RNA Interference-Mediated Repression of MtCCD1 in Mycorrhizal Roots of Medicago truncatula Causes Accumulation of C27 Apocarotenoids, Shedding Light on the Functional Role of CCD1\[W][OA]

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Tailoring carotenoids by plant carotenoid cleavage dioxygenases (CCDs) generates various bioactive apocarotenoids. Recombinant CCD1 has been shown to catalyze symmetrical cleavage of C40 carotenoid substrates at 9,10 and 9',10' positions. The actual substrate(s) of the enzyme in planta, however, is still unknown. In this study, we have carried out RNA interference (RNAi)-mediated repression of a Medicago truncatula CCD1 gene in hairy roots colonized by the arbuscular mycorrhizal (AM) fungus Glomus intraradices. As a consequence, the normal AM-mediated accumulation of apocarotenoids (C13 cyclohexenone and C14 mycorradicin derivatives) was differentially modified. Mycorradicin derivatives were strongly reduced to 3% to 6% of literature data implicate C27 apocarotenoid cleavage as the general functional role of CCD1 in planta. A revised scheme of plant carotenoid cleavage in two consecutive steps is proposed, in which CCD1 catalyzes only the second step in the cytosol (C27 → C14 + C13), while the first step (C40 → C27 + C13) may be catalyzed by CCD7 and/or CCD4 inside plastids.

Carotenoids are a large class of isoprenoid pigments, which are produced by plants and certain microbes. Conjugated double bonds in a C40 tetraterpene skeleton form the basis of their chromophores, which are extended and modified by cyclization, isomerization, and oxidation. Carotenoids fulfill important roles as accessory photosynthetic pigments harvesting light and preventing photooxidative damage, as general antioxidants, and as mediators of membrane fluidity. They also serve as yellow, orange, and red pigments in many flowers and fruits. In plants, biosynthesis of C40 carotenoids is compartmentalized to plastids, comprising not only chloroplasts but also chromoplasts and other plastid types (Hirschberg, 2001). A plastidial location also applies for the early steps of biosynthesis carried out by enzymes of the methylerythritol phosphate (MEP) pathway, generating C5 isopentenyl diphosphate precursors (Rodríguez-Concepcion and Boronat, 2002).

Carotenoid catabolism is not simply responsible for balancing steady-state carotenoid levels but forms many new bioactive compounds. While unspecific chemical oxidative cleavage is possible, the vast majority of known cleavage products (apocarotenoids) are thought to originate from defined cleavage (“tailoring”) of carotenoids by carotenoid cleavage dioxygenase (CCD) enzymes (Giuliano et al., 2003; Bouvier et al., 2005). In addition, many signaling molecules comprising abscisic acid (ABA), strigolactones, and a shoot-branching inhibitor with unknown structure are derived from cleavage products of carotenoids (Akiyama et al., 2005; Auldridge et al., 2006b; Kloer and Schulz, 2006).

1 This work was supported in part by the Deutsche Forschungsgemeinschaft (Bonn) in the priority program SPP 1084 (MolMyk).
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www.plantphysiol.org/cgi/doi/10.1104/pp.108.125062

Plant Physiology, November 2008, Vol. 148, pp. 1267–1282, www.plantphysiol.org © 2008 American Society of Plant Biologists 1267
Molecular identification has by now defined nine different clades of plant CCDs, which are named after their representatives in Arabidopsis (*Arabidopsis thaliana*). AtCCD1, AtCCD4, AtCCD7, and AtCCD8 cleave a variety of trans-carotenoid substrates at specific double bonds, but only CCD1 and CCD7 have confirmed cleavage specificities. Recombinant AtCCD1 cleaves symmetrically at 9,10 (9',10') positions to yield two C_{13} ketones and one C_{14} dialdehyde from one C_{40} parent carotenoid molecule (Schwartz et al., 2001; Fig. 1). Recently, additional in vitro activity of CCD1 on 5,6 (5',6') double bonds of lycopene has been reported (Vogel et al., 2008). Recombinant AtCCD7 exhibits regioselectivity for the 9,10 position similar to AtCCD1, yet it cleaves only once asymmetrically, resulting in C_{13} and C_{27} products (Schwartz et al., 2004). AtCCD8 seems to further cleave this C_{27} product to C_{18} and C_{9}, which are the assumed precursors of a still elusive shoot-branching inhibitor (Schwartz et al., 2004; Alder et al., 2008). While these four CCD classes are structurally only distantly related to each other, the remaining five CCDs fall into one class of specialized CCDs. These are called NCEDs, a term derived from their specificity for 9-cis-epoxycarotenoid substrates. Arabidopsis NCED2, NCED3, NCED5, NCED6, and NCED9 are all related to VP14, the prototype NCED/CCD characterized first from the maize (*Zea mays*) mutant *vp14*, which is impaired in the earliest step of ABA biosynthesis from carotenoids (Schwartz et al., 1997; Tan et al., 2003). All NCEDs cleave the 11,12 double bond of 9-cis isomers of neoxanthin or violaxanthin to yield the ABA precursor xanthoxin. Import studies, immunolocalization experiments, and transit peptide prediction algorithms suggest a plastidial location for all NCEDs and CCDs except CCD1, which has been shown in several plant systems to act in the cytosol (Bouvier et al., 2003; Tan et al., 2003; Simkin et al., 2004a; Atulridge et al., 2006a).

