliottii) and loblolly pine. Additionally, the authors showed that inoculations with a virulent fungus increased ethylene production over nonpathogenic fungal inoculations, indicating that higher levels of ethylene production were likely a result of continuing fungal growth through stem tissues. Popp et al. (1997) also found that loblolly pine cell cultures exposed to virulent fungus showed increased levels of ethylene production, further demonstrating the relationship between fungal invasion and elicitation of ethylene production. It is possible that fungal toxins or enzymes may result in oxidative processes leading to the formation of jasmonates, which are known to elicit ethylene formation and ACC oxidase activity as shown here and by others (Kim et al., 1998; Watanabe and Sakai, 1998; Ievinsh et al., 2001), but this remains to be tested.

The similarity of responses to MJ and ethylene in the absence of wounding implies these compounds may both be directly involved in the inducible defense response pathway of conifers, similar to the genetically defined defense pathways of Arabidopsis (Pieterse et al., 1998; McDowell and Dangl, 2000). In angiosperms, several models have been proposed where jasmonates and ethylene produce a synergistic effect on certain defense related genes (Xu et al., 1994; O’Donnell et al., 1996; Penninckx et al., 1998) or resistance to multiple organisms (Bostock, 1998). The connection between MJ and ethylene may involve several potential pathways, but evidence provided here indicates that MJ acts upstream of ethylene. We propose a general model for the role of jasmonates and ethylene in the anatomical-based defense responses that we have been investigating (Fig. 9), and this will provide a basis for further investigations resolving the signaling pathway and transduction mechanisms and cellular response in the bark and cambial zone of conifers.

Our previous studies demonstrated that MJ or wounding can elicit a response in the untreated regions above or below the treated regions of mature trees or saplings (Franceschi et al., 2000, 2002; Hudgins et al., 2003a, 2004; Krekling et al., 2004), which is important to systemic resistance in conifers. Krekling et al. (2004) suggested a diffusible signal was responsible for a developmental wave away from a wound that moved at 2.5 cm per day with respect to TD formation. This study shows that ethylene can elicit the same systemic response as wounding and MJ.

Branches do not respond, indicating gaseous MJ or ethylene diffusing from the treated internodes is not at a high enough level to penetrate the periderm and is, thus, not responsible for the effect seen in untreated internodes. These observations support the concept that an internal signal is generated that can travel some distance from the region of initial elicitation, via the phloem (downward) and xylem (upward). Ethylene is more soluble than MJ or jasmonic acid in aqueous solutions, and we suggest it may be the primary signaling agent. However, given the constitutive levels of ACC oxidase, it is also possible that ACC (or a conjugate) may be transported as suggested by O’Neill et al. (1993), and a rapid flux of ACC to distant parts of the stem gives rise to a transient ethylene burst. These alternatives remain to be tested.

CONCLUSIONS

This study provides evidence for the activity of MJ and ethylene as key signaling compounds for inducible defense mechanisms and systemic defense involving the phenylpropanoid and terpenoid resin pathways in conifers. This up-regulation parallels at least part of the wound response that is seen in a number of conifers following abiotic or biotic injury and is relevant to acquired resistance against bark beetles and pathogenic fungi. Interestingly, exogenous MJ, ethylene, or MS applications did not induce lesion formation or characteristic tissue damage associated with a hypersensitive response, which implies that other signaling agents are required for the establishment of necrotic tissue and the necrophylactic periderm. When taken together, the data on exogenous ethylene induction of anatomical defense responses, MJ induction of ethylene synthesis and ACC oxidase protein, and 1-MCP inhibition of these responses provide a strong case for ethylene as the immediate signal for PP cell activation and cambial zone reprogramming for TD formation. These studies provide valuable new information on defense response mechanisms and experimental approaches for further work to resolve the signaling and signal transduction pathways of inducible defenses in conifer stems.

MATERIALS AND METHODS

Plant Materials and Greenhouse Conditions

Four-year-old Douglas fir (Pseudotsuga menziesii) and giant redwood (Sequoia sempervirens) saplings from a commercial nursery (Forest Farm, Williams, OR) were grown in a Washington State University-Pullman greenhouse. They were approximately 1 m in height, with a diameter of 1.5 to 2.0 cm at the first internode. Saplings were grown in 3-L containers at 23°C day and 18°C night temperatures, under a 14-h-light/10-h-dark regime for 3 months prior to treatments. They received 500 mL of water daily and fertilizer weekly [Peters fertilizer (20:20:20 N:P:K); Scotts Horticulture, Marysville, OH].

