Novel Roles for the Polyphenol Oxidase Enzyme in Secondary Metabolism and the Regulation of Cell Death in Walnut


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The enzyme polyphenol oxidase (PPO) catalyzes the oxidation of phenolic compounds into highly reactive quinones. Polymerization of PPO-derived quinones causes the postharvest browning of cut or bruised fruit, but the native physiological functions of PPOs in undamaged, intact plant cells are not well understood. Walnut (juglans regia) produces a rich array of phenolic compounds and possesses a single PPO enzyme, rendering it an ideal model to study PPO. We generated a series of PPO-silenced transgenic walnut lines that display less than 5% of wild-type PPO activity. Strikingly, the PPO-silenced plants developed spontaneous necrotic lesions on their leaves in the absence of pathogen challenge (i.e. a lesion mimic phenotype). To gain a clearer perspective on the potential functions of PPO and its possible connection to cell death, we compared the leaf transcriptomes and metabolomes of wild-type and PPO-silenced plants. Silencing of PPO caused major alterations in the metabolism of phenolic compounds and their derivatives (e.g. coumaric acid and catechin) and in the expression of phenylpropanoid pathway genes. Several observed metabolic changes point to a direct role for PPO in the metabolism of tyrosine and in the biosynthesis of the hydroxycoumarin esculetin in vivo. In addition, PPO-silenced plants displayed massive (9-fold) increases in the tyrosine-derived metabolite tyramine, whose exogenous application elicits cell death in walnut and several other plant species. Overall, these results suggest that PPO plays a novel and fundamental role in secondary metabolism and acts as an indirect regulator of cell death in walnut.

Polyphenol oxidases (PPOs) are a group of copper-containing enzymes that catalyze the o-hydroxylation of monophenols to o-diphenols (tyrosinase activity, Enzyme Commission 1.14.18.1) as well as the oxidation of o-diphenols to quinones (catecholase activity, Enzyme Commission 1.10.3.2) in the presence of oxygen. These enzymes are broadly distributed among animals, fungi, and plants, though many plant PPOs appear to lack tyrosinase activity (Steffens et al., 1994). The study of PPOs in plants has focused primarily on their role in the process of postharvest browning, whereby cut or damaged plant tissues turn brown due to the polymerization of PPO-generated quinones, generating phytomelanins (Mesquita and Queiroz, 2013). Classically, PPOs and their potential phenolic substrates have been considered to be physically separated from one another in intact plant cells, with most PPOs targeted to the chloroplasts, while phenolic compounds accumulate primarily in the vacuole and cell wall (Vaughn et al., 1988; Steffens et al., 1994). Thus, PPO activity in vivo has typically been associated with senescing, wounded, or damaged plant tissues in which cellular compartmentalization is lost (Steffens et al., 1994).

PPOs are most commonly encoded by multigene families in plants, ranging in size from 13 PPO genes in Physcomitrella patens to two PPO genes in rice (Oryza sativa). However, several plant species, including cucumber (Cucumis sativus), cassava (Manihot esculenta), and walnut (juglans regia), have only a single PPO gene, and Arabidopsis (Arabidopsis thaliana) lacks PPO altogether (Escobar et al., 2008; Tran et al., 2012). Within a single plant species, different members of the PPO gene family can display quite divergent regulatory patterns, with expression in different tissues or in the same tissue but at different developmental stages (Thygesen et al., 1995; Thipyapong and Steffens, 1997; Kim et al., 2001). Many PPO genes have been shown to be up-regulated by wounding, pathogens, and/or stress-associated hormones such as methyl jasmonate and salicylic acid (SA), but responsiveness to these cues again varies between species and within PPO gene families (Constabel and Ryan, 1998). For example, in the five-member PPO gene family in dandelion (Taraxacum officinale), only ppo2 is responsive to pathogen infection (Richter et al., 2012).

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Although PPO was once described as “the chloroplast oxidase with no established function” (Vaughn et al., 1988), more recent studies have shed some light on possible functions of PPOs in plants. Best characterized is a role of some PPO genes in plant defense against insects and pathogens. As previously discussed, many PPO genes are up-regulated upon pathogen challenge, and overexpression of a potato (Solanum tuberosum) PPO in tomato (Solanum lycopersicum) resulted in decreased susceptibility to Pseudomonas syringae pv tomato (Li and Steffens, 2002). Likewise, silencing of PPOs in tomato led to increased disease susceptibility (Thiypayong et al., 2004). Similarly, overexpression of PPO genes in tomato and hybrid aspen (Populus tremula × Populus alba) resulted in increased insect resistance, and silencing of PPOs resulted in increased susceptibility to insect herbivory (Wang and Constabel, 2004; Mahanal et al., 2008; Bhonwong et al., 2009). PPO also plays a central role in the polymerization of glandular trichome exudates in wild potato (Solanum berthaultii), a process that entraps small insects such as aphids (Kowalski et al., 1992).

PPOs have also been implicated in the biosynthesis of some specialized pigments and other secondary metabolites. Specific PPOs are required for the synthesis of aurone pigments in snapdragons (Antirrhinum majus) and betalin pigments in the order Caryophyllales (Nakayama et al., 2001; Gandia-Herrero et al., 2005). PPO also has a central role in the biosynthesis of 8-8’ linked lignans in creosote bush (Larrea tridentata; Cho et al., 2003). Given the diverse regulation and characterized functions of PPO genes, it appears likely that different PPOs within a gene family may play a variety of roles in the physiology and development of a single plant.

Walnut presents an interesting model to further explore the function(s) of PPOs in plants, in that walnuts generate an exceptionally diverse array of phenolic compounds (Colaric et al., 2005; Solar et al., 2006) and possess a single PPO gene, JrPPO1 (Escobar et al., 2008). JrPPO1 appears to be constitutively expressed at a high level in all green tissues and is not responsive to wounding or methyl jasmonate treatment (Escobar et al., 2008). To better understand the functional role of the single PPO enzyme in walnut, we used RNA interference (RNAi) to generate a series of transgenic plants, extracted total protein, and performed PPO enzyme activity assays, using L-3,4-dihydroxy-Phe (l-DOPA) as substrate (Fig. 1). All of the RNAi lines showed more than 95% reduction in leaf PPO activity, demonstrating highly efficient silencing of JrPPO1. Corresponding reductions in JrPPO1 mRNA levels were also demonstrated via real-time reverse transcription-PCR (RT-PCR) analysis of selected RNAi lines (Supplemental Fig. S1). Surprisingly, the “overexpression” lines (40-1-1, 78-2-8) also showed large reductions in leaf PPO activity, indicating the activation of cosuppression rather than successful overexpression of JrPPO1.