Root colonization by arbuscular mycorrhizal (AM) fungi frequently leads to the accumulation of large amounts of apocarotenoids of a cyclic C_{13} type and a linear C_{14} type (Klingner et al., 1995; Maier et al., 1995; Walter et al., 2000; Strack and Fester, 2006; Fig. 1). This phenomenon occurs preferentially during late stages of the AM symbiotic plant-microbe interaction and is often accompanied by the appearance of a yellowish root coloration. The chromophore responsible for this coloration is mycorradicin, a C_{14} polyenic dicarboxylic acid. Derivatives of mycorradicin accumulate concomitantly with structural variants of C_{13} cyclohexenones, the latter being glycosylated and deposited in vacuoles. Both of these AM-induced apocarotenoid types were predicted to be generated by the action of a CCD1-type enzyme on a common C_{40} carotenoid precursor (Walter et al., 2000, 2007; Hans et al., 2004; Strack and Fester, 2006; Akiyama, 2007; Fig. 1). Attempts to isolate pathway intermediates and particularly the proposed C_{40} carotenoid precursor have so far only led to the detection of a minor increase in phytoene upon application of the carotenoid biosynthesis inhibitor norfluorazon (Fester et al., 2005). This indicates that carotenoid cleavage is unlikely to be a rate-limiting step in the biosynthesis of these apocarotenoids in mycorrhizal roots and that the carotenoids formed during the symbiosis are probably instantly cleaved. A different class of plant apocarotenoids (strigolactones) occurring in low quantities plays a role in early signaling steps of the AM symbiosis (Akiyama et al., 2005; Matusova et al., 2005), but whether there is any relationship to the biogenesis of abundant C_{13}- and C_{14}-type apocarotenoids is currently unknown.

![Figure 1](image_url)
Previous molecular studies on the accumulation of C13 and C14 apocarotenoids in mycorrhizal roots have focused on the first two steps of the MEP pathway. Both 1-deoxy-d-xylulose 5-phosphate synthase (DXS) and 1-deoxy-d-xylulose 5-phosphate reductoisomerase genes were shown to be transcriptionally up-regulated in cereal roots upon mycorrhization, emphasizing the importance of precursor supply activation (Walter et al., 2000; Hans et al., 2004). Further investigations revealed the widespread occurrence of two differentially regulated DXS isogenes, of which only DXS2 was up-regulated in mycorrhizal roots (Walter et al., 2002). Recent studies have shown local promoter activity of a DXS2 gene in cells hosting fungal arbuscules (Floss et al., 2008), in accordance with earlier findings of local apocarotenoid deposition in arbusculated cells (Fester et al., 2002a).

Arbuscules are highly branched fungal organs outgrowing from hyphae within host cortical root cells, which constitute the predominant site of nutrient exchange between the plant and the fungal symbiont. Their proper function can determine the development of symbiotic structures (Javot et al., 2007). Interestingly, strong silencing of DXS2 expression in hairy roots of Medicago truncatula leads to reduced levels of both types of apocarotenoids (cyclohexenone and mycorradicin derivatives), accompanied by an abolishment of the normal expression of many arbuscule-specific plant marker genes (Floss et al., 2008). However, whether this impairment of symbiotic functions is indeed linked to the lack of either of these mycorrhizal apocarotenoids is still unresolved, since effects from the absence of other DXS2-dependent isoprenoids cannot be excluded.

This still inconclusive result led us to target the later steps in the pathway, namely carotenoid cleavage, for a new RNA interference (RNAi) approach to elucidate the function of C13 cyclohexenone and/or C14 mycorradicin apocarotenoids in mycorrhizal roots. Both 1-deoxy-d-xylulose 5-phosphate synthase (DXS) and 1-deoxy-d-xylulose 5-phosphate reductoisomerase genes were shown to be transcriptionally up-regulated in cereal roots upon mycorrhization, emphasizing the importance of precursor supply activation (Walter et al., 2000; Hans et al., 2004). Further investigations revealed the widespread occurrence of two differentially regulated DXS isogenes, of which only DXS2 was up-regulated in mycorrhizal roots (Walter et al., 2002). Recent studies have shown local promoter activity of a DXS2 gene in cells hosting fungal arbuscules (Floss et al., 2008), in accordance with earlier findings of local apocarotenoid deposition in arbusculated cells (Fester et al., 2002a).

