

# Class II Cytochrome P450 Reductase Governs the Biosynthesis of Alkaloids<sup>1</sup>

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Expansion of the biosynthesis of plant specialized metabolites notably results from the massive recruitment of cytochrome P450s that catalyze multiple types of conversion of biosynthetic intermediates. For catalysis, P450s require a two-electron transfer catalyzed by shared cytochrome P450 oxidoreductases (CPRs), making these auxiliary proteins an essential component of specialized metabolism. CPR isoforms usually group into two distinct classes with different proposed roles, namely involvement in primary and basal specialized metabolisms for class I and inducible specialized metabolism for class II. By studying the role of CPRs in the biosynthesis of monoterpene indole alkaloids, we provide compelling evidence of an operational specialization of CPR isoforms in *Catharanthus roseus* (Madagascar periwinkle). Global analyses of gene expression correlation combined with transcript localization in specific leaf tissues and gene-silencing experiments of both classes of CPR all point to the strict requirement of class II CPRs for monoterpene indole alkaloid biosynthesis with a minimal or null role of class I. Direct assays of interaction and reduction of P450s *in vitro*, however, showed that both classes of CPR performed equally well. Such high specialization of class II CPRs in planta highlights the evolutionary strategy that ensures an efficient reduction of P450s in specialized metabolism.

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C.P., E.F., A.L., M.A.L., I.C., N.P., and M.C. performed the biochemical characterization of CPRs; E.F. studied protein subcellular localization and interaction; F.K. carried out silencing experiments; V.B., S.M., and B.S.-P. analyzed transcript distribution by *in situ* hybridization; T.D.d.B. and S.B. performed bioinformatics analyses; A.O. and G.G. conducted analysis of gene expression; N.G.-G., B.S.-P., L.A., and M.C. assisted in the supervision of this work; S.E.O. and V.C. conceived, supervised, and coordinated this study and wrote the article.

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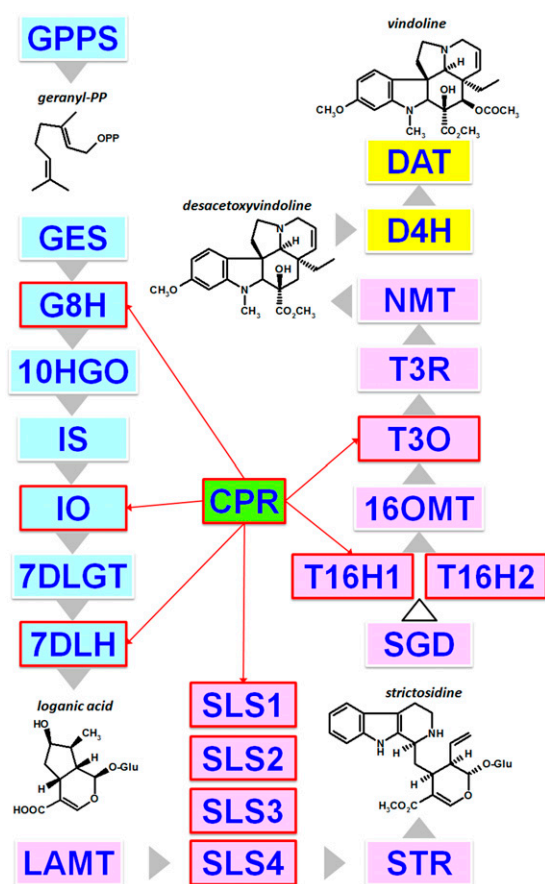
With more than 200,000 distinct molecules, the specialized metabolism of plants constitutes one of the main sources of bioactive natural compounds, with this large number reflecting the capacity of these sessile organisms to adapt to and interact with the environment. This singular complexity of compounds results from an evolutionary process that involves a dramatic diversification of plant metabolic pathways and the genes encoding the associated metabolic enzymes. From this perspective, cytochromes P450 (P450s) are a prototypical example of ubiquitous enzymes encoded by a gene superfamily that can carry out multiple types of reactions, among them hydroxylation, epoxidation, oxygenation, dealkylation, decarboxylation, C-C cleavage, and ring opening (Bak et al., 2011; Guengerich and Munro, 2013). P450s catalyze a considerable array of chemically challenging reactions in the biosynthesis of plant specialized metabolites, including phenylpropanoids, terpenoids, cyanogenic glycosides, and alkaloids (Mizutani and Ohta, 2010; Mizutani and Sato, 2011), in addition to various roles in plant primary metabolism, such as the biosynthesis of hormones

(Takei et al., 2004; Bak et al., 2011). Most characterized P450s perform single oxidation reactions, but growing evidence now also points to the existence of multistep oxidations catalyzed by a single P450 (Guengerich et al., 2011). However, in all P450-catalyzed reactions, the catalytic cycle always requires two one-electron transfer steps from NADPH into the prosthetic heme. This transfer occurs through the FAD and FMN domains of NADPH-cytochrome P450 reductases (CPRs), although, in some cases, the second electron transfer also can arise from cytochrome *b*<sub>5</sub> (Munro et al., 2013). The absolute requirement of CPR thus makes this flavoprotein a cornerstone in P450 activities and, consequently, in specialized and primary metabolic pathways. As such, the physical interactions between P450s and CPRs that guide electron shuffling directly influence P450 activities and are facilitated by membrane anchoring of both types of proteins, which are typically localized to the endoplasmic reticulum (ER; Hasemann et al., 1995; Ro et al., 2002; Denisov et al., 2007). Elucidation of the three-dimensional structure of the rat CPR suggested that P450-CPR interactions occur through both ionic interactions involving the FMN domain of CPRs and hydrophobic interactions between the membrane domains of CPRs and P450s (Wang et al., 1997). The low CPR:P450 ratio, measured at approximately 1:15 in liver, also implies a potential competition between P450s for these accessory proteins in plants; as a consequence, some kind of logistic control must be employed to ensure the coordinated operation of all P450s belonging to similar biosynthetic pathways (Shephard et al., 1983).

In contrast to yeasts and mammals that harbor only a single CPR, vascular plants have evolved two or three CPR isoforms, as reported for instance in Jerusalem artichoke (*Helianthus tuberosus*; Benveniste et al., 1991), poplar (*Populus* spp.; Ro et al., 2002), parsley (*Petroselinum crispum*; Koopmann and Hahlbrock, 1997), cotton (*Gossypium hirsutum*; Yang et al., 2010), and winter cherry (*Withania somnifera*; Rana et al., 2013). The *Arabidopsis* (*Arabidopsis thaliana*) genome contains two genes encoding functionally active CPR genes (*ARABIDOPSIS THALIANA* P450 REDUCTASE), named *ATR1* and *ATR2*, and a third more distant gene (*ATR3*) whose expression product was not able to reduce P450s in vitro (Urban et al., 1997; Mizutani and Ohta, 2010; Varadarajan et al., 2010). By contrast, poplar and *Nothapodytes foetida* possess three genes encoding genuine CPRs that are similar to *ATR1* and *ATR2* and an additional predicted *ATR3*-like isoform (Ro et al., 2002; Huang et al., 2012). Homologs of *ATR1* and *ATR2* in flowering plants are highly conserved (65%–80%) and are clustered into two distinct phylogenetic groups on the basis of their N-terminal sequences. The first cluster (*ATR1* homologs, class I) contains sequences from eudicotyledons, while the second (class II) are found in both monocotyledons and eudicotyledons (Ro et al., 2002). Homologs of *ATR3* fall into a distinct third cluster (Varadarajan et al., 2010). While CPRs from both class I and class II can reduce P450 with an apparent similar efficiency, their expression profiles are different

(Urban et al., 1997; Ro et al., 2002; Mizutani and Ohta, 2010). CPR1s belonging to class I, such as *ATR1*, are constitutively expressed, while CPR2s from class II (e.g. *ATR2*) are inducible by environmental stimuli such as wounding, pathogen infection, or light exposure (Koopmann and Hahlbrock, 1997; Ro et al., 2002; Schwarz et al., 2009; Mizutani and Ohta, 2010; Yang et al., 2010; Rana et al., 2013). These observations suggest that each CPR isoform has dedicated physiological roles, ensuring that plants can meet the reductive demand of P450-mediated reactions. It is now believed that constitutively expressed CPRs (CPR1s, class I) ensure basal P450 activities in primary metabolism or in the constitutive synthesis of specialized metabolism, while inducible CPRs (CPR2s, class II) serve in adaptation mechanisms or in defense reactions, including the elicited biosynthesis of specialized metabolites. This hypothesis has been partially confirmed by establishing a correlation between *ATR2* activity and lignin biosynthesis in *Arabidopsis* as well as between *CPR1* expression and basal pungent alkaloid synthesis in *Capsicum* spp. (Mazourek et al., 2009; Sundin et al., 2014). In contrast, CPR-like (*ATR3*) homologs are poorly characterized but appear to be essential for embryo development (Varadarajan et al., 2010). However, more investigations are required to firmly establish the functional specificity of all CPRs in plants.

For more than 40 years, specialized metabolism has been investigated using the Madagascar periwinkle (*Catharanthus roseus*), which serves as an excellent case study for the analysis of plant specialized metabolism. This plant most notably synthesizes alkaloids from the monoterpene indole alkaloid (MIA) family that includes valuable compounds such as the antineoplastic vinblastine and vincristine. These MIAs result from a long and complex biosynthetic pathway whose characterization has made great progress over the last 5 years due to the development of large sets of transcriptomic data and a draft genome sequence (Góngora-Castillo et al., 2012; Van Moerkercke et al., 2013; Xiao et al., 2013; Dugé de Bernonville et al., 2015a; Kellner et al., 2015b). Within the 30 to 50 enzymatic steps predicted to form the network of reactions of the MIA biosynthetic pathway, no less than 11 P450s have been identified to date (Fig. 1): geraniol 8-hydroxylase (G8H [CYP76B6]; Collu et al., 2001), iridoid oxidase (IO [CYP76A26]; Miettinen et al., 2014; Salim et al., 2014), 7-deoxyloganic acid 7-hydroxylase (7DLH [CYP72A224]; Salim et al., 2013; Miettinen et al., 2014), four isoforms of secologanin synthase (SLS1–SLS4 [CYP72A1]; Irmeler et al., 2000; Brown et al., 2015; Dugé de Bernonville et al., 2015b), two isoforms of tabersonine 16-hydroxylase (T16H1 and T16H2 [CYP71D12 and CYP71D351]; Schröder et al., 1999; Guirimand et al., 2011b; Besseau et al., 2013), tabersonine 19-hydroxylase (T19H [CYP71B1]; Giddings et al., 2011), and 16-methoxytabersonine 3-oxygenase (T3O [CYP71D1]; Kellner et al., 2015a; Qu et al., 2015b). While some of these enzymes catalyze a single oxygenation reaction, including hydroxylation (7DLH, T16H1,