The existence of transgenic walnut lines showing near-complete suppression of PPO activity allowed us to examine which of the diverse group of phenolic compounds generated by walnuts could potentially serve as substrates for JrPPO1. Caffeic acid, chlorogenic acid, and catechin are o-diphenols that are naturally synthesized in walnut leaves and/or hulls (Solar et al., 2006). These phenolics were utilized as substrates in polarimetric PPO activity assays (catecholase activity), using total protein extracts from either wild-type leaves or leaves from a randomly selected PPO-silenced line (9-5-1). The PPO-silenced line served as a negative control, assuring that any observed activity (oxygen consumption) could be attributed specifically to PPO. As summarized in Figure 2A, each of these native substrates could be somatic embryos were transformed with two binary vectors: one designed to constitutively overexpress the jrPPO1 gene and one designed to silence the jrPPO1 gene. We recovered and germinated two lines transformed with the overexpression vector and nine lines transformed with the RNAi vector. Nontransformed walnut somatic embryos were germinated in parallel to provide matched wild-type controls. Following transfer to soil, we collected leaves from these transgenic plants, extracted total protein, and performed PPO enzyme activity assays, using L-3,4-dihydroxy-Phe (l-DOPA) as substrate (Fig. 1).

**RESULTS**

**Generation and Characterization of PPO-Silenced Walnut Lines**

To manipulate the levels of active PPO enzyme in walnut and study the resulting phenotypic effects, walnut

![Figure 1](image-url)
effectively oxidized by PPO in vitro, with no activity detected in protein extracts from the silenced line. Substrate kinetics for each of these o-diphenols is presented in Table I. We also examined the native monophenols Tyr, tyramine, and p-coumaric acid as potential substrates for JrPPO1. In general, PPOs display much lower tyrosinase activities than catecholase activities (i.e. the hydroxylation of monophenols is much slower than the oxidation of o-diphenols; Selinheimo et al., 2007; Goldfeder et al., 2013), and this was also true for JrPPO1 (Supplemental Fig. S2). To simplify measurement of the relatively low in vitro tyrosinase activities, PPO assays utilizing monophenol substrates were performed spectrophotometrically. As summarized in Figure 2B, all three monophenols could be utilized as substrates by JrPPO1, though activity was substantially lower for p-coumaric acid than for Tyr and tyramine.

Phenotyping PPO-Silenced Walnut Lines

Given previous research suggesting a role for some PPOs in plant defense (Li and Steffens, 2002), we initially planned to perform a series of pathogen challenge bioassays on the PPO-silenced walnut lines and matched wild-type controls. These plans were almost immediately abandoned, however, because shortly after transfer to soil, the PPO-silenced lines (but none of the wild-type controls) spontaneously developed scattered necrotic lesions on their leaves (Fig. 3). The development of these lesions has been observed in the PPO-silenced lines in four consecutive growing seasons, with trees grown at two separate locations (Davis, CA; San Marcos, CA). Once formed, the lesions are typically static, though they can coalesce late in the season, causing premature leaf drop (Fig. 3). We initially believed that the leaf necrosis was caused by naturally occurring pathogen infection, given that PPO-silenced tomatoes have previously been shown to be hypersusceptible to bacterial speck disease (Thiyavong et al., 2004). Two walnut diseases can generate scattered leaf necrosis symptoms: walnut blight (causal agent, Xanthomonas arboricola pv juglandis) and walnut anthracnose (causal agent, Gnomonia leptostyla; Cline and Neely, 1983; Belisario et al., 1999). Attempts to isolate X. arboricola pv juglandis from the necrotic lesions on PPO-silenced lines were unsuccessful, though the pathogen could be readily isolated from artificially inoculated wild-type plants (data not shown; Belisario et al., 1999). The lesions also lacked the small fruiting bodies (acervuli) that are diagnostic features of walnut anthracnose, and attempts to isolate G. leptostyla mycelia from leaf discs containing necrotic tissue were also unsuccessful (Belisario et al., 2008). In addition, the lesions were completely static when detached leaves were incubated in a moist chamber, with no formation of fungal fruiting bodies, bacterial ooze, or increase in lesion size. Thus, the necrotic spots that form on the leaves of PPO-silenced walnut plants appear to be independent of pathogen challenge and were thus classified as a lesion mimic phenotype.

Transcript and Metabolite Profiling of PPO-Silenced Walnut Lines

To gain a better understanding of how the silencing of jrPPO1 leads to cell death, we performed global transcript profiling and global metabolite profiling of leaves collected from matched wild-type and PPO-silenced
edgeR (Robinson et al., 2010) demonstrated clustering of twodimensional scaling of log-transformed count data using detected for 53,994 of these contigs in our data set. Multidimensional scaling of log-transformed count data using edgeR (Robinson et al., 2010) demonstrated clustering of the three wild-type replicates and clustering of the three PPO-silenced replicates (Supplemental Fig. S6). As expected, sequence reads aligning to the neomycin phosphotransferase II gene (the plant-selectable marker gene on the RNAi construct) were abundant in the PPO-silenced samples (3,363 total reads) and essentially absent in the wild-type samples (two total reads; presumably misalignments). In total, 93 contigs in the walnut transcriptome database showed statistically significant differential expression between the PPO-silenced lines and the wild-type controls (Supplemental Table S1). BLASTX analysis against The Arabidopsis Information Resource 10 Proteins database was performed to identify putative Arabidopsis homologs of the differentially expressed walnut genes. The Arabidopsis Genome Initiative numbers of these putative homologs (Supplemental Table S1) were then analyzed using BioMaps software, which identifies and statistically analyzes overrepresented functional categories in global data sets (Katari et al., 2010). This analysis revealed overrepresentation of cell wall-associated genes (P = 0.001), all of which were expressed at higher levels in the PPO-silenced lines (Supplemental Table S2). Visual inspection of the data also revealed that many of the overexpressed genes in PPO-silenced lines had predicted functions in the metabolism of phenolic compounds, especially the phenylpropanoid pathway, including putative cinnamic acid 4-hydroxylase, 4-coumarate-CoA ligase, hydroxycinnamoyl-CoA shikimate/quinate hydroxycinnamoyl transferase, and 4-coumaroyl ester 3′-hydroxylase genes (Fig. 4).