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This still inconclusive result led us to target the later steps in the pathway, namely carotenoid cleavage, for a new RNA interference (RNAi) approach to elucidate the function of C13 cyclohexenone and/or C14 mycorradicin apocarotenoids in the AM symbiosis. CCD1 genes have been characterized from several plants, all showing that in Escherichia coli recombinant CCD1 preferentially cleaves a variety of C40 carotenoids with cyclic and noncyclic end groups symmetrically at the 9,10 (9',10') double bonds (Schwartz et al., 2001; Mathieu et al., 2005; Ibdah et al., 2006; Fig. 1). However, it has not been proven whether these biochemical cleavage properties in vitro also represent the normal biological role of CCD1 enzymes. In fact, little is known about the actual substrates of native CCD1 in compartmentalized plant cells containing plastids. Previous knockdown approaches on CCD1 expression in transgenic plants have resulted in strong suppression of CCD1 transcript levels but to only limited effects on C40 carotenoid levels, prompting speculations about additional players (Simkin et al., 2004a, 2004b; Auldridge et al., 2006a). In this study, we have silenced the expression of a MtCCD1 gene in mycorrhizal hairy roots of M. truncatula. Surprisingly, the accumulation of the two hitherto known mycorrhiza-induced apocarotenoid types was reduced to different extents and novel C27 apocarotenoid derivatives appeared as a consequence of MtCCD1 repression. Together, these and other observations lead to a new view of carotenoid cleavage in planta, reconciling previous contradictions regarding the presumed substrate and the CCD1 enzyme being in different cellular compartments.

RESULTS
Cloning of a MtCCD1 cDNA and Proof of Functionality of the Protein Encoded

A full-length MtCCD1 cDNA was isolated from a M. truncatula mycorrhizal root cDNA library using an 825-bp MtCCD1 fragment as a probe. DNA sequencing of several clones revealed complete identity of the cDNA (accession no. FM204879) to TC100912 of the Dana-Farber Cancer Institute (DFCI) Medicago Gene index (http://compbio.dfci.harvard.edu/tgi/cgi-bin/ tgi/gimain.pl?gudb=medicago). The deduced MtCCD1 protein consists of 540 amino acids and has a calculated molecular mass of 60.9 kD. The MtCCD1 amino acid sequence exhibits strong similarity to many other CCD1 proteins, exemplified by 87.2% identity (91.1% similarity) to Phaseolus vulgaris CCD1, 80.2% identity (86.8% similarity) to Arabidopsis CCD1, and 76.5% identity (82.8% similarity) to maize CCD1 (Supplemental Fig. S1). This indicates that MtCCD1 belongs to the well-studied class of CCD1 proteins and genes (Schwartz et al., 2001; Simkin et al., 2004a, 2004b; Mathieu et al., 2005; Auldridge et al., 2006a; Ibdah et al., 2006; Kato et al., 2006; Sun et al., 2008; Vogel et al., 2008; Supplemental Fig. S1). Application of prediction algorithms (http://www.cbs.dtu.dk/services/) suggests a subcellular localization of the MtCCD1 protein in the cytosol, in accordance with reports on other CCD1 proteins (data not shown). Searching EST databases of M. truncatula revealed 29 ESTs of the MtCCD1 type from all kinds of tissues, including roots and mycorrhizal roots. Four types of MtCCD1-related ESTs of 83% to 86% deduced amino acid sequence identity to MtCCD1 transcribed from putative MtCCD1 paralogs were much less abundant (one to three ESTs each) and originated from pod or shoot libraries but not from roots.

To prove the functionality of the protein encoded in the MtCCD1 cDNA, two constructs with or without a His tag were expressed in E. coli cells engineered to produce β-carotene (pAC-BETA) or zeaxanthin (pAC-ZEAX; Cunningham et al., 1996; Sun et al., 1996). Using both constructs, a MtCCD1 protein band of the calculated size could be observed after induction by isopropyl β-D-thiogalactopyranoside and separation on SDS gels. The expression of both constructs led to the expected decoloration of the two carotenogenic E. coli strains, which was absent in controls expressing the empty vector (EV) pET28a (Fig. 2). This demonstrates carotenoid cleavage activity of the recombinant protein encoded in the MtCCD1 cDNA.
Up-Regulation of MtCCD1 Transcript Levels in Mycorrhizal Roots of M. truncatula

Previous analyses have indicated a minor to moderate up-regulation of CCD1 transcripts in roots upon colonization by mycorrhizal fungi for both M. truncatula and maize (Lohse et al., 2005; Sun et al., 2008). To confirm these results for this study, we conducted an AM fungal colonization experiment on wild-type roots of M. truncatula for 7 and 9 weeks and for several biological replicates. An early gene of apocarotenoid biosynthesis, known to be AM inducible (MtDXS2; Walter et al., 2002; Floss et al., 2008), and a molecular marker of mycorrhizal colonization (the AM-specific expression of the phosphate transporter gene MtPT4; Harrison et al., 2002) were included in the analysis. Although MtCCD1 transcript levels were not elevated as strongly as those of either MtDXS2-1 or MtPT4, they exhibited a slight increase after 7 weeks and a still minor but significant elevation after 9 weeks (3-fold) in mycorrhizal roots compared with nonmycorrhizal control roots (Table I). As discussed previously for ZmCCD1, local induction of MtCCD1 transcript levels in arbusculated cells, the site of mycorrhizal apocarotenoid biosynthesis, is likely to be considerably higher than the induction factor calculated from extracts (Sun et al., 2008).