**Figure 1.** Main characterized steps of MIA biosynthesis in *C. roseus* aerial organs highlighting several steps involving P450s. The color code indicates the cellular organization of the pathway as follows: blue, pink, and yellow rectangles for compartmentation in internal phloem-associated parenchyma (IPAP), epidermis, and laticifers/idioblasts, respectively. Noteworthy metabolites are depicted, such as geranyl diphosphate, which constitutes the entry point of MIA biosynthesis; loganic acid and desacetovindoline, which are transported from IPAP to epidermis and from epidermis to laticifers/idioblasts, respectively; strictosidine, the first MIA; and vindoline, which is the main MIA accumulated in leaves. P450s are highlighted by red rectangles, and reducing power by CPR is indicated by red arrow lines. The white arrowhead indicates multiple not yet discovered enzymes. GPPS, Geranyl diphosphate synthase; GES, geraniol synthase; G10H (CYP76B6), geraniol 10-hydroxylase; 10HGO, 10-hydroxygeraniol oxidoreductase; IS, iridoid synthase; 7DLGT, 7-deoxyloganic acid glucosyltransferase; 7DLH (CYP72A224), 7-deoxyloganic acid hydroxylase; LAMT, loganic acid *O*-methyltransferase; STR, strictosidine synthase; SGD, strictosidine  $\beta$ -glucosidase; 16OMT, 16-hydroxytabersonine *O*-methyltransferase; NMT, 3-hydroxy-16-methoxy-2,3-dihydrotabersonine *N*-methyltransferase; T3R, tabersonine 3-reductase; D4H, desacetovindoline 4-hydroxylase; DAT, deacetylindoline 4-acetyltransferase.

T16H2, and T19H) or epoxidation (T3O), unusual reactions also have been reported, such as the ring opening of loganin to yield secologanin (SLS) or the three-step oxidation of nepetalactol to form 7-deoxyloganic acid (IO). In addition to this diversity of reactions, a complex spatiotemporal organization of these P450-mediated enzymatic steps also has been

reported in periwinkle leaves, with the first MIA biosynthetic conversions (G8H to 7DLH) occurring in internal phloem-associated parenchyma (IPAP) and the remaining reactions up to T3O and the next two enzymatic steps in epidermis (Fig. 1; Courdavault et al., 2014; Miettinen et al., 2014; Salim et al., 2014; Qu et al., 2015b). As a matter of fact, this multicellular organization of the MIA biosynthetic pathway constitutes the first layer of the physiological processes regulating MIA formation. Interestingly, the periwinkle CPR was one of the first plant CPRs to be purified and cloned (Madyastha and Coscia, 1979; Meijer et al., 1993). Due to the presence of a hydrophobic residue region at its N-terminal end, this protein groups with class II CPRs, in agreement with its transcriptional regulation in response to fungal elicitor preparation and to jasmonate (Meijer et al., 1993; Collu et al., 2001; Ro et al., 2002). Although only one CPR had been cloned in *C. roseus* at the start of this study, the occurrence of multiple isoforms was expected and is now supported by transcriptomic and genomic data (Canto-Canché and Loyola-Vargas, 2001). However, no formal relationship had been established between the periwinkle CPRs and the biosynthesis of MIA. This prompted us to take advantage of the richness of the *C. roseus* MIA metabolism to accurately explore the role of each class of CPR in specialized metabolism. By combining biochemical characterization, protein interaction analyses, mapping of cellular gene expression profiles, and gene-silencing approaches, our data provide new evidence for the predominant role of class II CPRs in MIA/specialized metabolism.

## RESULTS

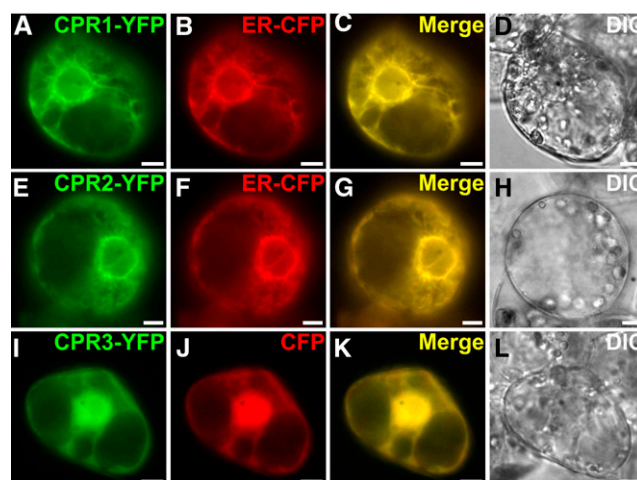
### Identification of *C. roseus* CPR Homologs

In addition to a contig corresponding to the CPR cDNA (X69791), which was cloned previously by Meijer et al. (1993), interrogation of *C. roseus* transcriptomic resources led to the identification of two additional contigs homologous to CPR (Supplemental Table S1). Proteins deduced from contig CPR candidate 1 and contig CPR candidate 2 displayed 67% and 25% identity with the originally identified CPR, respectively (Supplemental Fig. S1). According to previously published classifications of CPRs (Ro et al., 2002; Jensen and Moller, 2010; Varadarajan et al., 2010), phylogenetic analyses demonstrated that each deduced protein clusters in distinct phylogenetic subgroups (Supplemental Fig. S2). While the original CPR is categorized as a class II CPR, contig CPR candidate 1 falls into the class I CPRs, which is hypothesized to be mostly dedicated to basal metabolism. Contig CPR candidate 2 clusters in class III CPRs, whose prominent member in Arabidopsis corresponds to ATR3. Based on this result and on the Arabidopsis nomenclature, contig CPR candidate 1, the original CPR, and contig CPR candidate 2 were renamed CPR1, CPR2, and CPR3/diflavin reductase (DFR), respectively. *C. roseus* genome

analysis also revealed that the three corresponding genes were present at one copy per haploid genome and spanned over 18 exons for CPR1 (CRO\_T001672) and CPR2 (CRO\_T031702) and 12 exons for CPR3 (CRO\_T033752; Supplemental Fig. S3). The genomic organization of CPR1 and CPR2 is similar regarding intron positions and intron/exon sizes, except for the first intron of CPR1, which is roughly 7-fold longer than the first intron of CPR2. Such similarity may reflect the gene duplication event that led to both CPR appearances. Therefore, these results suggest that *C. roseus* contains two CPRs, CPR1 (KJ701028) and CPR2 (X69791), potentially associated with basal and inducible/specialized metabolism, respectively, and one more distant copy, CPR3/DFR (KM111538), a likely ortholog of ATR3, the function of which remains unclear.

### Sequence Analysis and Subcellular Localization of *C. roseus* CPRs

Analysis of the deduced protein sequences of *C. roseus* CPRs revealed that both CPR1 and CPR2 are characterized by (1) the presence of conserved FMN-, FAD-, and NADPH-binding domains that have been implicated in electron transfer and (2) identical residues predicted to be involved in interactions with P450s (Jensen and Moller, 2010; Supplemental Figs. S4 and S5). Both proteins also bear a predicted membrane-spanning domain at their N-terminal ends (Supplemental Fig. S6) that has been shown to mediate ER anchoring in poplar (Ro et al., 2002). To investigate whether this sequence does in fact anchor the protein to the ER membrane, the subcellular localizations of *C. roseus* CPR1 and CPR2 were expressed as C-terminal yellow fluorescent protein (YFP) fusion proteins (CPR1-YFP and CPR2-YFP) to avoid interference with the membrane-spanning domain. In *C. roseus* transiently transformed cells, the fusion proteins exhibited a network-shaped fluorescent signal that perfectly merged with the signal of the ER-cyan fluorescent protein (CFP) marker (Fig. 2, A–H), suggesting that both CPR1 and CPR2 are anchored to the ER, in agreement with the classical localization pattern of P450s. When mutants of CPR1 and CPR2 that lacked this predicted transmembrane domain were expressed as YFP-labeled fusions, the previously observed localization pattern was disrupted, while the fusion of these membrane domains to YFP enabled ER anchoring. These experiments clearly demonstrate that this predicted membrane-spanning domain is necessary and sufficient to ensure ER localization/anchoring (Supplemental Fig. S7). In addition, in agreement with the assigned CPR classification, we noted that CPR1 and CPR2 differ in the length of the protein sequence preceding the membrane-spanning domain: CPR1 has only a short stretch of residues, while CPR2 exhibits an extended amino acid sequence, enriched in Ser residues, that was initially, but wrongly, considered to be a plastid-targeting sequence (Supplemental Fig. S1; Ro et al., 2002). CPR3/DFR also displays conserved FMN-, FAD-, and NADPH-binding domains but shows

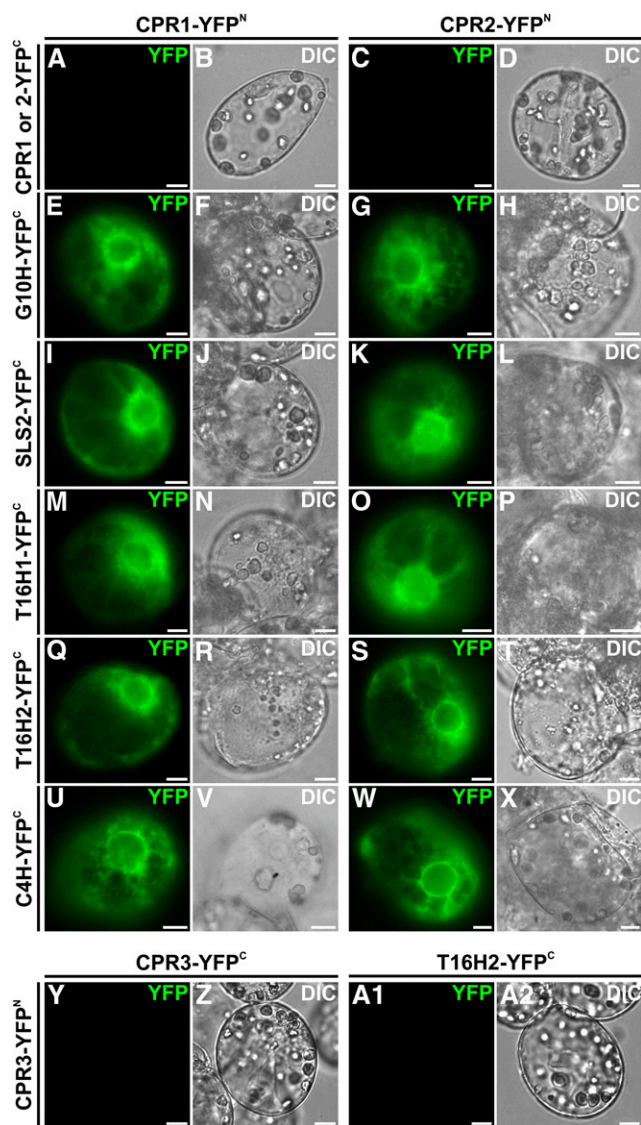


**Figure 2.** CPR1 and CPR2 are located to the ER, while CPR3 is a nucleocytoplasmic protein. *C. roseus* cells were transiently transformed with the CPR1-YFP (A), CPR2-YFP (E), or CPR3-YFP (I) expressing vectors in combination with plasmids expressing either an ER-CFP marker (B and F) or a nucleocytoplasmic marker (CFP; J). Colocalization of the fluorescent signals appears on the merged images (C, G, and K). Cell morphologies (D, H, and L) were observed with differential interference contrast (DIC) imaging. Bars = 10  $\mu$ m.