Metabolite profiling revealed a total of 83 small molecules showing statistically significant differences in abundance between wild-type and PPO-silenced leaf samples (Supplemental Table S3). Strikingly, out of the 10 metabolites that showed the most significant differences in abundance between wild-type and PPO-silenced lines (i.e. lowest associated P values), seven are phenolic compounds or derivatives of phenolic compounds. Large increases in many metabolites derived from the phenylpropanoid pathway were observed in the PPO-silenced lines, most notably flavonoids such as catechin, gallocatechin, and eriodictyol (Fig. 4). The phenylpropanoid pathway also produces the precursors to lignin, including sinapic acid and ferulic acid, which were elevated 3.6- and 1.6-fold, respectively, in the PPO-silenced lines. However, subsequent analyses of H lignin, S lignin, G lignin, and total lignin content in wild-type and PPO-silenced walnut leaves revealed no significant alterations in lignin composition (Supplemental Fig. S7). Regardless, both RNA-seq data and metabolite data suggest an increased activity of the phenylpropanoid pathway in PPO-silenced walnut plants.

In addition to Phe (the starting point of the phenylpropanoid pathway), metabolism involving the other aromatic amino acids was also altered in the PPO-silenced walnut lines. For example, the Trp derivatives tryptamine and serotonin displayed large increases in abundance in the PPO-silenced lines (6.1- and 8.6-fold, respectively).

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Because the walnut genome has not yet been sequenced, reads from RNA-seq analysis were aligned to a previously assembled and annotated walnut transcriptome database containing 85,045 contigs (You et al., 2012). Substantial expression (≥ 20 aligning reads) was detected for 53,994 of these contigs in our data set. Multidimensional scaling of log-transformed count data using edgeR (Robinson et al., 2010) demonstrated clustering of the three wild-type replicates and clustering of the three PPO-silenced replicates (Supplemental Fig. S6). As expected, sequence reads aligning to the neomycin phosphotransferase II gene (the plant-selectable marker gene on the RNAi construct) were abundant in the PPO-silenced samples (3,363 total reads) and essentially absent in the wild-type samples (two total reads; presumably misalignments). In total, 93 contigs in the walnut transcriptome database showed statistically significant differential expression between the PPO-silenced lines and the wild-type controls (Supplemental Table S1). BLASTX analysis against The Arabidopsis Information Resource 10 Proteins database was performed to identify putative Arabidopsis homologs of the differentially expressed walnut genes. The Arabidopsis Genome Initiative numbers of these putative homologs (Supplemental Table S1) were then analyzed using BioMaps software, which identifies and statistically analyzes overrepresented functional categories in global data sets (Katari et al., 2010). This analysis revealed overrepresentation of cell wall-associated genes (P = 0.001), all of which were expressed at higher levels in the PPO-silenced lines (Supplemental Table S2). Visual inspection of the data also revealed that many of the overexpressed genes in PPO-silenced lines had predicted functions in the metabolism of phenolic compounds, especially the phenylpropanoid pathway, including putative cinnamic acid 4-hydroxylase, 4-coumarate-CoA ligase, hydroxycinnamoyl-CoA shikimate/quinate hydroxycinnamoyl transferase, and 4-coumaroyl ester 3′-hydroxylase genes (Fig. 4).

Figure 3. Silencing of jpppo1 induces a lesion mimic phenotype. A, Wild-type walnut leaf. B, PPO-silenced line, early season (June). Inset, magnified view of necrotic lesions from abaxial side of leaf. C, PPO-silenced line, late season (September). The development of the necrotic lesions was not associated with the presence of any detectable pathogen. Bars = 4 cm.
levels of the L-DOPA derivative 5,6-dihydroxyindole were decreased more than 95% in the PPO-silenced lines. Dopamine can be synthesized via either by the L-DOPA pathway or by the tyramine pathway through successive o-hydroxylation and decarboxylation of Tyr (the order of reactions depends on the pathway; Fig. 5). Dopamine levels were also dramatically reduced in PPO-silenced lines (>80% reduction). The simplest interpretation of these findings is that the o-hydroxylation of Tyr and tyramine is blocked in the PPO-silenced lines, leading to the accumulation of tyramine and the shunting of Tyr metabolism into alternative metabolic pathways, such as 4-hydroxyphenylpyruvate pathway (Fig. 5).

The accumulation of aromatic amines such as tyramine, tryptamine, and serotonin is often associated with the synthesis of hydroxycinnamic acid amides (HCAAs), which are deposited into the cell wall, increasing its strength and reducing its digestibility (Jang et al., 2004; Ishihara et al., 2008). HCAAs are produced through the condensation of aromatic amines and hydroxycinnamoyl-CoA thioesters, often in response to wounding or pathogen challenge (Guillet and De Luca, 2005; Muroi et al., 2009). Because both aromatic amines and phenylpropanoid-derived hydroxycinnamoyl-CoA thioesters are presumably elevated in PPO-silenced lines, we examined HCAA accumulation in situ by staining walnut leaf sections with Neu’s reagent and visualizing HCAA fluorescence (Alemanno et al., 2003). Both wild-type and PPO-silenced lines showed clear staining in leaf vasculature, but the PPO-silenced lines also displayed strong HCAA accumulation in the living cells surrounding necrotic lesions (Fig. 6). Thus, it appears that the lesion mimic phenotype in PPO-silenced walnut leaves is associated with the localized accumulation of HCAAs.