Methyl Jasmonate and Methyl Salicylate Treatments

On April 26, 2002, three replicate saplings of each species were treated with 10, 25, 50, or 100 mM MJ (Sigma-Aldrich, St. Louis; provided by Dr. J.
Gershenzon) in Millipore (Billerica, MA) Nanopure water with 0.1% (v/v) Tween 20 (Fischer Biotech, Fairlawn, NJ), or 10, 25, 50, or 100 mM MS (Sigma-Aldrich) in water with 0.1% (v/v) Tween 20. Three control trees from each species were treated with 0.1% (v/v) Tween 20 in water. The solutions were applied evenly around the second internode using a small brush and repeated after 5 min to ensure a uniform coating. A small rubber tube was tightened around the bottom of all treated internodes to ensure fluid did not reach the lower untreated internode. Tween 20 is a biologically nonactive detergent used to emulsify MJ and MS and to act as a surfactant to evenly spread the solution on the bark surface (Franceschi et al., 2002). On May 24, 2002, stem samples were collected for microscopy from the middle of the internode below the treated internode, the treated internode, and both internodes (10 and 20 cm, respectively) immediately above the treatment site. Samples were prepared for microscopy as described below. Representative images and statistical analysis are only presented for the application internode from 100 mM MJ, 100 mM MS, Ethaphon, and 0.1% Tween 20 treatments described above for Figures 1 to 3. Statistical analysis was conducted on all MJ concentrations for the dose-response study presented in Figure 4.

Ethylene Treatments

Three saplings of each species were treated with exogenous application of an Ethaphon (Carolina Biological, Burlington, NC) solution, and three received water (control). Ethaphon releases ethylene in the presence of water. At the first internode of all trees, a 5-cm length of stem was wrapped with thin cheesecloth and then wrapped twice with plastic (Saran Wrap: polyvinylidene chloride). Rubber bands were cut and tensioned over the plastic wrap at both ends to create a tight seal without injuring the stem. Ethylene treatments were applied by injecting 5 mL of Ethaphon:water solution (1:1, v/v) under the plastic wrap with a small gauge hypodermic needle. Following application, the treated area was wrapped with plastic. Second time with plastic. Saplings were treated for 12 h, and the apparatus was then removed. Control (water) treatments used the same method. Stem samples were taken 24 h after treatment and were prepared for microscopy as described below.

Ethylene Inhibitor Treatment on Douglas Fir

To examine the potential role of ethylene as a downstream signaling agent from MJ, nine saplings of Douglas fir (three replicates of each of three treatments) were exposed to the ethylene response inhibitor 1-MCP (EthylBloc; Rohm & Haas, Philadelphia) prior to subsequent treatments. Three groups of three saplings were treated separately and sequentially in a sealed 225-L Plexiglas growth chamber. Each group of three saplings was first exposed to 1-MCP at 9 ppm for 12 h, and then the three saplings from each group were treated with Tween 20 (control), 100 mM MJ, or a 5-mm cork borer wound into the sapwood. Along with the treatment, a second dose of 1-MCP of the same concentration was released in the container, and after an additional 12 h of exposure the plants were transferred to a greenhouse. Control, MJ, and wound treatments were conducted independently to exclude additional 12 h of exposure the plants were transferred to a greenhouse. Controls were raised in goat against a peptide mapping at the amino terminus; Santa Cruz Biotechnology, Santa Cruz, CA) for 3 h. The membrane was rinsed with TBST + BSA (4 × 15 min), and then treated with rabbit anti-goat alkaline phosphatase-tagged secondary antibody (Sigma-Aldrich) diluted 1:2,500 in TBST + BSA for 1 h. After washing, the membrane was allowed to equilibrate in reaction buffer (100 mM Tris, 10 mM NaCl, 10 mM MgCl2, pH 9.5), and protein immunodetection was visualized by a reaction with p-Nitro-Blue tetrazolium chloride (NBT) and X-Phosphate in the reaction buffer (45 μL of NBT and 35 μL of X-Phosphate/10 mL of buffer).

Light Microscopy Staining

Copper Acetate Resin Staining

Samples from experiments were routinely stained to determine resin acid production and location. Tissue was fixed as described below, rinsed (four times) in 50 mM buffer (L-piperazine-N,N’-bis-(ethane sulfonic) acid, pH 7.2) and hand cut into semithin sections (0.1-0.5 mm). Sections were then soaked 12 h in a saturated (aqueous) copper acetate solution, which stains resin acids blue. Sections were rinsed with distilled water and examined by light microscopy to determine if resin acid accumulation was present.

Sampling and Microscopic Preparation of Stem Samples

Intact stem section samples (3 cm) were collected and directly placed in fixative solution [2% (v/v) paraformaldehyde and 1.25% (v/v) glutaraldehyde buffered in 50 mM 1-piperazine-N,N’-bis-(ethane sulfonic) acid, pH 7.2]. Each sample was subsequently cut into 2.5-mm² blocks and allowed to fix for 24 h at 4°C. Blocks were subsampled and rinsed with the same buffer solution, dehydrated with an ethanol series, and infiltrated with L.R. White acrylic resin (London Resin, Reading, UK). Samples were polymerized in fresh resin at 62°C for 12 h.