substantial differences in the P450-interacting region compared with CPR1 and CPR2 (Supplemental Fig. S5). Moreover, CPR3/DFR lacks the N-terminal membrane-spanning domain, explaining the nucleocytoplasmic localization observed in *C. roseus* cells transformed with the CPR3-YFP fusion protein (Fig. 2, I–L). This nucleocytoplasmic localization was observed previously with ATR3 (Varadarajan et al., 2010). Therefore, the distinct subcellular localization patterns, along with the sequence differences in the P450-interacting domain in CPR1 and CPR2 compared with CPR3/DFR, suggest the existence of distinct sets of interacting partners for these proteins.

### CPR1 and CPR2, But Not CPR3/DFR, Interact and Reduce *C. roseus* P450s Associated with Specialized Metabolism

To gain insight into possible selective interactions among the P450s and CPRs of *C. roseus*, bimolecular fluorescence complementation (BiFC) analyses were conducted by coexpressing each of the three CPRs with several P450s from the MIA biosynthetic pathway, including G8H, SLS2, T16H1, and T16H2 (Figs. 1 and 3). An additional test also was conducted with cinnamate 4-hydroxylase (C4H; CYP73A5) from *C. roseus*, since this P450 corresponds to the first P450 of the phenylpropanoid biosynthetic pathway, thus acting in a distinct specialized metabolic pathway. Moreover, the Arabidopsis C4H ortholog has been shown to be reduced with a similar efficiency by both ATR1 and ATR2, thereby providing a point of reference (Hotze et al., 1995; Mizutani and Ohta, 2010).



**Figure 3.** CPR1 and CPR2 but not CPR3 interact with G10H, SLS2, T16H1, T16H2, and C4H. CPR and P450 interactions were analyzed by BiFC in *C. roseus* cells transiently transformed by plasmids encoding fusions indicated on the top (CPR1 or CPR2 fused to the split YFP<sup>N</sup> fragment) and on the left (CPR1, CPR2, G10H, SLS2, T16H1, T16H2, and C4H fused to the split YFP<sup>C</sup> fragment). The YFP signal emanating from BiFC complex reformation is shown in green false color (A, C, E, G, I, K, M, O, Q, S, U, W, Y, and A1), and cell morphology is observed with differential interference contrast (DIC) imaging (B, D, F, H, J, L, N, P, R, T, V, X, Z, and A2). Bars = 10  $\mu$ m.

As described above, CPRs and these five P450s were fused to the N-terminal end of the split YFP fragments (N-terminal split YFP [YFP<sup>N</sup>] for CPRs and C-terminal split YFP [YFP<sup>C</sup>] for P450s) to preserve ER-anchoring capacities following transient expression in *C. roseus* cells. Interestingly, while no self-interactions were observed for either CPR1 or CPR2 (Fig. 3, A–D), the appearance of a strong fluorescent signal in the combinations of each CPR with all of the P450s suggested that CPR1 and CPR2

were capable of interaction with each of the five tested P450s (Fig. 3, E–X). By contrast, no interactions of CPR3/DFR, either with itself or with any of the tested P450s, were observed, as exemplified with T16H2 (Fig. 3, Y–A2). This may be linked to the neofunctionalization of class III CPRs and/or to the lack of the N-terminal membrane-spanning domain in CPR3/DFR.

Additionally, the capacity of the periwinkle CPR1 and CPR2 to reduce G8H, SLS2, T16H1, T16H2, and C4H was analyzed by conducting functional assays with proteins heterologously expressed in *Saccharomyces cerevisiae*. The resulting activities were compared with those measured in the WAT11 strain expressing an Arabidopsis CPR and with the activity engendered by the endogenous yeast (*S. cerevisiae*) CPR (Table I; Supplemental Table S2). Since partial losses of activity were observed when *C. roseus* CPRs were expressed with tags, as reported for CPR from *Camptotheca acuminata* (Qu et al., 2015a), assays were performed with untagged proteins using unpurified microsomes. Interestingly, in agreement with the BiFC results, we noted that both CPR1 and CPR2 were able to reduce the five tested P450s, as revealed by the conversion of the specific substrates of each of the five P450s. Moreover, although absolute quantification of CPR/P450 activity, which requires tagged protein, was not achievable, the relative activity of each of the P450s was measured to be within the same order of magnitude, suggesting that both CPR1 and CPR2 may reduce P450s *in vitro* with a similar efficiency, as reported previously for C4H in Arabidopsis (Mizutani and Ohta, 2010). Furthermore, we also noticed that CPR3/DFR was unable to reduce T16H, which is consistent with the lack of interaction observed in BiFC assays. To determine whether the inactivity resulted from a lack of ER membrane anchoring, the first 74 residues of CPR2 encompassing the membrane-spanning domain were fused to the N terminus of CPR3/DFR, which resulted in ER localization, as revealed by YFP imaging in *C. roseus* cells (Supplemental Fig. S8). However, this modification was unable to restore P450 activities (at least with SLS2, T16H1, and T16H2) in yeast (*S. cerevisiae*), strongly suggesting that CPR3/DFR reduces different enzymes and acts in different metabolic pathways. As a consequence, CPR3/DFR was renamed DFR in agreement with the presence of two flavin reductase domains and the apparent lack of P450 reduction capacity.

#### CPR1 and CPR2 Display Ubiquitous Expression Patterns Associated with Specific Sets of Genes

The putative specialization of CPRs with respect to MIA biosynthesis was investigated subsequently at the whole-plant level through gene expression analyses. By studying transcript abundance in *C. roseus* transcriptomic data sets (Góngora-Castillo et al., 2012; Dugé de Bernonville et al., 2015b), we observed that CPR1, CPR2, and DFR genes were expressed in all organs associated with MIA biosynthesis, including roots,

**Table 1.** *CPR1* and *CPR2* reduce *C4H*, *G8H*, *SLS2*, *T16H1*, and *T16H2* with an apparent similar efficiency

Substrate conversion rates (%) were determined using crude protein extract of the WT303 yeast strain expressing each CPR/P450 pair (*C4H*, *SLS2*, *T16H1*, and *T16H2* assays) or microsomes (*G8H* assay) by the addition of NADPH as an electron donor. Yeast endogenous CPR activity was estimated by measuring P450 activity in similar conditions without expression of the periwinkle CPRs and was subtracted from activities measured with *CPR1* and *CPR2*. All assays were conducted independently three times with at least three technical replicates.

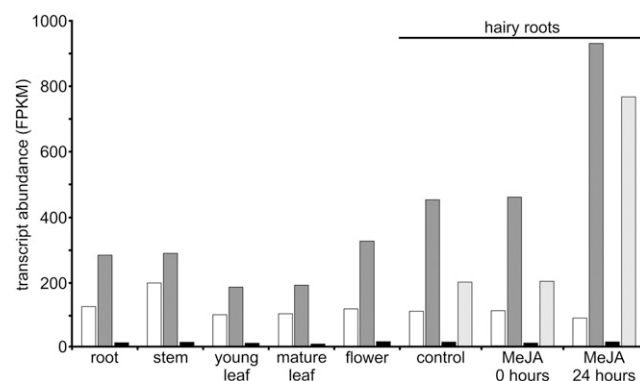
Assay	Substrate	CPR1	CPR2
C4H (CYP73)	Cinnamic acid	31.2 ± 0.8	26.2 ± 1.61
G10H (CYP76B6)	Geraniol	65.2 ± 0.95	70.3 ± 1.12
SLS2 (CYP71A2)	Loganin	25.6 ± 1.12	39.1 ± 3.7
T16H1 (CYP71D12)	Tabersonine	46 ± 0.83	47.3 ± 0.7
T16H2 (CYP71D351)	Tabersonine	60.9 ± 1.07	60.2 ± 0.25

stems, young/mature leaves, and flowers, although *DFR* expression was very low, as reported previously for *ATR3* (Fig. 4; Varadarajan et al., 2010). *CPR1* and *CPR2* exhibited a similar pattern of expression in organs, but *CPR2* expression was always almost 2-fold higher than *CPR1* expression. Furthermore, while *CPR1* levels were not affected by methyl jasmonate treatment, *CPR2* was induced by this hormone, which is consistent with the role of class II *CPR* in induced pathways. This ubiquitous distribution of *CPR1* and *CPR2* transcripts in periwinkle organs was confirmed by quantitative PCR (qPCR) and also was compared with the expression of *G10H*, *SLS2*, *T16H1*, *T16H2*, and *C4H* (Supplemental Fig. S9). Although we noted that CPR- and P450-encoding genes were coexpressed in the different MIA-producing organs with some profile specificities inherent to each P450, this cooccurrence of gene expression led us to speculate that *CPR1* as well as *CPR2* both can potentially reduce P450s associated with MIA biosynthesis.