Knockdown of MtCCD1 Expression Leads to Strong Reduction of Mycorradicin Derivatives But to Only Moderate Reduction of Cyclohexenone Derivatives

To reduce the levels of MtCCD1 transcripts and thereby generate a loss-of-function mutant of MtCCD1, an RNAi construct of 335 bp covering the 3' part of the coding region and 3' untranslated sequences was generated. After transfer into the binary vector pRedRoot, this construct was introduced and expressed in transgenic hairy roots of M. truncatula. Concomitantly, plants expressing EV sequences were cultivated as controls. After injection of Agrobacterium rhizogenes transgene carriers into hypocotyls, further cultivation of plants involved the selection of transgenic roots outgrowing from the injection site and the removal of residual wild-type as well as untransformed hairy roots to obtain individual, independently transformed composite plants, each with individual transgenic hairy roots and a wild-type shoot.

Figure 3 summarizes the results of two experiments comparing mycorrhizal RNAi root samples with mycorrhizal EV control roots after 7 or 9 weeks of colonization. While the average levels of EV controls differed somewhat between the experiments, a strong reduction was observed in MtCCD1 transcript levels in RNAi roots compared with EV control roots, resulting in about 8% residual transcripts in RNAi roots (Fig. 3A). Next, we determined the levels of both C14 mycorradicin and C13 cyclohexenone derivatives in these roots. Mycorradicin derivative concentrations were strikingly lower in RNAi roots (5 nmol g−1 fresh weight [7 weeks of colonization] and 3 nmol g−1 fresh weight [9 weeks of colonization]), corresponding to 6% and 3% residual levels of the amounts of EV controls (Fig. 3B). These reductions were thus in the same order of magnitude as the reduction in MtCCD1 transcript levels, suggesting a link between MtCCD1 activity and the accumulation of mycorradicin derivatives. However, C13 cyclohexenone derivative levels were not

| Time | Gene    | EΔCt × 10−3 | Nonmycorrhizal | Mycorrhizal |
|------|---------|-------------|----------------|-------------|
| 7 weeks | MtCCD1  | 28.0 ± 7.8  | 41.7 ± 14.6    |             |
|       | MtDXS2-1| 7.6 ± 3.1   | 16.3 ± 4.0     |             |
|       | MtPT4   | 0.02 ± 0.005| 273.3 ± 94.5   |             |
| 9 weeks | MtCCD1  | 20.7 ± 6.8  | 62.3 ± 21.5    |             |
|       | MtDXS2-1| 4.4 ± 1.7   | 32.3 ± 5.1     |             |
|       | MtPT4   | 0.03 ± 0.01 | 825.3 ± 386.9  |             |
reduced to a comparable extent. They exhibited a significant reduction in both experiments, but the residual levels in RNAi roots compared with EV control roots were still between 30% and 47% on average (Fig. 3C).

A Yellow-Orange Root Coloration Appears in MtCCD1-Repressed Roots

In addition to the biochemical changes in the levels of the C\textsubscript{13} and C\textsubscript{14} apocarotenoids, the roots of the mycorrhizal RNAi plants exhibited a striking phenotypic change in coloration (Fig. 4). The EV control roots also exhibit a coloration, related to mycorraticin accumulation, but this orange-brown color is typically rather faint. It is similarly observed with mycorrhizal wild-type roots of M. truncatula (Walter et al., 2007). However, the RNAi roots silenced for MtCCD1 expression exhibited a conspicuous change in color from the regular faint coloration in EV controls to a much more intense yellow-orange color in RNAi roots (Fig. 4). This indicates the accumulation of a compound(s) with a new chromophore caused by the absence of MtCCD1 activity. Change of root color was the only phenotypical change observed in the composite plants with MtCCD1-silenced roots. There were no differences in shoot or root biomass, flowering, seed set, or other plant growth parameters between plants with mycorrhizal RNAi roots or mycorrhizal EV control roots (data not shown).

HPLC and Mass Spectrometry Analyses Indicate Accumulation of C\textsubscript{27} Apocarotenoid Derivatives in MtCCD1-Silenced Roots

Methanolic extracts of EV control roots and RNAi roots exhibiting a yellow to yellow-orange coloration were subjected to HPLC analyses. Using detection at 440 nm, at least seven new compounds were recognized in the mycorrhizal RNAi roots. (Fig. 5). They could not be detected in mycorrhizal EV controls (Fig. 5A) or in nonmycorrhizal RNAi roots (Fig. 5B). Analysis of roots from several individual transgenic root systems indicated a similar general pattern of accumulation but with variable proportions of the new compounds (Fig. 5B). Their UV spectra exhibited absorption maxima between 418 and 445 nm resembling those of recently described C\textsubscript{27} apocarotenoids, which are known to be yellow-orange (Fig. 5A; Yokoyama and White, 1966; Cooper et al., 2003). None of these compounds exhibited UV light absorption signatures reduced to a comparable extent. They exhibited a significant reduction in both experiments, but the residual levels in RNAi roots compared with EV control roots were still between 30% and 47% on average (Fig. 3C).