The Relationship between PPO and Cell Death

Other than the recent demonstration that the tomato PPO B gene is strongly expressed in some tissues undergoing apoptosis (Newman et al., 2011), little previous evidence has linked PPO to programmed cell death. By contrast, several studies have previously implicated aromatic amines in the regulation of cell death and/or lesion mimic phenotypes (in addition to their role in HCAA production). Namely, elevating plant tyramine or serotonin levels by exogenous application or through transgenic approaches can lead to cell death, growth inhibition, and/or lesion mimic phenotypes (Negrel et al., 1993; Ueno et al., 2003; Kang et al., 2007; Fujiwara et al., 2010; Kim et al., 2011). Given that PPO-silenced lines display a lesion mimic phenotype and have highly elevated tyramine and serotonin levels (approximately 9-fold increases in each relative to the wild type), we were curious whether exogenous treatment with tyramine or serotonin could induce de novo lesion formation in wild-type walnut leaves. Thus, the petioles of detached wild-type walnut leaves were incubated in solutions of 5 mM tyramine or 5 mM serotonin, mimicking the transpirational uptake system and aromatic amine concentrations respectively. The most striking alterations, however, were observed in the metabolism of Tyr. As summarized in Figure 5, three major pathways of Tyr metabolism are the tyramine pathway, the L-DOPA pathway, and the 4-hydroxyphenylpyruvate pathway (Zhang et al., 2005; Maeda and Dudareva, 2012). PPO-silenced plants displayed a massive increase in tyramine levels (9.3-fold), suggesting increased activity of this pathway. 4-Hydroxyphenylpyruvate was not detected in our metabolite extracts, but its derivatives α-tocopherol, β-tocopherol, δ-tocopherol, and γ-tocopherol were all significantly elevated in PPO-silenced plants, suggesting increased activity of the 4-hydroxyphenylpyruvate pathway (Riewe et al., 2012). By contrast, the L-DOPA pathway, whose initial step involves the o-hydroxylation of Tyr by an as-yet uncharacterized enzyme (Lee and Facchini, 2011), appears to be essentially inactive in the PPO-silenced lines. Although L-DOPA itself was not detected, the
previously utilized by Ueno et al. (2003) in their investigations of the Sekiguchi lesion mutant of rice. Relative to detached leaves incubated in water, serotonin treatment typically caused a darkening of the walnut leaf midrib after 2 to 4 d (data not shown), potentially due to oxidation of serotonin by native peroxidases (Ishihara et al., 2008). By contrast, incubation with tyramine typically caused the spontaneous development of expanding necrotic lesions in the leaf lamina after 2 to 4 d, often in close association with leaf veins (Fig. 7). Thus, exposure of wild-type leaves to elevated levels of tyramine results in a near phenocopy of the lesion mimic phenotype observed in PPO-silenced lines.

DISCUSSION AND CONCLUSION

Although Vaughn et al. (1988) suggested that PPOs are not involved in the metabolism of phenolic compounds in healthy, intact cells, several more recent studies have demonstrated that some PPOs do serve as biosynthetic enzymes required for the production of specialized phenolic-derived pigments (aurones, betalins) and other secondary metabolites (Joy et al., 1995; Cho et al., 2003; Gandía-Herrero et al., 2005). Using a reverse genetics approach, we attempted to overexpress and silence the single PPO gene in walnut, *jrPPO1*, to determine its physiological function. Interestingly, we were unable to recover any *jrPPO1* overexpressing lines, while silencing of *jrPPO1* was very efficient (Fig. 1). Although PPOs have been overexpressed in a variety of other plant species, it is possible that successful overexpression of PPO in walnut leads to a lethal phenotype. Alternatively, given the very high basal level of JrPPO1 activity in walnut (Escobar et al., 2008), it is possible that attempts to express this gene at an even higher level simply result in cosuppression, as was observed in lines 40-I-1 and 78-2-8. In any case, further analysis was restricted to *jrPPO1*-silenced lines, all of which demonstrated more than 95% reduction in leaf PPO activity.

Several studies have shown that PPOs are quite recalcitrant to high-level recombinant protein production in nonplant systems (Sullivan et al., 2004; Wu et al., 2010; Richter et al., 2012). Thus, we decided to analyze the catalytic activity of JrPPO1 directly in crude leaf protein extracts, using the PPO-silenced lines as controls. Regardless of the type of PPO enzyme activity assay (polarimetric, spectrophotometric) or the substrate (monophenol, o-diphenol), PPO-silenced lines displayed essentially no activity, indicating that measured activity in wild-type protein extracts could be
attributed to JrPPO1. Catecholase activity was measured by monitoring the rate of oxygen consumption using a Clark oxygen electrode (polarimetric assay), and calculated kinetic parameters using native o-diphenols as substrates (Table I) were comparable to previously reported values in other plant species (Mesquita and Queiroz, 2013). Unlike many plant PPOs, JrPPO1 also displays tyrosinase activity and was capable of oxidizing all tested native monophenols. The rate of in vitro monophenol oxidation was only about 2% of the rate of o-diphenol oxidation (Supplemental Fig. S2), but this is comparable to characterized tyrosinase:catecholase activity ratios in apple (Malus domestica) and potato, two of the best-characterized plant tyrosinases (Selinheimo et al., 2007). Thus, JrPPO1 has the capability to oxidize a wide variety of native phenolic compounds in vitro, though this does not provide direct evidence that these same reactions occur in vivo.

Once PPO-silenced walnut plants were transferred from tissue culture to soil, it became apparent that many of the developing leaves displayed abnormal necrotic lesions. These lesions were absent from control plants germinated from the same somatic embryo line Chandler1 (CR1) and from walnut plants derived from CR1 embryos transformed with other RNAi constructs (Escobar et al., 2002). Based on extensive pathogen testing, we have concluded that these lesions are not associated with disease. Similar lesion mimic phenotypes have been observed in a variety of transgenic plants in which antioxidant levels (Takahashi et al., 1997), carbohydrate metabolism (Mittler and Rizhsky, 2000), defense-associated genes (Shirano et al., 2002), or phenolic metabolism (Di Fiore et al., 2002) have been manipulated. The absence of elevated reactive oxygen species levels, SA, and oxidative damage in PPO-silenced lines points to a subtle, potentially metabolic, connection between cell death and PPO in walnut.