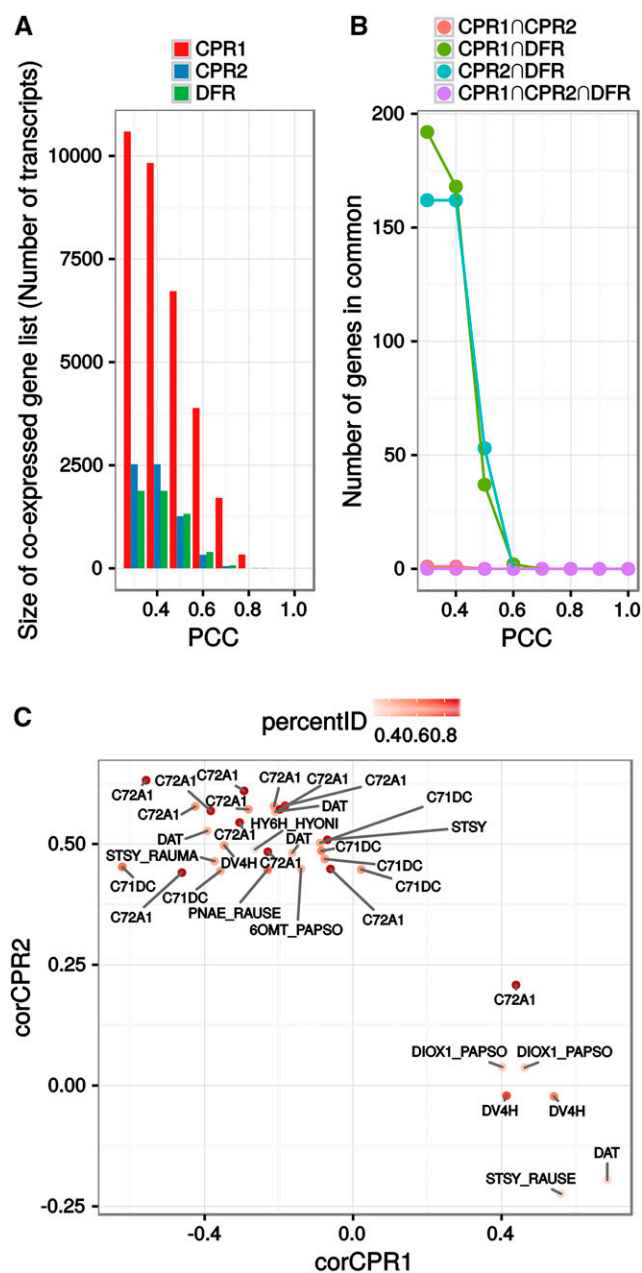
As a consequence, by taking advantage of the optimized transcriptomic *C. roseus* data set (Dugé de Bernonville et al., 2015b), a global analysis of the periwinkle genes coexpressed with each CPR was undertaken by calculating the Pearson correlation coefficient (PCC) of *CPR1*, *CPR2*, or *DFR* with each of the 58,338 other transcripts found in this assembly. This data set was obtained previously by combining individual transcriptomes assembled from sequencing runs available on the Sequence Read Archive. Expression levels within this transcriptome were computed with Salmon (Patro et al., 2015) on 39 samples (including 16 with paired-end sequencing design). This consensus assembly has the advantage of integrating expression measurements over a wider range of samples than was reported previously while integrating sequence polymorphisms by the use of sequences that were clustered from individual assemblies (Dugé de Bernonville et al., 2015b). PCC were calculated on  $\log_2$ -transformed TPM to switch the expected Poisson distribution of RNA sequencing expression data to a Gaussian shaped distribution (Supplemental Fig. S10). Different lists of

coexpressed genes for each CPR/*DFR* were then established at different PCC values (Fig. 5A), and we subsequently examined their similarities (Fig. 5B). Interestingly, the number of correlated genes differed strongly between the two CPRs and *DFR*. For instance, at PCC > 0.4 (FDR-corrected  $P < 0.05$ ), we counted 9,831 transcripts for *CPR1*, 2,525 for *CPR2*, and 1,881 for *DFR* (Fig. 5A). In fact, *CPR1* had the highest number of strongly coexpressed genes (for PCC > 0.8, 336 transcripts for *CPR1*, one for *CPR2*, and seven for *CPR3*). In addition, intersection sizes indicated that these coexpressed gene lists did not overlap unless considering genes with lower PCC values (e.g. PCC < 0.3), thus arguing for a specialization of CPR expression. This revealed once again that periwinkle CPRs and *DFR* are transcriptionally unrelated. In this analysis, coexpressed gene lists for each CPR were set with a PCC threshold of 0.4 and an FDR-corrected  $P$  of 0.05. While this PCC might keep weak correlations, our  $P$  threshold was expected to conserve only significant positive correlations. In addition, this PCC cutoff value was used to retrieve enough information (in particular for *CPR2* and *DFR*) in each list to perform Gene Ontology (GO) term and UniProt keyword enrichment as well as promoter motif analyses. Indeed, the transcriptional unrelatedness was further confirmed by an analysis of overexpressed GO terms represented in coexpressed genes with a chosen low PCC value and of keywords from UniProt (Supplemental Fig. S11, A and B).

While more genes coexpressed with *CPR1*, we noted that a higher proportion of P450s were associated with *CPR2* (54 out of 2,525 coexpressed genes) compared with *CPR1* (49 out of 9,831) or with *DFR* (six out of 1,881; Supplemental Table S3, A and B), based on a total of around 400 predicted P450 coding sequences (that may exhibit redundancy). A basic analysis of the predicted P450 functions revealed that more than 20% of



**Figure 4.** *CPR1*, *CPR2*, and *DFR* transcript abundance in *C. roseus* organs and response to methyl jasmonate (MeJA) treatment. The abundance of transcripts was determined by comparing fragments per kilobase of transcript per million mapped reads (FPKM) values of *CPR1*, *CPR2*, and *DFR* loci from the main *C. roseus* transcriptomic data set. In hairy roots, estimation of the abundance of the transcripts encoding strictosidine synthase (STR), a well-known MIA biosynthetic gene inducible by methyl jasmonate, was included as a control. *CPR1*, White bars; *CPR2*, dark gray bars; *DFR*, black bars; STR, light gray bars.



**Figure 5.** Analysis of CPR1, CPR2, or DFR gene coexpression correlation. Lists of coexpressed genes were obtained after calculating the PCC of *C. roseus* CPR expression levels with each other transcript found in the CDF97 assembly and setting different PCC threshold values. For each threshold, only associations with a false discovery rate (FDR)-corrected  $P < 0.05$  were kept. A, Sizes of coexpressed gene lists for each CPR at different PCC thresholds. B, Sizes of intersections between coexpressed gene lists. C, Comparison of PCCs of transcripts related to alkaloid metabolism (according to the UniProt annotation) with CPR1 and CPR2. The corresponding protein name is followed by the name of the initial plant (CATRO, *C. roseus*; HYONI, *Hyoscyamus niger*; PAPSO, *Papaver somniferum*; RAUMA, *Rauwolfia mannii*; and RAUSE, *Rauwolfia serpentina*). For full gene information, see Supplemental Table S3B.

the CPR1-associated P450s (10 within 49 identified P450s but 10 of 35 P450s displaying predicted functions) acted in primary metabolism such as hormone synthesis, whereas the remaining P450s were potentially involved in a specialized metabolism that appears to correspond to phenylpropanoid biosynthesis. By contrast, only three occurrences potentially associated with primary metabolism were found in the CPR2 list, whereas all the others (40 within 54 identified P450s but 40 of 43 with predicted functions) were associated with specialized metabolism, including MIA biosynthesis. Furthermore, the fact that only six P450s were associated with DFR strengthens the proposition that this protein is not involved in the reduction of P450s, in agreement with biochemical activity assays.

Such discrepancies also were confirmed by the analysis of GO terms of the whole gene list of each CPR/DFR. Indeed, GO terms found in the coexpressed gene list of CPR1 correspond to large cellular functions (nucleotide binding and protein binding), supporting its involvement in a basal/primary metabolism (Supplemental Fig. S11; Supplemental Table S3, A and B). By contrast, many genes coexpressed with CPR2 were associated with oxidation and metabolic processes, reflecting its transcriptional association with metabolic processes requiring many P450s, such as specialized metabolisms. In fact, more than 25 genes from the MIA biosynthetic pathway (including P450s) were identified in the coexpressed gene list of CPR2, whereas only a single MIA gene was found in CPR1 and DFR lists (Fig. 5C; Supplemental Table S3, A and B). Accordingly, a motif enrichment analysis in the promoter regions (500 bp upstream the ATG codon, when available) of genes coexpressed with each CPR revealed associations with specific metabolisms (Supplemental Table S3C). Upstream regions of genes coexpressed with CPR1 were enriched with the CACRTR and ATRYAC motifs and were found to be associated with primary metabolism and hormone response. The latter motif also was present in the promoter of CPR1, although only 300 bp was available in the 5' sequence. Additionally, promoter regions of genes coexpressed with CPR2 were enriched significantly with the CACGHG motif. CPR2 also contained this motif, which was shown previously in the promoter of strictosidine synthase and is able to bind MYC transcription factors, probably involved during jasmonic acid signaling (Chatel et al., 2003). Altogether, these results emphasize the occurrence of specific metabolic associations for each CPR (e.g. CPR1 with primary and basal specialized metabolisms and CPR2 with specialized metabolisms and, notably, the biosynthesis of MIA). Due to the absence of P450 gene expression association and reduction capacity, DFR was not investigated further in this study.

### CPR2 Is Expressed in Leaf and Cotyledon Tissues Hosting MIA Biosynthetic Steps Catalyzed by P450s

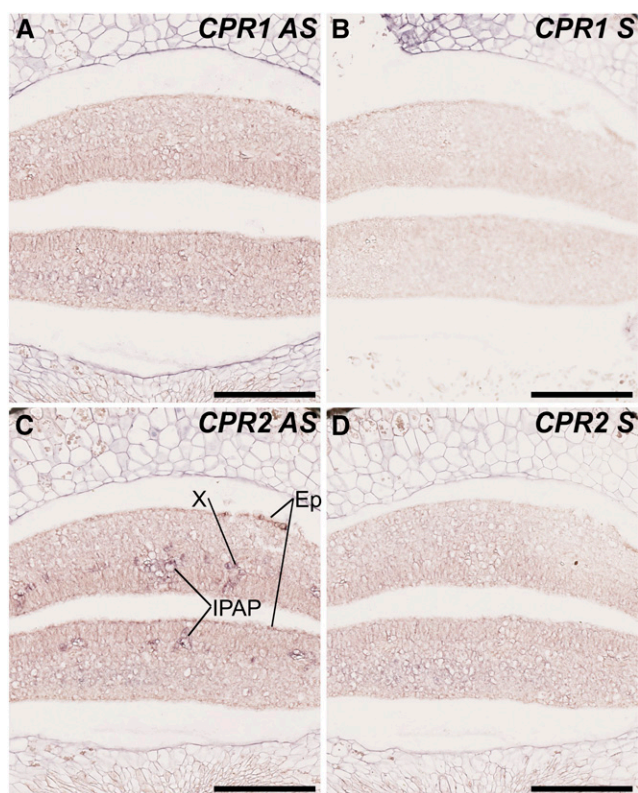
Although our global analysis of gene coexpression provides valuable information regarding CPRs/metabolic