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of C₄₀ carotenoids, which have absorption maxima at higher wavelengths.

Using our standard procedure for liberating mycorrhadacin from its derivatives (Floss et al., 2008), alkaline treatments of the extracts from the transgenic roots were performed. HPLC analyses revealed two major peaks, which exhibited UV light spectra with maxima at 428 and 434 nm, again reminiscent to C₂₇ apocarotenoids (Fig. 6). The two compounds, with retention times of 19.1 and 21.8 min, respectively, termed Apo1 and Apo2, were isolated by semipreparative HPLC from methanolic root extracts after alkaline hydrolysis. To get data for their structures, the Apo1- and Apo2-containing fractions were subjected to liquid chromatography electrospray ionization mass spectrometry (LC-ESI/MS) analyses. Unfortunately, Apo1 coeluted with a compound showing a MS fragmentation pattern identical to that of the known root saponin bayogenin dihexoside (Huhman and Sumner, 2002; Huhman et al., 2005; Schliemann et al., 2008a), precluding identification. However, it was possible to identify Apo2. ESI-Fourier transformation ion cyclotron resonance (FTICR)-MS analysis indicated a [M + H]⁺ ion with a mass-to-charge ratio (m/z) 733.37944 corresponding to C₃₉H₅₇O₁₃, compatible with a C₂₇ 3-hydroxy-α-apo-10'-carotenoic acid glycosylated by two hexose moieties at the 3-hydroxyl position (Fig. 7). Due to the fact that AM-induced cyclohexenone derivatives characterized to date exclusively contain an α-ionone ring (Strack and Fester, 2006), this ionone type is also tentatively proposed for the C₂₇ apocarotenoid acid identified. The 15-eV collision-induced dissociation (CID) mass spectrum of m/z 733 obtained from triple quadrupole MS shows a consecutive loss of

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**Figure 5.** HPLC separation of methanol extracts from hairy roots. A, Comparison of mycorrhizal EV control roots (bottom trace) and mycorrhizal RNAi roots (top trace) with compound detection at 440 nm. Separation of native constituents revealed seven novel compounds, which appear exclusively in RNAi samples. Corresponding UV absorption maxima are shown in the inset. B, Comparison between nonmycorrhizal (NM) and mycorrhizal (M) roots of three individual RNAi root systems each. Root extraction, separation conditions, and compound terminology were as in A.
two hexose units ($\Delta m = 162$). The corresponding ions appear at $m/z 571 [M + H − 162]^+$ and $m/z 409 [M + H − 2 \times 162]^+$ (Fig. 7). A $[M + H]^+$ signal at $m/z 409$ and a UV light absorption maximum at 431.9 nm in 85% methanol/15% hexane has been reported for a C$_{27}$ apocarotenoic acid aglycone (Cooper et al., 2003), which is almost identical with our data under our solvent conditions and provides evidence for the proposed aglycone structure (Fig. 7). A consecutive loss of two molecules of water from the hydroxy and carboxy groups may lead to $m/z 391$ (major peak) and $m/z 373$, respectively. Additional characterization of the aglycone was possible by the 25-eV ESI-CID mass spectrum. The ion at $m/z 269$ may result from splitting next to the ionone ring (a). Significant key ions at $m/z 201$ (b) and $m/z 187$ (c) may be formed by bond cleavages at the indicated positions. Based on these analyses, compound Apo2 has been tentatively identified as a C$_{27}$ 3-hydroxy-α-apo-10′-carotenoic acid, glycosylated with two hexose moieties at the 3-hydroxyl position of the apocarotenoid (Fig. 7).

**Root Colonization by AM Fungi and Transcript Levels of AM Molecular Markers Are Largely Unaltered upon MtCCD1 Repression**

Potential alterations in colonization by AM fungi of MtCCD1-silenced roots were analyzed by standard ink staining tests using the individual transgenic root systems of experiment I. Assessment and calculation of mycorrhizal colonization parameters were carried out according to Trouvelot et al. (1986). There were no significant changes in either the total colonization of roots (frequency of mycorrhizal structures; F%), the density of mycorrhizal structures (m%), or the abundance of ink-stainable arbuscules in the colonized part of the root (a%; Fig. 8, A–C). A slight upward trend for F% and a% was recognizable rather than an adverse effect, but whether this is a reproducible effect requires further experiments. Arbuscules are ephemeral structures that constantly decay and reemerge with a life span of 7 to 10 d (Alexander et al., 1988). Following up earlier assessments of the ratios of various stages of arbuscule development in arbuscule populations (Floss et al., 2008), an acid-fuchsin staining analysis was performed. Confocal images of arbuscules from these stainings were used to classify individual arbuscules into four developmental stages ranging from early development to fully degraded and dead arbuscules, as described (Floss et al., 2008). The most obvious alteration in MtCCD1-repressed roots was a moderate, yet significant, increase in degenerating arbuscules from 32% to 48% of the total population of arbuscules (Fig. 8D).