Lacking any obvious mechanistic connections between the observed lesion mimic phenotype and PPO, we decided to carry out comparative transcriptomics and metabolomics studies on PPO-silenced and wild-type walnut leaves. Substantial alterations in both the transcriptome and the metabolome were observed, but the inherent challenge of this approach is separating the direct effects of silencing JrPPO1 from the nonspecific, indirect responses triggered by cell death and the lesion mimic phenotype. Thus, data analysis was focused on the following questions. (1) Which of the observed changes are caused directly by the absence of PPO? (2) Which of these direct effects is tied to the activation of cell death? (3) Which of the observed changes are simply general responses to the damage caused by the activation of cell death in PPO-silenced lines? In an attempt to answer the first of these questions, we focused our initial

Figure 6. Staining HCAAs in walnut leaf sections. A, Wild-type walnut leaf showing HCAA staining (blue-white fluorescence) in the vasculature. B, Leaf from a PPO-silenced line displaying pattern of HCAA staining similar to the wild type. C, HCAA staining in a region of a PPO-silenced leaf displaying lesion development. D, Light micrograph of the leaf section shown in C. The brown areas are necrotic lesions. All fluorescence micrographs were taken with identical illumination conditions. Bars = 100 μm.

Figure 7. Treatment of wild-type walnut leaves with exogenous tyramine leads to the development of necrotic lesions. A, Detached wild-type leaf incubated in water for 4 d. B, Detached wild-type leaf incubated in a 5 mM solution of tyramine for 2 d. C, Detached wild-type leaf incubated in a 5 mM solution of tyramine for 4 d. Bars = 2 cm.
data analysis on the phenolic compounds native to walnut that can serve as substrates for JrPPO1 in vitro (Fig. 2).

Assuming that JrPPO1 actively oxidizes a phenolic compound in vivo, the most naive model would posit that plants lacking PPO would accumulate its phenolic substrate(s) with a corresponding decrease in the reaction products. Levels of the o-diphenols caffeic acid and cryptochlorogenic acid (an isomer of chlorogenic acid, which was not detected) were not significantly altered in the PPO-silenced lines, so these compounds were not further investigated. By contrast, the o-diphenol catechin was substantially increased in PPO-silenced lines. However, this increase occurred in concert with elevated levels of many other flavonoids (most of which are not feasible substrates for PPO) and an apparent up-regulation of the phenylpropanoid pathway as a whole (Fig. 4). Thus, increases in catechin are more likely attributable to a general increase in phenylpropanoid metabolism than to a specific role as an in vivo substrate for JrPPO1. Similarly, elevated levels of p-coumaric acid are likely attributable to a generalized increase in the activity of the phenylpropanoid pathway, including induction of a cinnamic acid 4-hydroxylase gene, which encodes the enzyme responsible for the synthesis of p-coumaric acid (Fig. 4). Thus, none of the aforementioned phenolic compounds are likely to be direct targets of JrPPO1 activity in vivo.

By contrast, the effects of PPO silencing on the metabolism of Tyr do point to a direct role for JrPPO1 in phenolic metabolism in walnut. To be clear, Tyr levels (and the levels of Phe and Trp) were unaltered in PPO-silenced lines, as might be expected considering the complex feedback mechanisms that maintain aromatic amino acid homeostasis (Tzin and Galili, 2010). However, PPO-silenced lines displayed dramatic alterations in all three major pathways of Tyr catabolism, with increases in metabolites derived from the 4-hydroxyphenylpyruvate pathway (tocopherols), elevated tyramine levels, and decreased levels of f-DOPA-derived 5,6-dihydroxyindole and dopamine (Fig. 5). Quantitatively, these are among the largest changes observed in the metabolite profiling data, with tyramine among the five metabolites showing the largest increases in PPO-silenced lines (9.3-fold) and dopamine and 5,6-dihydroxyindole among the five metabolites showing the largest decreases in PPO-silenced lines (0.16- and 0.01-fold, respectively). As summarized in Figure 5, the production of 5,6-dihydroxyindole requires the o-hydroxylation of Tyr, and the production of dopamine requires the o-hydroxylation of either Tyr or tyramine. The enzymes responsible for these activities have not been characterized in plants (Lee and Facchini, 2011). JrPPO1 has demonstrated these activities in vitro (Fig. 2), and the silencing of JrPPO1 leads to clear increases in substrate (tyramine) or alternative metabolic pathways (4-hydroxyphenylpyruvate pathway) and clear decreases in products (5,6-dihydroxyindole, dopamine). Thus, we propose that JrPPO1 catalyzes the o-hydroxylation of Tyr and tyramine in vivo as a fundamental feature of aromatic amino acid metabolism in walnut leaves.

If JrPPO1 catalyzes the o-hydroxylation of Tyr and tyramine in intact cells, then the enzyme and its substrates must be located in the same subcellular compartment, at least transiently. Almost all PPOs are chloroplast localized (Steffens et al., 1994), and subcellular localization predictors place JrPPO1 in the chloroplast with high confidence (Predotar: plastid, 0.9; MultiLoc: chloroplast, 0.97; Small et al., 2004; Höglund et al., 2006). Likewise, the major enzymes of Tyr biosynthesis have been localized to the chloroplast (Tzin and Galili, 2010), placing JrPPO1 and Tyr in the same compartment. Tyr decarboxylase, the enzyme that converts Tyr to tyramine, has been localized to the cytosol in Arabidopsis (Lehmann and Pollmann, 2009), so tyramine would presumably need to be transported back to the chloroplast for catalysis by PPO.

The recent characterization of an Arabidopsis Tyr aminotransferase mutant provides an interesting parallel to our findings related to Tyr metabolism (Riewe et al., 2012). Tyr aminotransferase catalyzes the conversion of Tyr to 4-hydroxyphenylpyruvate (Fig. 5). As expected, the Tyr aminotransferase mutant showed decreased flux through the 4-hydroxyphenylpyruvate pathway, as demonstrated by large reductions in tocopherol levels. Tyramine was elevated approximately 2.5-fold in the mutant, demonstrating metabolic counterbalancing of the Tyr catabolic pathways, as was also observed in our PPO-silenced walnut plants. Dopamine was not detected in the samples, which is interesting given that Arabidopsis is one of the few plants that lack PPO genes (Tran et al., 2012).