pathway associations, these analyses cannot actually reflect the intricate gene coexpression networks occurring at the tissue and/or cellular levels and are required to ensure efficient P450 reduction. Therefore, since the MIA and phenylpropanoid biosynthetic pathways display complex spatial organizations in *C. roseus* (St-Pierre et al., 1999; Burlat et al., 2004; Mahroug et al., 2006; for review, see Courdavault et al., 2014), we next studied the cellular distribution of *CPR1* and *CPR2* transcripts in cotyledons of germinating seedlings. Given the high expressions of *CPR1/CPR2* in this tissue, RNA in situ hybridization was used to monitor cellular distribution. No specific zones of high accumulation of *CPR1* mRNAs were detected, but a diffuse barely detectable signal was detected in the whole cotyledon tissues (mostly visible in the spongy parenchyma) with the antisense probe (Fig. 6A), while no apparent signal was revealed with the sense probe (Fig. 6B). By contrast, intense signals were observed with the *CPR2* antisense probe in distinct cotyledon cell types, including IPAP, xylem, and epidermis (Fig. 6C), which were not revealed with the sense probe (Fig. 6D). This complex organization of *CPR2* transcript distribution was further compared with the expression of representative *P450s*

from both MIA and phenylpropanoid pathways that also exhibited compartmented expression, such as *C4H*, *G8H*, and *SLS2*. In young leaves, *CPR2* mRNAs were detected in IPAP, epidermis, and xylem once again (Fig. 7A). Remarkably, this profile corresponds to a superimposition of the expression patterns of each P450 tested, including *C4H*, whose transcripts were detected in epidermis and xylem (Fig. 7B), *G8H* expressed in IPAP, and *SLS2* displaying mRNA accumulation in epidermis (Fig. 7D). Such observations also were confirmed in cotyledons of germinating seedlings (Fig. 7, E–H) and support a preferential involvement of *CPR2* in the reduction of P450s expressed in tissues hosting high levels of MIA and phenylpropanoid biosynthesis. In contrast, the low ubiquitous expression of *CPR1* argues again for a preferential involvement in basal specialized and primary metabolism. These results also suggest that the 79 predicted P450s coexpressed with *CPR2* may be involved in metabolic pathways located to one of these three tissues (i.e. IPAP, xylem, and/or epidermis), whereas no such conclusion may be drawn for the P450s coexpressed with *CPR1*.

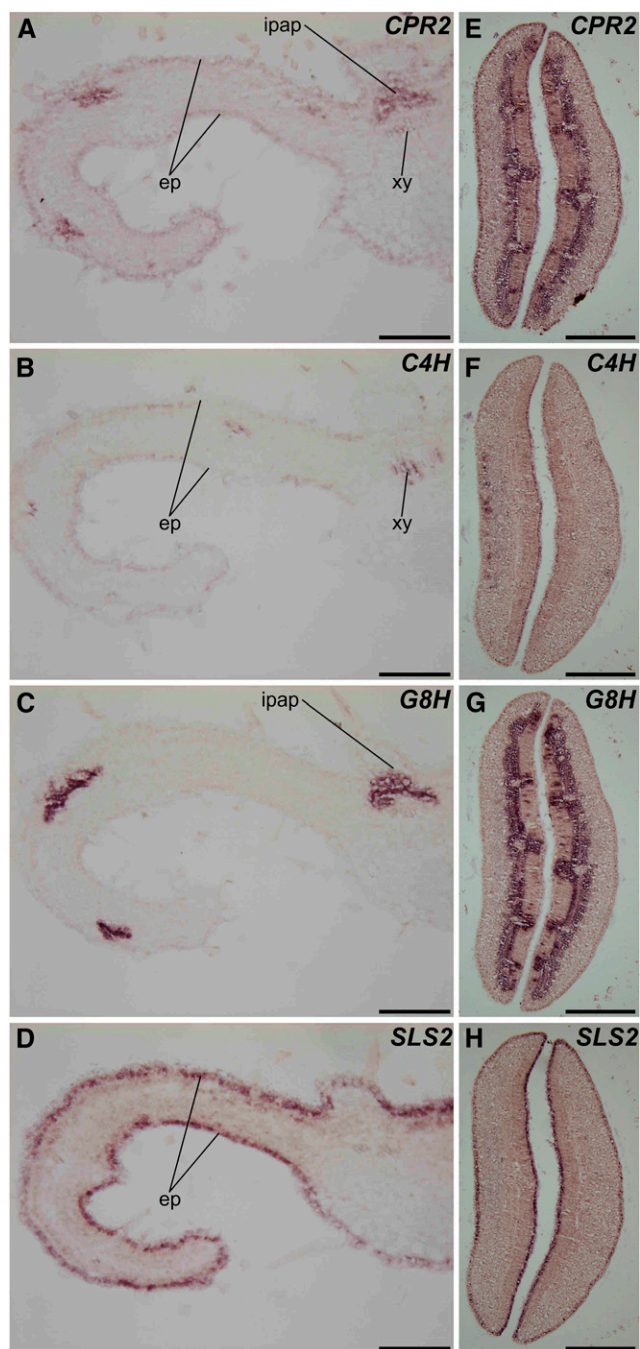
#### Silencing of *CPR1* Does Not Impact the Biosynthesis of MIA

To determine the role of *CPR1* and *CPR2* in the reduction of P450s associated with specialized metabolism in planta, we used virus-induced gene silencing (VIGS) to target each protein. Given the relatively high sequence similarity between *CPR1* and *CPR2* and the presence of highly conserved domains, the identification of silencing sequences that did not result in cross silencing remained tricky. For instance, these silencing sequences were first designed in the 3' untranslated region of each *CPR* but did not lead to substantial transcript down-regulation (data not shown). Therefore, we subsequently used partial coding sequences of *CPR1* or *CPR2*, but it was difficult to avoid some degree of cross silencing, although the most divergent sequence regions were selected. qPCR of mRNA isolated from silenced plants indicated that *CPR1* could be silenced to an expression level of approximately 40% of the empty vector control levels with minimal cross silencing of *CPR2* (Fig. 8A). However, when *CPR2* was silenced (an expression level down to 30% of empty vector control levels), we also observed a decrease in the levels of *CPR1* transcripts (an expression level down to 60% of empty vector control levels; Fig. 8A). Nevertheless, we used these plants to monitor the levels of four monoterpene indole alkaloids that are abundant in leaf, namely catharanthine, vindoline, vindorosine, and serpentine (Fig. 8B). First, we noted that *CPR1* silencing did not result in any quantitative modifications of the MIA content of the silenced plants, strongly suggesting that *CPR1* is not required for MIA biosynthesis and/or that *CPR2* can fully compensate the lack of *CPR1* to reduce P450s associated with the production of MIA. By contrast, in *CPR2* silenced plants, a 45% decrease of the total analyzed MIA content was observed, which



**Figure 6.** Localization of *CPR1* and *CPR2* transcripts in cotyledons of *C. roseus* germinating seedlings. The analysis of *CPR1* and *CPR2* transcript distribution was performed by in situ RNA hybridization. Serial sections of germinating seedlings were hybridized with *CPR1* antisense (AS) probes (A), *CPR2* antisense probes (C), *CPR1* sense (S) probes (B), or *CPR2* sense probes (D) used as negative controls. Ep, Epidermis; X, xylem. Bars = 100  $\mu$ m.





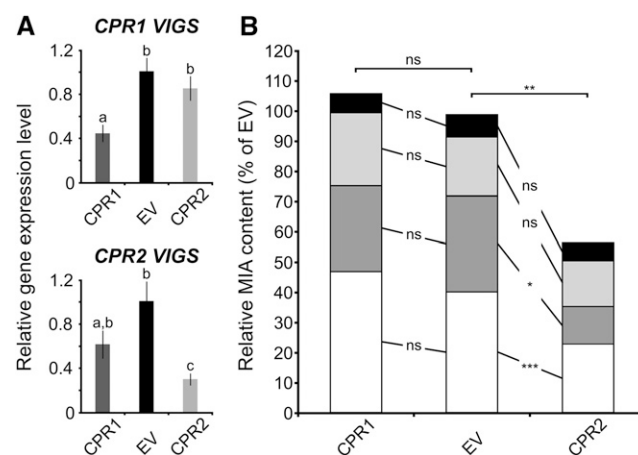
**Figure 7.** *CPR2* is expressed in leaf and cotyledon tissues hosting transcripts of P450s involved in MIA and phenylpropanoid metabolism. *CPR2*, *C4H*, *G8H*, and *SLS2* transcript localizations were carried out by RNA in situ hybridization performed on young leaves (A–D) and cotyledons (E–H). Serial sections were hybridized with *CPR2* antisense probes (A and E), *C4H* antisense probes (B and F), *G8H* antisense probes (C and G), or *SLS1* antisense probes (D and H). ep, Epidermis; xy, xylem. Bars = 100 μm.

mainly resulted from reduction in the levels of catharanthine and vindorosine. This result also was reinforced by the untargeted metabolite analysis, showing that silencing *CPR2* has a much stronger effect on metabolites. In *CPR2* silenced tissue, 37 peaks were altered

significantly (13 up-regulated and 24 down-regulated; *P* value threshold of 0.005), while in *CPR1*, only four were altered (one up-regulated and three down-regulated; Supplemental Table S4). Although the precise identity of these compounds cannot be established from exact mass data, it is possible that some up-regulated metabolites in *CPR2* silenced plants correspond to MIA biosynthetic intermediates. Finally, although we cannot exclude that this MIA decrease is a consequence of both *CPR1* and *CPR2* transcript down-regulation, the more pronounced *CPR2* silencing suggests that *CPR2* impacts MIA biosynthesis to a greater extent than *CPR1*.