In addition, the transcript levels of several AM molecular marker plant genes were determined by real-time reverse transcription (RT)-PCR. These markers include MtDXS2-1 as an AM-inducible gene involved in early steps of apocarotenoid biosynthesis (Walter et al., 2002; Floss et al., 2008) and MtPT4 as an AM-specific phosphate transporter (Harrison et al., 2002; Table I). Other AM markers have been specified by transcriptome analyses of mycorrhizal roots (Hohnjec et al., 2005) and have been used previously to assess alterations in mycorrhizal physiology upon MtDXS2 repression (Floss et al., 2008). None of the AM plant markers used in this study showed significant alterations in their transcript levels in mycorrhizal RNAi roots relative to mycorrhizal EV controls (Table II). The only exception exhibiting a minor, yet significant, decrease was MtLEC7, an AM-inducible lectin gene (Frenzel et al., 2005). Taken together, there was no indication of a major impairment of the AM symbiosis by silencing MtCCD1 expression.

**DISCUSSION**

In our studies on apocarotenoid biosynthesis, an experimental system of a symbiotic plant-microbe interaction (AM) is employed, which has turned out to be very useful (Walter et al., 2002; Hans et al., 2004; Floss et al., 2008). First, apocarotenoid biosynthesis in roots is inducible upon colonization by AM fungi and leads to massive accumulation of apocarotenoids. This proceeds in an apparently cell-specific manner related...
to the occurrence of fungal arbuscules (Fester et al., 2002a; Floss et al., 2008). Second, this accumulation involves transcriptional up-regulation of various steps of carotenoid biosynthesis and cleavage (Walter et al., 2000, 2002; Fester et al., 2002b; Hans et al., 2004; Sun et al., 2008; Table I). Third and most important for this

Figure 7. Mass spectral analysis of compound Apo2. A, Scheme of the proposed fragmentation of compound Apo2, whose proposed structure is shown at top left. Apo2 exhibits a m/z of 733 [M + H]+. Removing one or two hexose residues (increments of 162) results in fragments with m/z 571 and 409. Splitting of the aglycone next to the cyclohexene ring is proposed to yield fragment a (m/z 269). Additional fragments may be derived from fragmentation in the linear side chain of the Apo2 aglycone (m/z 391) at the positions indicated, resulting in fragments of m/z 201 (b) or m/z 187 (c). B and C, ESI-CID mass spectra of Apo2 (B, 15 eV; C, 25 eV).
study, the metabolic fate of primary carotenoid cleavage products is different from that in other systems. In the mycorrhizal root system used here, both the C\textsubscript{13} and C\textsubscript{14} cleavage products are easily traceable as oxidized and further modified downstream metabolites. In particular, the central C\textsubscript{14} linear product of C\textsubscript{40} carotenoid cleavage is converted to a stable yellow dicarboxylic acid (mycorradicin) usually accumulating in esterified form (Fester et al., 2002a). The ratio of C\textsubscript{13} to C\textsubscript{14} apocarotenoid end products in mycorrhizal roots exhibits large variations ranging from 0.4 to 0.5 in the system used here (Fig. 3) through values of 1 to 2 in AM wild-type roots of \textit{M. truncatula} or in AM cereal roots (Walter et al., 2000; Schliemann et al., 2008a) to as high as 27 in AM \textit{Allium porrum} roots (Schliemann et al., 2008b). These fluctuations argue for a variable metabolic fate of apocarotenoids in mycorrhizal roots of individual plant species. In other nonmycorrhizal systems, the C\textsubscript{14} primary cleavage product often escapes detection, since it is further metabolized to unknown products or is not volatile enough for gas chromatography-MS analysis (Auldridge et al., 2006a; Ibdah et al., 2006; Sun et al., 2008). One alternative system, in which the fate of the C\textsubscript{14} product has been investigated to some extent, is the rose petal, where a reduced C\textsubscript{14} dialdehyde downstream product has been identified: the diol rosalfluene. This dialcohol has been characterized as being colorless and unstable (Eugster and Märtik-Fischer, 1991), as opposed to the oxidized C\textsubscript{14} metabolite mycorradicin and its derivatives.