If JrPPO1 plays a fundamental role in Tyr metabolism in walnut, then a direct product of disrupted Tyr metabolism should presumably be responsible for the observed lesion mimic phenotype in PPO-silenced lines. Previous studies have shown that the exogenous application of tyramine to rice cell cultures or to tobacco (Nicotiana tabacum), soybean (Glycine max), corn (Zea mays), and sunflower (Helianthus annuus) callus causes cell death (Christou and Barton, 1989; Negrel et al., 1993). Similarly, overexpression of Tyr decarboxylase in rice leads to massively elevated tyramine levels and stunted growth, fertility, and other toxicity symptoms (Kang et al., 2007). Exogenous application of tyramine to wild-type walnut leaves triggers the development of spontaneous necrotic lesions that are a near phenocopy of the lesion mimic phenotype observed in PPO-silenced lines (Fig. 7). Thus, the hyperaccumulation of the toxic metabolite tyramine may underlie the activation of cell death pathways in PPO-silenced walnut lines.

Many of the other observed changes in the leaf transcriptome and metabolome of PPO-silenced walnut lines may simply be stress responses associated with the activation of cell death and the lesion mimic phenotype. For example, the accumulation of HCAAs is typically associated with wounding, pathogen challenge, or other stressors (Facchini et al., 2002; Guillet and De Luca, 2005). Thus, the observed accumulation of HCAAs around necrotic lesions in the PPO-silenced walnut lines (Fig. 6) is likely a general response to the damage caused
by the lesions, not a direct effect of the absence of JrPPO1. Azelaic acid, a dicarboxylic acid that was recently characterized as a metabolic marker of oxidative membrane damage and cell death (Zoeller et al., 2012), is also highly elevated (22-fold) in PPO-silenced lines (Supplemental Table S3). The phenylpropanoid pathway and its products (e.g. flavonoids) are strongly induced by stress, including wounding and pathogen challenge (Dixon and Paiva, 1995; Dixon et al., 2002; Mikulic-Petkovsek et al., 2011). Likewise, synthesis of the Trp derivatives tryptamine and serotonin is up-regulated by pathogen challenge and in tissues undergoing abscission (Ramakrishna et al., 2011). It is also possible that Phe- and Trp-associated pathways are affected by the disruption of Tyr metabolism in PPO-silenced lines through feedback to the shikimate pathway, which generates the precursors of all aromatic amino acids (Tzin and Galili, 2010). Regardless of the regulatory mechanisms, alterations in HCAAs, azelaic acid, Phe metabolism, and Trp metabolism are most likely secondary effects of the silencing of JrPPO1.

An interesting exception to the general up-regulation of the phenylpropanoid pathway and its derivatives in PPO-silenced lines is the hydroxycoumarin esculentin (Fig. 4). The silencing of JrPPO1 caused dramatic reductions in esculentin, which was readily quantified in all wild-type walnut leaf samples, but was detectable in only one of the 16 samples derived from PPO-silenced lines (Fig. 4; Supplemental Table S3). Although the metabolism of esculentin is not well understood, proposed biosynthesis pathways start with coumaric acid (Blagbrough et al., 2010), an in vitro substrate of JrPPO1. Further, Sato (1967) previously implicated a chloroplast-localized phenolase (PPO) in the synthesis of esculentin based on in vitro studies using protein extracts from Saxifraga stolonifera. Our results provide support for the hypothesis that PPO is directly involved in esculentin biosynthesis in vivo, potentially via the oxidation of caffeic acid (Sato, 1967) or via the hydroxylation of umbelliferone (Garcia-Molina et al., 2013; Supplemental Fig. S8). Esculetin can act as an antioxidant and may contribute to plant defense responses (Blagbrough et al., 2010), but very little is known about its specific functional role(s). It is therefore unclear how the loss of esculetin might alter plant physiology and/or influence the lesion mimic phenotype in JrPPO1-silenced walnuts.

Overall, our results implicate JrPPO1 as a central enzyme in the metabolism of Tyr and esculetin in walnut, though further research (e.g. radioactive pulse labeling) will be required to definitively assign its role. It is unclear whether PPO enzymes serve similar functions in other plant species, but several previous studies suggest that our findings are not universal. Expression of an antisense PPO gene in tomato resulted in complete silencing of all members of the PPO gene family, but these PPO-silenced plants were phenotypically normal (Thiyapong et al., 2004). In an earlier study, the treatment of mung bean (Vigna radiata) seedlings with the PPO inhibitorotentoxin had no apparent effect on levels of the flavonoid rutin or total soluble hydroxyphenolics (Duke and Vaughn, 1982). As previously mentioned, Arabidopsis lacks PPO entirely (Tran et al., 2012). Thus, PPO may not play an important role in the metabolism of phenolic compounds in these plant species, as it does in walnut. Previous studies that have defined a metabolic role for PPOs in vivo have typically implicated biochemical pathways of limited taxonomic distribution. For example, PPO is involved in the biosynthesis of aurone pigments, but this pathway is primarily associated with the Compositae and Leguminosae (Nakayama et al., 2001). In creosote bush, PPO hydroxylates the phenolic compound larretricin, a key step in the biosynthesis of a group of 8′-lignans with very limited distribution (Cho et al., 2003). The biosynthesis of betalain pigments also involves a PPO, but this pathway is limited to the order Caryophyllales (Gandía-Herrero and García-Carmona, 2013). Both the PPO involved in betalain synthesis and JrPPO1 appear to utilize Tyr as a substrate, but the betalain biosynthesis pathway does not involve the production of dopamine (Gandía-Herrero and García-Carmona, 2013). As discussed above, our data suggests that JrPPO1 is the previously uncharacterized 6-hydroxylase that is required for the synthesis of dopamine from either Tyr or tyramine (Fig. 5). Dopamine is a central intermediate in the biosynthesis of isoquinoline alkaloids, a diverse and economically important group of secondary metabolites (e.g. berberine, morphine, codeine; Tzin and Galili, 2010). Unfortunately, almost nothing is currently known about the alkaloid composition of walnut, so future studies will be required to define the potential role for PPO in isoquinoline alkaloid biosynthesis, both in walnut and in other plant species that produce these biomedically significant secondary metabolites.