## DISCUSSION

While the specific functions of class I and class II CPRs in plants have not been clearly deciphered to date, the Madagascar periwinkle and its complex MIA biosynthesis constitute an attractive model in which to study the relationship between CPRs and specialized metabolism. In contrast to fungi and animals that contain a single CPR gene, most vascular plants possess



**Figure 8.** Silencing of *CPR1* does not impact MIA biosynthesis. A, Down-regulation of *CPR1* and *CPR2* transcript by VIGS. The relative expression of each gene was determined by real-time quantitative reverse transcription-PCR analyses performed on total RNA extracted from *C. roseus* leaves of *CPR1* or *CPR2* silenced plants (dark gray or light gray bars, respectively) or plants transformed with an empty virus control (EV; black bars). CrRPS9 (for RIBOSOMAL PROTEIN S9) was used as a reference gene. Data were normalized to CrRPS9 expression and correspond to average values ( $n = 6$ )  $\pm$  sd of independently transformed plants. Letters indicate statistical classes (Wilcoxon rank-sum test, FDR-adjusted  $P < 0.05$ ). B, Relative MIA contents of *CPR1*- and *CPR2*-VIGS plants expressed relative to that of EV plants. The relative MIA content was determined by measuring peak areas (normalized by sample fresh weight) of catharanthine (white bars), vindorosine (dark gray bars), vindoline (light gray bars), and serpentine (black bars) performed by liquid chromatography-mass spectrometry. The amount of each MIA in silenced plants (eight plants per silencing construct) was expressed relative to that measured in EV plants (eight plants; normalized to 100%). Asterisks denote statistical significance (\*,  $P < 0.005$ ; \*\*,  $P < 0.0005$ ; and \*\*\*,  $P < 0.00005$ ). ns, Not significant.

two functional CPRs, originating from an ancestral gene duplication, along with a more distant CPR without a transmembrane domain (DFR) that does not reduce the P450s tested in this study. Based on their distinct N-terminal sequences, the two functional periwinkle CPRs are assigned to class I and class II CPRs. Their responses to jasmonate suggest that class I and class II play active roles in basal primary metabolism/constitutive specialized metabolism and inducible specialized metabolism, respectively (Jensen and Moller, 2010). In support of this hypothesis, functional biochemical assays and tests of interaction between CPR1 or CPR2 and the P450s G8H, SLS2, T16H1, T16H2, and C4H clearly demonstrated that both CPRs interact with and reduce each P450 with an apparent similar efficiency, suggesting that both types of CPR do not display any structural specificity toward individual P450s (Fig. 3; Table I). However, analysis of gene coexpression patterns with CPR1 and CPR2 sheds light on how CPR2 plays a dedicated role in specialized metabolism, in this case MIA biosynthesis. First, CPR1 expression is correlated with a higher number of genes than CPR2 (9,831 versus 2,525 transcripts), suggesting a broader role of CPR1 in general plant physiology (Fig. 5B). Moreover, the genes coexpressed with CPR1 and CPR2 differed strongly, even when considering lower correlations ( $PCC < 0.3$ ), further supporting the distinct specialization of CPR1 and CPR2 (Fig. 5A). Such a hypothesis is reinforced by the fact that the UniProt keywords associated with genes correlated to CPR1 primarily include basal cellular functions (Supplemental Fig. S10). For example, among the P450s with expression profiles related to CPR1, almost 20% act in primary metabolic pathways such as hormone biosynthesis, while the remaining P450s are mostly associated with phenylpropanoid metabolism (Supplemental Table S3, A and B). In contrast, almost all of the P450s with expression profiles correlating with CPR2 are known to be associated with, or have UniProt keywords associated with, specialized metabolism, especially the MIA metabolism that predominates in *C. roseus* (Supplemental Fig. S10; Supplemental Table S3, A and B). The strong correlation between the expression profiles of CPR2 and MIA biosynthetic genes is reinforced by the identification of nearly all known MIA biosynthetic genes in the gene list associated with CPR2 (Fig. 5C; Supplemental Table S3B). Interestingly, although numerous P450s associated with other specialized metabolic pathways such as sesquiterpene metabolism were correlated to CPR2, only a few involved in phenylpropanoid metabolism were identified. The functional specialization of CPRs in other plant species can be less stringent. For example, this is slightly different from the situation described in *Arabidopsis*, in which previous transcriptomic data analyses showed that both ATR1 and ATR2 were coexpressed with lignin biosynthetic genes during inflorescence stem development, even if, for ATR2, the data set was generated from *Arabidopsis* leaves subjected to cold treatment under high-light conditions or from leaves of lignin biosynthetic gene mutants (Soitamo et al., 2008; Sundin et al., 2014). Taken

altogether, specific gene coexpression with each class of CPR adds another layer of complexity to CPR specialization and the control of specialized metabolism. In periwinkle, the proposed partition of class I and class II CPRs between basal and inducible specialized metabolism cannot be ruled out for phenylpropanoid biosynthesis, but the exclusive association of class II CPRs with subclasses of specialized metabolism such as MIA biosynthesis seems to have been acquired.

The localization pattern of these CPRs also supports a specialized role for CPR2 expression in MIA biosynthesis. Although qPCR demonstrated that CPR1 and CPR2 were both expressed in the same organs, RNA in situ hybridization established cell-specific colocalization of CPR2 transcripts with the tested P450s involved in specialized metabolism, while CPR1 did not show such a correlation (Figs. 4, 6, and 7). The biosynthesis of MIA in *C. roseus* leaves is a highly compartmentalized process involving at least four distinct cell types: IPAP, hosting all the early steps of the monoterpene precursor synthesis up to loganic acid; epidermis that carries out the incorporation of loganic acid into MIA including desacetoxyvindoline; and the specialized cells laticifers and idioblasts, where the last two steps of vindoline formation from desacetoxyvindoline take place. The expression of P450s known to be associated with MIA biosynthesis has been shown to be restricted to IPAP (G8H, IO, and 7DLH) or to epidermis (SLS1–SLS4, T16H1, T16H2, and 16T3O) cell types. In our localization studies, while the CPR1 transcripts were barely detectable in cotyledons and leaves with an apparent general distribution, we observed that CPR2 mRNAs were highly enriched, together with G8H, in IPAP; in epidermis, similar to SLS2 and T16H2; and to a lower extent in xylem, where C4H transcripts from phenylpropanoid biosynthesis also were detected. Thus, the CPR2 expression profile appears to be highly compartmentalized, being superimposed with selected MIA and phenylpropanoid biosynthetic genes. This high CPR2 expression in IPAP suggests that class II CPR is required for G8H activity in this tissue, and by extension to IO and 7DLH, while class I CPR seems to play a housekeeping role, given its uniform distribution and low transcript level. The same rationale could be applied to leaf epidermis, in which CPR2, SLS2, and T16H2 are highly transcribed, reinforcing the specialization of CPR2 in MIA biosynthesis. Data concerning the tissue-specific expression of CPRs in plants are scarce. Therefore, the demonstration of the specific cooccurrence of CPR and P450 transcripts in similar restricted tissues provides a plausible explanation of how CPR specialization is established in planta, based on similar transcriptional regulatory processes. This also may shed light on a predictive tissue-specific expression for the list of P450s coexpressed with CPR2 and on the colocalization of various specialized metabolic pathways in given specialized cell types, as exemplified for epidermis, which harbors MIA, indole, monoterpene-secoiridoid, and phenylpropanoid pathways (Mahroug et al., 2006).

CPR specialization also was supported by VIGS assays conducted in *C. roseus*. While the characterization of CPR mutants was already reported in Arabidopsis (Sundin et al., 2014), gene-silencing approaches have only been applied to CPRs in animal systems (Mackenzie et al., 2010; Tang et al., 2012; Liu et al., 2015). Here, we established that silencing CPR1 in *C. roseus* has no substantial effect on MIA biosynthesis (Fig. 8). This result again suggests that CPR1 does not play a significant role in the reduction of P450s associated with MIA production. Although we cannot exclude a possible compensation by CPR2 in CPR1 silenced plants, this seems unlikely, given their highly specific gene expression profiles (Figs. 6 and 7); moreover, such a complete compensation between ATR1 and ATR2 has not been reported in Arabidopsis systems (Sundin et al., 2014). Due to the high sequence identity, we were not able to specifically down-regulate CPR2; while CPR2 silenced plants had a huge decrease of CPR2 transcripts, this was accompanied by a less marked but substantial decrease (40%) of CPR1 (Fig. 8A). Such transcript down-regulation engendered a significant decrease of the total leaf MIA content (around 45%) in which the most abundant leaf MIAs were most affected (Fig. 8B). Given the absence of such a decrease in CPR1 VIGS lines, it seems likely that this decrease in MIAs is largely the result of CPR2 down-regulation. Interestingly, the lignin content measured in Arabidopsis CPR2 null is altered only slightly (Sundin et al., 2014), suggesting that CPR1 can compensate by providing electrons for the basal lignin metabolism level in Arabidopsis. In contrast, there does not appear to be a compensatory effect for MIA metabolism in *C. roseus*. Notably, in this plant, the biosynthesis of MIA is a constitutive process that also is stimulated by external stimuli or defense-stimulated phytohormones (El-Sayed and Verpoorte, 2007). However, although CPR1 has been proposed to control basal specialized metabolism, it does not have a major impact on MIA biosynthesis in *C. roseus*. Thus, this distinction between the roles of class I and class II CPRs with respect to basal/inducible specialized metabolism may not always hold. Overall, our results suggest that, in *C. roseus*, CPR2 has a distinct function in the specialized metabolism at both the basal and developmentally regulated levels as well as in the response to environmental signals. By contrast, gene expression correlations reinforce the potential involvement of class I CPR with a different set of P450s involved in other metabolic pathways.

## CONCLUSION

In summary, our study gives compelling evidence that the acquisition of a second isoform of CPR (class II CPR) in the plant lineage provides an additional level of control of redox power homeostasis. With respect to the hazard of CPR overexpression (causing reactive oxygen species production; Bassard et al., 2012; Paddon et al., 2013), strict adjustment of cytochrome reducing power

to metabolic demand may take advantage of the transcriptional regulation of two independent genes, each with their specific regulatory control. The current prevailing hypothesis is that class I CPR provides reducing power required for low-level, basal, primary, and secondary metabolism, while class II CPR provides electrons for highly expressed P450s involved in tissue-specific and intensive/induced specialized metabolism. However, as more work is conducted in this area, this hypothesis will continue to be refined. For example, it was demonstrated recently that, in the Apiales lineage, class I CPR genes have been lost and compensated by the acquisition of additional isoforms of class II CPR (Andersen et al., 2016). In this study, we capitalize on the well-studied, complex MIA pathway of *C. roseus* to address the role of CPRs in specialized metabolism. With the indubitable requirement of class II CPR in the biosynthesis of MIA, our work provides new insight into the so-far poorly understood specialization of the CPR classes, suggesting another possible strategy of evolution for redox control.

## MATERIALS AND METHODS

### Chemicals

Secologanin and tabersonine were purchased from Phytoconsult and ChromaDex, respectively. Cinnamic acid, geraniol, and 10-hydroxygeraniol were purchased from Sigma-Aldrich.