In this system of easy traceability of all predicted carotenoid cleavage products, we could clearly recognize a differential reduction of C\textsubscript{13} and C\textsubscript{14} metabolites upon knockdown of \textit{MtCCD1} expression in mycorrhizal hairy roots of \textit{M. truncatula}. The C\textsubscript{14} cleavage products (mycorradicin derivatives) were strongly reduced almost to the limits of detection upon strong repression of \textit{MtCCD1}, whereas the C\textsubscript{13} cleavage products (cyclohexenone derivatives) were only reduced to about 30% to 50% of the levels of EV controls (Fig. 3). This suggests that mycorradicin biogenesis depends on \textit{MtCCD1} activity but also that only one of the two C\textsubscript{13} cleavage products expected from the presumed symmetrical cleavage of C\textsubscript{40} carotenoid precursors (Fig. 1) originates from this activity. The other one appears to be generated by the action of a different enzyme. The results obtained from two extensive experiments of \textit{MtCCD1} silencing with many biological replicates are clearly contradictory to the hitherto discussed scheme of CCD1 action (Fig. 1).
Concomitant with the differential reduction of C13 and C14 apocarotenoids, a conspicuous color change was observed as a consequence of MtCCD1 repression. The rather faint orange-brown coloration of mycorrhizal E. coli control roots related to mycorradicin derivative accumulation was modified to a clearly more intense yellow-orange root color, indicating the accumulation of a different abundant chromophore (Fig. 4). This coloration also differed from the bright yellow color frequently observed in many cereal roots after mycorrhization, indicating massive accumulation of mycorradicin derivatives (Klingner et al., 1995; Walter et al., 2000). Root extraction and analysis by HPLC revealed the emergence of at least seven novel compounds in mycorrhizal RNAi roots (Fig. 5). Their UV spectra were similar to those of C27 apocarotenoid structural variants recently reported with single absorption maxima between 428 and 440 nm (Cooper et al., 2003). In particular, these spectra matched most closely to several C27 carboxylic acids reported by those authors. There was no indication of an accumulation of C9 carotenoids, which had been expected at the outset of our studies based on the previous hypothesis of CCD1 action (Fig. 1).

The methanolic extracts of RNAi roots were subjected to alkaline hydrolysis, resulting in only two major compounds again exhibiting characteristics of C27 apocarotenoids (Fig. 6). Based on ESI-MS and MS/MS, the structure of Apo2 was proposed to be a C27 carboxylic acid (3-hydroxy-α-apo-10′-carotenoid acid) linked with two hexose moieties at the 3-hydroxy position of the cyclohexene ring (Fig. 7). At least three key fragments (m/z 409, 391, and 269) comport precisely to major fragments obtained earlier by atmospheric pressure chemical ionization MS and MS/MS from a nonglycosylated 3-hydroxy-β-apo-10′-carotenoid acid (Cooper et al., 2003), corroborating the proposed structure. While there is still ambiguity on the position of the double bond in the cyclohexene ring (β- or α-ionone type), the α-ionone (ε-rings) type appears to be more likely. It has always been found in structural analyses of C13 cyclohexenone derivatives obtained from mycorrhizal roots (Strack and Fester, 2006; Walter et al., 2007). In addition, there is no explanation yet for the origin of the keto group in these derivatives. Therefore, the tentatively proposed but still unknown precursor for the AM-induced apocarotenoids is likely to be neither zeaxanthin (β,β-carotene rings) nor lutein (β,ε-rings). Whether the less common xanthophyll luctacaxanthin (Cunningham and Gantt, 2001) derived from ε,ε-carotene containing two α-ionone-like rings (ε-rings) is involved in this pathway is currently a rather speculative assumption. The C27 3-hydroxy-α-apo-10′-carotenoid acid is an intermediate predicted to be formed analogous to mycorradicin by a single instead of a double cleavage of this proposed xanthophyll precursor. The 3-hydroxy position of the C27 compound was found to be glycosylated in the extracts from RNAi roots, as opposed to the C13 cyclohexenone derivatives containing a keto group, indicating that its conversion into the keto group in the cyclohexene moiety may not be possible with the C27 apocarotenoid intermediate. MtCCD1 thus appears to catalyze the C27 → C14 + C13 cleavage but not the C27 → C20 + C13 cleavage. Taken together, there is ample evidence for the accumulation of several C27 apocarotenoids, most probably differently esterified, in mycorrhizal roots caused by silencing of MtCCD1 expression. C27 apocarotenoids have never been observed in wild-type mycorrhizal roots. Derivatives of C27 apocarotenoid core structures have only occasionally been identified in plants, perhaps due to the abundant activities of CCD1 enzymes in many plant tissues (Simkin et al., 2004b; Auldridge et al., 2006a). Examples for natural C27 apocarotenoid occurrence, either as carboxylic acids or as alcohols, are flowers of Boronia megastigma (Cooper et al., 2003), rose petals (Eugster and Märki-Fischer, 1991), and roots of Escobedia scabridifolia (azafrin; Kuhn and Brockmann, 1935).