General Anatomy Staining

Sections (1 μm thick) were cut with a diamond knife on a Sorvall MT 5000 Ultramicrotome (Asheville, NC) and dried from a drop of water onto gelatin-coated slides. Stevener’s blue staining (Del Cerro et al., 1980) was used for general observation of tissue structure.

Image Analysis

Scion Image 1.62 imaging software (Scion, Frederick, MD; www.scioncorp.com) was used to make measurements of PP cells and TD from stem cross-sections of each of three replicate saplings for each treatment. The cross-sectional areas of PP cells and their polyphenolic inclusions were measured, whereas both the number of TDs and their cross-sectional areas were determined. Forty to 50 PP cells per section were measured from three different sections and combined to represent one treated internode per sapling. Eight to 20

Protein Extraction and Western-Immunoblot Analysis

And Immunocytochemical Analysis of Douglas Fir

Twenty-seven Douglas fir stems were treated as described above for ethylene analysis. Three saplings were collected following each treatment at 6, 48, and 96 h. Stem secondary phloem and cambial zone tissue was separated from the xylem, frozen in liquid nitrogen, and stored at -80°C. Protein extractions were conducted as described previously by Martin et al. (2002), with the addition of the protease inhibitors phenylmethylsulfonyl fluoride (0.5 mM) and Complete Tables (Roche, Nutley, NJ). Total protein concentrations were determined using a Protein Assay kit (Pierce Biotechnology, Rockford, IL) with bovine serum albumin (BSA) as a standard. Equal masses (18 μg) of protein were loaded in each lane, and the samples were subjected to SDS-PAGE using 10% polyacrylamide gels. Gels were either stained with Coomassie Brilliant Blue R-250 for imaging, or proteins were transferred to a PVDF membrane (Bio-Rad, Hercules, CA). The membrane was incubated for 1 h with TBST + BSA [Tris-buffered saline-Tween 20; 10 mM Tris, 250 mM NaCl, 0.1% (v/v) Tween 20 + 1% (v/v) BSA, pH 7.2] and then treated with 1.5% dilution of an affinity-purified polyclonal anti-Arabidopsis ACC oxidase (anN-19, raised in goat against a peptide mapping at the amino terminus; Santa Cruz Biotechnology, Santa Cruz, CA) for 3 h. The membrane was rinsed with TBST + BSA (4 × 15 min), and then treated with rabbit anti-goat alkaline phosphatase-tagged secondary antibody (Sigma-Aldrich) diluted 1:2,500 in TBST + BSA for 1 h. After washing, the membrane was allowed to equilibrate in reaction buffer (100 mM Tris, 10 mM NaCl, 10 mM MgCl2, pH 9.5), and protein immunodetection was visualized by a reaction with p-Nitro-Blue tetrazolium chloride (NBT) and X-Phosphate in the reaction buffer (45 μL of NBT and 35 μL of X-Phosphate/10 mL of buffer).
10 TDs per section were measured from three different sections and combined to represent a treated internode per sapling. For statistical analysis, data was calculated based on combined averages from three individual saplings (n = 3).

Immunocytochemistry

Stem section tissue was collected from MJ-, wound-, and control (water)-treated Douglas fir following treatments at 6 and 96 h for immunocytochemical comparison with western-blot analysis. Tissue was fixed and embedded in L.R. White as described above and was sectioned at 1 μm. Tissue was first blocked for nonspecific binding of antibodies by incubation for 1 h in TBST + BSA containing 0.05% (w/v) polyvinylpirrolidone (10,000 molecular weight). Sections were incubated for 4 h at room temperature with goat anti-Arabidopsis ACC oxidase polyclonal antibody diluted 1:75 in TBST + BSA. Following incubation with the primary antibody, the sections were rinsed with TBST + BSA (4 × 15 min) and incubated for 1 h at room temperature with rabbit anti-goat gold-tagged secondary antibody diluted 1:150 with TBST + BSA. After sequential rinsing with TBST + BSA, TBST, and distilled water, the sections were silver enhanced (Amersham Silver Enhancement kit; Amersham Life Science, Buckinghamshire, UK) and counter-stained with 0.5% (w/v) aqueous Safranin O. The sections were coverslipped with immersion oil as a mounting media and imaged with a confocal microscope (Bio-Rad 1024), using a combination of reflected and transmitted imaging.

Statistical Analysis

All statistical analyses were conducted with SAS software (SAS Institute, Cary, NC) with significance judged at P < 0.05 in all cases. Dose-response studies were treated as two-way analysis of variance (ANOVA) in a completely randomized design with protected LSD to compare number of resin ducts and transmitted imaging. For effect of various compounds between ethylene and MJ treatment groups. For effect of various compounds terthwaite adjustments were made to the tests 3).

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