### Plant and Cell Culture Growth

Fully expanded *Catharanthus roseus* plants ('Pacifica Pink' and 'Sunstorm Apricot') were used for microscopy fixation (RNA in situ hybridization experiments) and RNA extraction (cloning experiments), respectively. VIGS assays were performed on cv Sunstorm Apricot. The *C. roseus* C20A cell suspension culture used for subcellular localization studies was propagated in Gamborg's B5 medium (Duchefa) at 24°C under continuous shaking (100 rpm) for 7 d as described previously (Guirimand et al., 2009).

### Transcriptomic Resources

Identification of CPR homologous sequences was achieved by interrogating *C. roseus* transcriptomic databases, including the consensus *C. roseus* transcriptome CDF97 (Dugé de Bernonville et al., 2015b; [http://bbv-ea2106.sciences.univ-tours.fr/images/files\\_to\\_download/BAFC94.fasta](http://bbv-ea2106.sciences.univ-tours.fr/images/files_to_download/BAFC94.fasta)), the Medicinal Plant Genomic Resource database (Góngora-Castillo et al., 2012; <http://medicinalplantgenomics.msu.edu>), and the Phytometasyn database (Xiao et al., 2013; <http://www.phytometasyn.ca>).

### RNA Extraction and Reverse Transcription

Total RNA was extracted from young leaves using the NucleoSpin RNA Plant Kit (Macherey-Nagel). First-strand cDNA was synthesized from 0.5 µg of total RNA using oligo(dT)<sub>18</sub> primers (0.5 µM) and 15 units of ThermoScript reverse transcriptase (Invitrogen). Following reverse transcription, complementary RNA was removed by treatment with *Escherichia coli* RNase H (Invitrogen) for 20 min at 37°C.

### Subcellular Localization, Fusion/Deletion Experiments, and Protein Interaction Studies

The subcellular localizations of CPR1 (KJ701028), CPR2 (X69791), and CPR3/DFR (KM111538) were determined by expressing YFP fusion proteins. The full-length open reading frame of each CPR was amplified using specific pairs of

primers (Supplemental Table S5) and cloned into the *SpeI* restriction site of the pSCA-cassette YFPi plasmid in frame with the 5' extremity of the YFP coding sequence, to generate the CPR1-YFP, CPR2-YFP, and CPR3-YFP fusion proteins (Guirimand et al., 2009). The functionalities of the membrane-anchoring domains of CPR1 and CPR2 were studied by fusion/deletion experiments as follows. The coding sequences of the first 53 and 74 residues of CPR1 and CPR2 were amplified using primers helixCPR1-for/helixCPR1-rev and helixCPR2-for/helixCPR2-rev (Supplemental Table S5) and cloned into the *SpeI* restriction site of the pSCA-cassette YFPi plasmid to express the hxCPR1-YFP and hxCPR2-YFP fusion proteins, respectively. To express the CPR1 and CPR2 proteins deprived of their membrane-anchoring domain, the coding sequence of the remaining part of each protein (residues 54–691 for CPR1 and 75–714 for CPR2) was amplified using primers delCPR1/CPRnewrev and delCPR2/CPRoldrev (Supplemental Table S5) and cloned into the *SpeI* restriction site of pSCA-cassette YFPi. The addition of the CPR2 membrane-anchoring domain to CPR3 was achieved through amplification of the coding sequence of the first 74 residues of CPR2 using primers helixCPR-for and helixCPR-rev, harboring *XbaI* and *SpeI* restriction sites at their extremity, respectively. The resulting PCR product was cloned into the *SpeI* site of pSCA-YFP to generate pSCA-helixYFP. This plasmid was linearized subsequently by *SpeI* to allow the introduction of the CPR3 coding sequence, yielding the pSCA-hxCPR3-YFP plasmid. For protein interaction analyses, the same CPR amplification products were cloned into the *SpeI* restriction site of the pSCA-SPYNE 173 plasmid in frame with the 5' end of the sequence encoding YFP<sup>N</sup> to express the CPR1-YFP<sup>N</sup>, CPR2-YFP<sup>N</sup>, and CPR3-YFP<sup>N</sup> proteins (Guirimand et al., 2010). The coding sequences of the five tested P450s (G10H, SLS2, T16H1, T16H2, and C4H) were amplified using appropriate primers (Supplemental Table S5) and cloned into the *SpeI* and/or *BglII* restriction sites of the pSCA-SPYCE(M) plasmid in frame with the 5' end of the sequence encoding YFP<sup>C</sup> to express G10H-YFP<sup>C</sup>, SLS2-YFP<sup>C</sup>, T16H1-YFP<sup>C</sup>, T16H2-YFP<sup>C</sup>, and C4H-YFP<sup>C</sup>. These recombinant plasmids were used for the transient transformation of *C. roseus* cells by particle bombardment and YFP imaging according to Guirimand et al. (2009, 2010) and Foureau et al. (2016). Briefly, plated *C. roseus* cells were bombarded with DNA-coated gold particles (1  $\mu\text{m}$ ) and a 1,100-p.s.i. rupture disc at a stopping-screen-to-target distance of 6 cm, using the Bio-Rad PDS1000/He system. Cells were cultivated for 16 to 38 h before being harvested and observed. The subcellular localization was determined using an Olympus BX-51 epifluorescence microscope equipped with an Olympus DP-71 digital camera and a combination of YFP and CFP filters. The pattern of localization presented in this work is representative of approximately 50 observed cells. The ER or nucleocytoplasmic localization of the different fusion proteins was confirmed by cotransformation experiments using an ER-CFP marker (CD3-954; Nelson et al., 2007) and a nucleocytoplasmic CFP marker (Guirimand et al., 2011a). Such plasmid cotransformations were performed using 400 ng of each plasmid or 100 ng for BiFC assays.

## Tissue Fixation, Embedding in Paraffin, and Sectioning

RNase-free conditions were strictly observed for all steps. All glassware was baked for 8 h at 180°C, and nondisposable plastic wares were incubated for 10 min in an aqueous 3% hydrogen peroxide solution before washing in diethyl pyrocarbonate-treated water. Leaves from mature *C. roseus* plants grown in a greenhouse and young germinating seedlings were rapidly fixed in formalin (10%), acetic acid (5%), and ethanol (50%) and embedded in Paraplast (Dominique Dutscher) as described previously (Mahroug et al., 2006; Guirimand et al., 2011b). Serial sections (10  $\mu\text{m}$ ) were spread on aminopropyltriethoxysilane-coated slides overnight at 40°C, and paraffin was removed using xylene (twice for 15 min) before rehydration in an ethanol gradient series up to diethyl pyrocarbonate-treated water.

## In Situ RNA Hybridization of *C. roseus* Leaves and Cotyledons

Full-length CPR1 and CPR2 cDNAs cloned into pGEMT-easy vector (Promega) were used for the synthesis of sense and antisense RNA probes as described previously (Mahroug et al., 2006). For G8H, SLS, and C4H, previously described plasmids were used for the riboprobe in vitro transcription (Irmeler et al., 2000; Burlat et al., 2004; Mahroug et al., 2006). Paraffin-embedded serial longitudinal sections of young leaves and cross sections of emerging cotyledons were hybridized with digoxigenin-labeled transcripts and localized with anti-digoxigenin-alkaline phosphatase conjugates according to Mahroug et al. (2006).

## Heterologous Expression of *C. roseus* CPRs and P450s in *Saccharomyces cerevisiae*

Full-length CPR1, CPR2, and CPR3 cDNAs were amplified using the specific yeast expression primers described in Supplemental Table S5. Each pair of primers includes appropriate restriction sites at their extremities to allow cloning of the resulting PCR products into the *SpeI* site of pESC-Leu yeast expression plasmid harboring the LEU2 nutritional marker (Agilent Technologies). Addition of the CPR2 membrane-anchoring domain to CPR3 was achieved as described in the previous section, except that cloning was performed into the pESC-Leu plasmid. G8H, SLS2, T16H1, T16H2, and C4H coding sequences were amplified using primers including *BglII* restriction at their extremities to allow cloning into the *BamHI* site of the pYEDP60 plasmid harboring URA3 and ADE2 nutritional markers (Pompon et al., 1996). pESC-Leu CPRs, pYEDP60 P450 recombinant vectors, and/or empty plasmids were associated by pair and used to transform either the *Saccharomyces cerevisiae* strain WT303 (containing the endogenous yeast CPR) or the WAT11-expressing ATR1 (Pompon et al., 1996). Leu+, Ura+/Ade+, or Ura+/Ade+/Leu+ transformants were selected onto solid complete synthetic medium plates containing 0.67% yeast nitrogen base, 2% agar, 2% dextrose, and 0.05% DOB-Leu, DOB-Ura-Ade, or DOB-Ura-Ade-Leu as required. Yeast transformants were grown in 10 mL of appropriate liquid complete synthetic medium until reaching the stationary phase of culture and then harvested by centrifugation. Protein expression was induced by cultivating the harvested yeast in 50 mL of liquid YPGal medium (1% bactopectone, 1% yeast extract, and 2% Gal) for 6 h as described by Besseau et al. (2013).

## Enzyme Assays

Following the induction of protein expression, 50 mL of yeast culture was harvested by centrifugation and resuspended in 2 mL of buffer R (50 mM Tris-HCl, pH 7.5, and 1 mM EDTA) in a 50-mL centrifugation tube. An equal volume of glass beads was added (425–600  $\mu\text{m}$ ; Sigma-Aldrich), and cells were broken by vigorous shaking. For this purpose, tubes were shaken by hand during 30 s in a cold room (4°C) before being put on ice for an additional 30 s. This operation was repeated 10 times before the addition of 2 volumes of buffer R, allowing the recovery of the yeast crude extracts prior to protein quantification using the Bio-Rad protein microassay. P450 activities were analyzed in a final volume of 100  $\mu\text{L}$  containing 300  $\mu\text{g}$  of protein, 100  $\mu\text{M}$  NADPH(H<sup>+</sup>), and 20  $\mu\text{M}$  of loganin for SLS2, tabersonine for T16H1 and T16H2, or cinnamate for C4H. Reactions were initiated by the addition of NADPH(H<sup>+</sup>), incubated at 30°C for 0, 5, 15, and 30 min (T16H1, T16H2, and C4H) or for 10, 30, 60, or 120 min (SLS2), and quenched by the addition of 100  $\mu\text{L}$  of methanol prior to ultra-performance liquid chromatography-mass spectrometry analysis. G10H activity was tested using microsomal membranes purified according to Heitz et al. (2012). Enzymatic assays were conducted according to Höfer et al. (2013) in a final volume of 100  $\mu\text{L}$  of 20 mM citrate-phosphate buffer, pH 7.4, containing 400  $\mu\text{M}$  NADPH, 200  $\mu\text{M}$  geraniol, and normalized amounts of microsomes. After 60 min of reaction, products were extracted with 500  $\mu\text{L}$  of ethyl acetate. The ethyl acetate phase was collected with a glass pipette and dried under a nitrogen gas flow, and products were analyzed by gas chromatography-flame ionization detector. For all P450 tests performed in the WT303 yeast strain including CPR1, CPR2, or CPR3, activities resulting from the yeast endogenous CPR were subtracted from total activity to estimate the activity resulting from each periwinkle CPR. All results were expressed as substrate conversion rate for 1 h.