MATERIALS AND METHODS

Plant Transformation and Growth

An Agrobacterium tumefaciens binary vector designed to silence JrPPO1 was constructed by cloning the full-length JrPPO1 sequence into the pRi7 plasmid (Gleave, 1992) in the antisense orientation. The initial 1.2 kb of JrPPO1 was then cloned directly in front of the antisense sequence, generating two 1.2-kb complementary sequences (the 5′ end of JrPPO1 separated by a 0.6-kb spacer (the 3′ end of JrPPO1). This self-complementary cassette was then cloned into the binary vector pDU97.1005 (A. Dandekar, unpublished data), generating the JrPPO1 RNAi vector pDE00.0224. The JrPPO1 overexpression vector pDE98.0806 has been previously described (Escobar et al., 2008). Both vectors were electroporated into the disarmed A. tumefaciens strain C58.C1. A. tumefaciens-mediated transformation of the walnut (Juglans regia) somatic embryo line CR1 and subsequent regeneration of transgenic plants was performed as previously described (McGranahan et al., 1990). After transfer to soil (pots), plants were placed in a lath house (at the University of California, Davis) or outdoor enclosure (at California State University, San Marcos) under ambient conditions with regular irrigation.

PPO Enzyme Activity Assays

Total protein isolation and quantification from walnut leaves was performed as previously described (Escobar et al., 2008). To compare JrPPO1 activity against different 6-diphenol substrates, oxygen consumption was measured via polarimetric assay using a Clark oxygen electrode as previously described (Stewart et al., 2001). For typical polarimetric assays, substrate concentrations were set at 8 mM, but decreasing concentrations were utilized for selected substrates to calculate $K_m$ and $V_{max}$ using a Lineweaver-Burk plot. Due to relatively low tyrosinase activity of JrPPO1, activity assays utilizing monophenol substrates were performed spectrophotometrically, as previously described (Escobar et al., 2008). For all assays involving wild-type protein extracts, protein extracts from one or
more JrPPO1-silenced lines were utilized as negative controls, as activity measured in the wild type but absent from a PPO-silenced line should be attributable to JrPPO1. In addition, control activity assays were performed in the presence of 2 mM 1-kojic acid, a specific inhibitor of PPO (Constabel and Ryan, 1998).

Real-Time RT-PCR

Total RNA was extracted from the leaves of wild-type and PPO-silenced walnut lines using PureLink Plant RNA Reagent (Ambion) according to the manufacturer’s specifications. Isolated RNA was further purified using an RNeasy MiniElute Cleanup Kit (Qiagen), and then RNA concentration and quality were determined by spectrophotometric analysis and electrophoretic separation using an Experion Automated Electrophoresis System (Bio-Rad). Complementary DNA was synthesized with a QuantiTect Reverse Transcription Kit (Qiagen) using 2-μg aliquots of total RNA. Real-time PCR was performed using an Applied Biosystems 7000 machine with Platinum SYBR Green qPCR SuperMix (Applied Biosystems). The primers PPO1-F (5′-TCGTTGAAGGTAAAGCTGC-3′) and PPO1-R (5′-CCTTTTCTCTCTGCTCGAT-3′) were used for analysis of the JrPPO1 transcript. Primers for the 18S ribosomal RNA transcript, which was utilized as an internal standard, were as previously described (Huang et al., 2011). Cycle threshold values were used to calculate relative changes in gene expression using the 2^(-ΔΔCt) method (Livak and Schmittgen, 2001).

Physiological Characterization of Transgenic Lines

To measure H2O2 levels, leaves were first frozen in liquid nitrogen and crushed using a mortar and pestle. Two hundred milligrams of leaf powder were mixed with 1 mL of 5% (v/v) trichloroacetic acid, 50 mg of insoluble poly(vinyl-polypyrrolidone) (Sigma), and 50 mg of activated charcoal (Norit SA2, Acros Organics). The slurry was vortexed for 30 s and then centrifuged at 10,000g for 10 min and 4°C. The supernatant was diluted 1:10 in 0.25M sodium phosphate buffer (pH 7.0) and then assayed using an Amplex Red Hydrogen Peroxide/Peroxidase Assay Kit (Invitrogen) as previously described (Shin et al., 2015). As a means to measure lipid peroxidation/general oxidative damage, a thiobarbituric acid reactive substances assay was performed (Umbach et al., 2005). Frozen tissue powder was lyophilized and sent to Metabolon for targeted metabolite profiling. Metabolite extraction was performed as previously described (Evans et al., 2009). Metabolite extracts were subjected to three analyses: gas chromatography and mass spectrometry, ultra HPLC and tandem mass spectrometry optimized for basic metabolites, and ultra HPLC and tandem mass spectrometry optimized for acidic metabolites (Casool et al., 2013). Relative abundance was determined for 194 identified (named) metabolites. Wilcoxon’s rank sum tests were used to identify metabolites whose abundance differed significantly between grouped wild-type samples and grouped PPO-silenced samples. P values were corrected for multiple hypothesis testing (Benjamini and Hochberg, 1995), and adjusted P values ≤ 0.01 were considered statistically significant for the metabolite data set.

RNA-Seq

Leaf samples were collected from three wild-type trees and three JrPPO1-silenced trees (lines 7-8-1, 9-5-1, and 9-14). Each leaf sample consisted of three mature leaves and was immediately frozen in liquid nitrogen. RNA isolation and assessment of RNA quality was performed as described above for real-time RT-PCR. Samples utilized for library construction had an RNA Quality Indicator number greater than 8.0. Six sequencing libraries (three independent wild-type biological replicates and three independent PPO-silenced biological replicates) were made with Illumina TruSeq RNA Kit (Illumina) starting with 4 μg of total RNA, according to the manufacturer’s specifications. Each sequencing library was barcoded, pooled on a single Illumina flow cell lane, and then sequenced using an Illumina HiSeq 2000 (100-nucleotide paired-end reads) at the University of California, Davis Genome Center. Demultiplexed sequence reads were trimmed with custom scripts, including Sfffile (unpublished, available at https://github.com/ucdavis-bioinformatics/sfffile) to remove Illumina adapter contamination and sickle (unpublished, available at https://github.com/ucdavis-bioinformatics/sickle) to trim low-quality bases at the 3′ end of reads. Trimmed reads were aligned to a walnut transcriptome assembly consisting of 85,045 contigs using Burrows-Wheeler Aligner version 0.6.1 (Li and Durbin, 2009). The walnut transcriptome was previously assembled from approximately 1 billion 85-nucleotide RNA-seq reads derived from RNA isolated from 20 different walnut tissue samples (You et al., 2012). The BLAST2GO program (Conesa et al., 2005) was utilized for putative functional annotation of the walnut sequences. BLASTX alignment of the walnut sequences to the Arabidopsis (Arabidopsis thaliana) proteome was used to identify putative homologous genes, with an E value cutoff of 1e-05. Differential expression analysis of count data were performed using DESeq version 1.6 (Anders and Huber, 2010), with P values corrected for multiple hypothesis testing (Benjamini and Hochberg, 2000). Adjusted P values ≤ 0.05 were considered statistically significant for the RNA-seq data set. Contigs with less than 20 total counts were excluded from analysis.

Pathogen Testing

Walnut blight and walnut anthracnose can cause the development of necrotic lesions on walnut leaves. Thus, we attempted to isolate the causal agents of these diseases (Xanthomonas arboricola pv juglandis and Gnomonia leptostyla, respectively) from the lesions on the leaves of JrPPO1-silenced plants. Leaves were surface disinfested by immersion in 15% (v/v) household bleach and 0.1% (v/v) Tween 20 for 1 min followed by three rinses in sterile deionized water. One leaf disc (approximately 20 mm²) was then punched from the leaf (in tissue around a lesion), and the leaf disc was homogenized in sterile distilled water. Dilutions of the slurry were then plated on Difco nutrient agar for isolation of X. arboricola pv. juglandis (Belisario et al., 1999) or water agar supplemented with 40 mg L⁻¹ chloramphenicol and 40 mg L⁻¹ streptomycin for isolation of G. leptostyla (Belisario et al., 2008). Difco agar plates were incubated at 28°C for 3 d, and water agar plates were incubated at 25°C for 5 d. These experiments were repeated at least three times across multiple growing seasons.

To promote the rapid growth of potential pathogens, detached leaves from PPO-silenced plants were incubated in a moist chamber consisting of a sealed petri plate containing filter paper saturated with sterile water. These plates were placed in a 25°C incubator for 1 week and were monitored daily for lesion expansion, bacterial ooze formation, or the development of acervuli (fruiting bodies). Small black acervuli are diagnostic features of G. leptostyla (Cline and Neely, 1983).
expressed. This was because one of the three analyzed RNAi lines showed a relatively high level of \( \text{jrPPO1} \) counts compared with both of the other transgenic lines. (The three wild-type lines had 1,230, 2,545, and 1,252 \( \text{jrPPO1} \) counts, and the three RNAi lines had 301 [line 9-5-1], 549 [line 9-14], and 2,284 [line 7-8-1] counts). Despite this unexpected result, several lines of evidence suggest that the 7-8-1 transgenic line is silenced for \( \text{jrPPO1} \). First, PPO activity assays performed using the same tissue sample that was subsequently utilized for RNA isolation, and RNA-seq analysis showed less than 5% of wild-type-PPO activity in line 7-8-1 (in agreement with previous PPO activity assays performed on this line; Fig. 1). Second, subsequent analyses of six separate RNA samples isolated from 7-8-1 and analyzed by real-time PCR showed dramatically reduced \( \text{jrPPO1} \) transcript levels, mirroring the activity data (Supplemental Fig. S1). Finally, multidimensional scaling analysis of the overall RNA-seq transcriptome profile of 7-8-1 clearly groups this line with the other two PPO-silenced lines, separate from the wild-type samples (Supplemental Fig. S6). Thus, we ultimately decided to retain the RNA-seq data from line 7-8-1 in our final analysis.

To investigate the possibility of off-target effects of RNAi, TBLASTX was used to align the \( \text{jrPPO1} \) gene sequence to the sequences of all identified differentially expressed genes. No significant hits were found, suggesting that the observed changes in gene expression were not associated with the silencing of nontarget genes possessing a high degree of sequence similarity to \( \text{jrPPO1} \).

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession number SRP036081.

Supplemental Data

The following materials are available in the online version of this article.

Supplemental Figure S1. Real-time RT-PCR analysis of \( \text{jrPPO1} \) transcript levels in wild-type and PPO-silenced leaves.

Supplemental Figure S2. Comparison of tyrosinase and catecholase activities of \( \text{jrPPO1} \).

Supplemental Figure S3. SA levels in metabolite extracts from wild-type (CR1) and PPO-silenced walnut leaves.

Supplemental Figure S4. \( \text{H}_2\text{O}_2 \) levels in leaf extracts from wild-type walnut (CR1) and PPO-silenced lines.

Supplemental Figure S5. Malondialdehyde levels in metabolite extracts from wild-type (CR1) and PPO-silenced walnut leaves.

Supplemental Figure S6. Multidimensional scaling plot of log-transformed RNA-seq count data.

Supplemental Figure S7. Lignin analysis of wild-type (CR1) and PPO-silenced walnut lines.

Supplemental Figure S8. Two proposed esculetin biosynthesis pathways involving PPO.

Supplemental Table S1. List of differentially regulated contigs between wild-type and \( \text{jrPPO1} \)-silenced plants.

Supplemental Table S2. BioMaps analysis of genes differentially expressed in PPO-silenced walnut lines.

Supplemental Table S3. Comparison of metabolite levels in wild-type and PPO-silenced walnut leaves.

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LITERATURE CITED


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