## Ultra-Performance Liquid Chromatography-Mass Spectrometry Analyses

All samples were centrifuged, and the supernatants were stored at 4°C prior to injection. The ultra-performance liquid chromatography system consisted of an ACQUITY UPLC device (Waters). Separation was performed using a Waters Acquity HSS T3 C18 column (150 mm  $\times$  2.1 mm, 1.8  $\mu\text{m}$ ) with a flow rate of 0.4 mL min<sup>-1</sup> at 55°C. The injection volume was 5  $\mu\text{L}$ . The mobile phase consisted of solvent A (0.1% formic acid in water) and solvent B (0.1% formic acid in acetonitrile). Chromatographic separation was achieved using an 8-min linear gradient from 10% to 24% solvent B. Mass spectrometry detection was performed using an SQD mass spectrometer equipped with an electrospray ionization source controlled by Masslynx 4.1 software (Waters). The capillary and sample cone voltages were 3,000 and 30 V, respectively. The cone and desolvation gas flow rates were 60 and 800 L h<sup>-1</sup>. Data collection was carried out in negative mode for secologanin ([M + HCOOH-H]<sup>-</sup> = 433, retention time =

7.12 min), cinnamate ( $[M - H]^- = 147$ ), and 4-hydroxycinnamate ( $[M - H]^- = 163$ ) and in positive mode for loganin ( $[M + Na]^+ = 413$ , retention time = 5.61 min), tabersonine ( $[M + H]^+ = 337$ , retention time = 4.72 min), and 16-hydroxytabersonine ( $[M + H]^+ = 353$ , retention time = 3.87 min).

## Gas Chromatography Analyses

Monoterpenes were analyzed by gas chromatography-flame ionization detection (Alpha-MOS). Samples were injected in the split mode ( $50 \text{ mL min}^{-1}$ ), and compounds were separated on a BPX5 capillary column. The injector was heated at  $250^\circ\text{C}$ , and the oven was set at  $45^\circ\text{C}$  for 60 s. The temperature was next increased to  $250^\circ\text{C}$  at a rate of  $8^\circ\text{C min}^{-1}$ , followed by an increase to  $320^\circ\text{C}$  at a rate of  $30^\circ\text{C min}^{-1}$ . The oven was maintained at  $320^\circ\text{C}$  for 1 min before the end of analysis. The flame ionization detector was set at  $280^\circ\text{C}$ . Peaks were identified by comparing the retention times of authentic standards.

## Gene Expression Correlation Analyses

The abundance of transcripts in the CDF97 assembly was estimated by pseudoaligning reads with Salmon (Patro et al., 2015) in the variational bayesian optimized (-vbo) quasi-mapping mode with bias correction (-biasCorrect). Expression was  $\log_2$  transformed to distribute TPM normally (Supplemental Fig. S10). A total of 16 paired-end samples (SRR1144633, SRR1144634, SRR1271857, SRR1271858, SRR1271859, SRR342017, SRR342019, SRR342022, SRR342023, SRR646572, SRR646596, SRR646604, SRR648705, SRR648707, SRR924147, and SRR924148) and 23 SE samples (SRR122239, SRR122240, SRR122241, SRR122242, SRR122243, SRR122244, SRR122245, SRR122246, SRR122247, SRR122248, SRR122249, SRR122250, SRR122251, SRR122252, SRR122253, SRR122254, SRR122255, SRR122256, SRR122257, SRR122258, SRR122259, SRR122260, and SRR122261) were used as described by Dugé de Bernonville et al. (2015b). PCCs and their significance were calculated with R (R Development Core Team, 2015) for each CPR with each transcript predicted in the CDF97 consensus assembly. Annotation of transcripts was performed by following the Trinotate pipeline (<https://trinotate.github.io/>). GO terms and UniProt keywords were attributed according to the UniProt database. Gene set enrichment analyses were made by comparing the observed representation of GO terms with a theoretical hypergeometric distribution (phyper function in R). A promoter analysis was performed with the MEME suite (version 4.11; Bailey et al., 2009). R scripts were designed to retrieve promoter sequences of each predicted transcript in the *C. roseus* genome sequence (Kellner et al., 2015b). Promoter sequences of genes putatively associated with each CPR (PCC > 0.4 and FDR-adjusted PCC  $P < 0.05$ ) were analyzed with DREME, and the resulting motifs were analyzed for GO term association with GOMO. The most promising motifs were next inspected manually in promoter sequences of each CPR.

## Gene Expression Analyses

CPR1, CPR2, and CPR3 expression was first determined by comparing transcript abundance in *C. roseus* transcriptomic data sets (Góngora-Castillo et al., 2012; Dugé de Bernonville et al., 2015b) and subsequently measured by real-time reverse transcription-qPCR using the primers listed in Supplemental Table S6. The corresponding PCR products were cloned in pGEMT-easy (Promega) and sequenced to ensure the specificity of amplification. Primer efficacy was evaluated on plasmids containing the appropriate cDNA. Distinct *C. roseus* organs (such as roots, stems, young and mature leaves, and petals; cv Apricot Sunstorm) were immediately frozen in liquid nitrogen after sampling. Samples (50 mg) were ground with a mortar and pestle in liquid nitrogen, and total RNA was extracted with Nucleospin RNA (Macherey-Nagel), quantified with a Nanodrop spectrophotometer (Thermo Fisher), and treated ( $1.5 \mu\text{g}$ ) with RQ1 RNase-free DNase (Promega) before being used for first-strand cDNA synthesis by priming with  $0.5 \mu\text{M}$  oligo(dT)<sub>18</sub>. Retrotranscription of  $1.5 \mu\text{g}$  of total RNA was carried out using the SuperScript III reverse transcriptase kit (Invitrogen) at  $50^\circ\text{C}$  for 1 h according to the manufacturer's instructions. Real-time PCR was run on the CFX96 Touch Real-Time PCR System (Bio-Rad) using SYBR Green I technology. Each reaction was performed in a total reaction volume of  $25 \mu\text{L}$  containing an equal amount of cDNAs (1:3 dilution),  $0.05 \mu\text{M}$  forward and reverse primers, and the  $1 \times$  DyNAmo ColorFlash Probe qPCR Kit (Thermo Fisher Scientific). The amplification program was  $95^\circ\text{C}$  for 7 min (polymerase heat activation) followed by 40 cycles containing two steps,  $95^\circ\text{C}$  for 10 s and  $60^\circ\text{C}$  for 40 s. At the end of the amplification, a melt curve was performed to check amplification specificity. Relative quantification of

transcripts was performed with calibration curves and normalization with the *C. roseus* 40S RPS9 (AJ749993.1) reference gene. All amplifications were performed in triplicate with at least two independent biological repeats.

## VIGS

CPR1 and CPR2 silencing fragments were amplified using the primers described in Supplemental Table S5 and cloned into the pTRV2u vector described by Geu-Flores et al. (2012). The resulting plasmids and the empty vector were used to perform the VIGS assays on *C. roseus* seedlings as described by Liscombe and O'Connor (2011) or Carqueijeiro et al. (2015). Leaves from the first two leaf pairs to emerge following inoculation were harvested from eight plants transformed with each construct and subjected to gene expression analysis by real-time reverse transcription-PCR (primers are given in Supplemental Table S6). The alkaloid content of silenced leaves was determined by liquid chromatography-mass spectrometry as described previously (Liscombe and O'Connor, 2011; Geu-Flores et al., 2012). A detailed version of the silencing procedure is given in Supplemental Methods S1.

## Accession Numbers

Sequence data of CPR1, CPR2 and CPR3/DRF from this article can be found in the GenBank/EMBL data libraries under accession numbers KJ701028, X69791, and KM111538, respectively.

## Supplemental Data

The following supplemental materials are available.

**Supplemental Figure S1.** Alignment of periwinkle CPR1, CPR2, and CPR3 deduced amino acid sequences.

**Supplemental Figure S2.** CPR phylogenetic tree.

**Supplemental Figure S3.** Gene organization of CPR1, CPR2, and CPR3/DRF.

**Supplemental Figure S4.** Alignment of CPR1 and CPR2 amino acid sequences, highlighting characteristic membrane anchor, FMN, FAD, NADPH, and P450-binding domains.

**Supplemental Figure S5.** Alignment of periwinkle CPR1, CPR2, and CPR3/DRF deduced amino acid sequences and determination of the putative functional domains of CPR3 according to Varadarajan et al. (2010).

**Supplemental Figure S6.** Detection of a putative transmembrane helix at the N-terminal end of CPR1, CPR2, and CPR3/DRF.

**Supplemental Figure S7.** Characterization of the membrane-anchoring domain of CPR1 and CPR2.

**Supplemental Figure S8.** Addition of the CPR2 membrane-anchoring domain to CPR3 enables ER anchoring.

**Supplemental Figure S9.** Transcript distribution of CPR1, CPR2, C4H, G8H, SLS2, T16H1, and T16H2 in various *C. roseus* organs.

**Supplemental Figure S10.** Distribution of TPM.

**Supplemental Figure S11.** Functional classification of the nearest coexpressed genes with *C. roseus* CPR.

**Supplemental Table S1.** Identification of contigs potentially encoding CPR candidates in the Medicinal Plant Genomic Resource database, the Phytometasyn database, and the *C. roseus* consensus transcriptome.

**Supplemental Table S2.** Evaluation of the efficiency of P450 reduction by CPR1, CPR2, CPR3/DRF, hxCPR3/DRF, the yeast endogenous CPR (WT303 yeast strain), and the codon-optimized ATR1 of the WAT11 yeast strain.

**Supplemental Table S3A.** P450 genes coexpressed with CPR1, CPR2, or DRF.

**Supplemental Table S3B.** Whole list of genes coexpressed with each CPR.

**Supplemental Table S3C.** Promoter analysis of coexpressed genes with each CPR.

- Supplemental Table S4.** Untargeted metabolite analysis in silenced plants.
- Supplemental Table S5.** Primers used for cDNA cloning.
- Supplemental Table S6.** Primers used for qPCR analyses.
- Supplemental Methods S1.** Detailed VIGS procedure.

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