The emergence of C27 apocarotenoids and not, as expected, C40 carotenoids upon blocking MtCCD1 gene expression in mycorrhizal RNAi roots clearly suggests the C27 apocarotenoids as the substrates of the MtCCD1 enzyme in planta. It further suggests a consecutive two-step cleavage mechanism of carotenoids following the schemes C40 → C27 + C13 and C27 → C14 + C13. Such a mechanism was proposed a number of years ago (Eugster and Märki-Fischer, 1991), but this scheme was abandoned in the wake of many studies on recombinant CCD1 proteins in E. coli showing symmetrical cleavage of a wide variety of C40 carotenoids in vitro. However, the in planta substrate(s) of the enzyme has remained elusive. Interestingly, a recent study on an Arabidopsis CCD1 loss-of-function mutant (ccd1-1) also came up with surprisingly few, if any, changes in precursor C40 carotenoid accumulation in the mutant. Leaf carotenoid levels were completely unchanged. Total seed carotenoid levels were significantly elevated by about 30% (Auldridge et al., 2006a). While this appears to indicate that CCD1 acts on C40 carotenoids at least in seeds, the result could also be explained by a feedback mechanism down-regulating

| DFCI Identifier | Gene | \(10^{-6}\times E_{cT} \) |
|-----------------|------|---------------------|
| TC94453         | MtPT4 | 146.5 ± 49.2        |
| TC95018         | MtGLP1 | 169.3 ± 41.2        |
| TC95567         | MtLEC7 | 80.3 ± 19.7         |
| TC95651         | MtDXS2-1 | 12.2 ± 3.0         |
| TC96500         | MtBCP1 | 5.0 ± 1.2           |
| TC100720        | MtGST1 | 96.3 ± 21.6         |
| TC106954        | MtSCP1 | 22.0 ± 5.0          |
the first cleavage step ($C_{40} \rightarrow C_{27} + C_{13}$) upon abnormal accumulation of the $C_{27}$ apocarotenoid intermediate, resulting from failure of the second step ($C_{27} \rightarrow C_{14} + C_{13}$; CCD1). A potential accumulation of $C_{27}$ apocarotenoids in the ccd1-1 mutant was not investigated, and a phenotype was not observed. Moreover, in an antisense approach on CCD1 expression in tomato, two transgenic lines with more than 90% reduction in CCD1 transcript levels were obtained, yet the reduction in C13 apocarotenoid volatile emission was only about 50% (Simkin et al., 2004a), reminiscent of the cyclohexenone derivative contents obtained in our study (Fig. 3). Fruit carotenoid content was not affected in these antisense plants. The authors concluded the likely existence of redundant activities for $C_{40}$ carotenoid cleavage (Simkin et al., 2004a). Similarly, strong cosuppression of PtCCD1 led to unexpectedly high residual levels of 24% to 42% in C13 β-ionone emission from petunia (Petunia hybrida) flowers (Simkin et al., 2004b). The conclusion of these authors for additional players in C40 carotenoid cleavage is supported by our data, but with the crucial difference that this is likely not a redundant but a complementary activity of at least two players in a cleavage process of $C_{40}$ to C14 and C13 compounds in two consecutive steps. The results obtained with the mycorrhizal root system, therefore, may be of general importance for the mechanisms of carotenoid cleavage in plants. Further supportive evidence comes from an in vitro study on recombinant AtCCD1 showing C13 β-ionone formation using a C30 apocarotenoid as substrate (Schmidt et al., 2006). Moreover, recent studies in the cyanobacterium Nostoc sp. PCC 7120 demonstrated the preference of a recombinant NosCCD (identical to previous NSC1) for cyanobacterial monomeric xanthophylls like zeaxanthin or canthaxanthin (Marasco et al., 2006; Scherzinger and Al-Babili, 2008). The NosCCD, structurally related to plant CCD1, also exhibits strong apocarotenoid cleavage activity for both C30 3-hydroxy-β-apo-8′-carotenal and C27 3-hydroxy-β-apo-10′-carotenal (Scherzinger and Al-Babili, 2008). These findings corroborate the notion that the major substrates for CCD1 enzymes in planta may not be C40 carotenoids but rather C27 apocarotenoid compounds.

The new view on carotenoid cleavage in planta brought forward here and summarized in Figure 9 may also reconcile a long-standing contradiction of how CCD1 enzymes get access to their substrates. CCD1 enzymes have been shown in several experimental systems and by several criteria to be located in the cytosol (Bouvier et al., 2003; Tan et al., 2003; Simkin et al., 2004a; Auldridge et al., 2006a). So how should CCD1 get access to their presumed C40 carotenoid substrates located within the plastid and separated from the CCD1 location by the plastidial membrane? The solution may be that in vivo C40 is a substrate for CCD1 only after damage or breakdown of plastids and release of C40 carotenoids into the cytosol. In intact plastids, CCD1 probably never gets access to C40 carotenoids. The primary $C_{40} \rightarrow C_{27} + C_{13}$ conversion is thus likely to take place inside the plastid, after which these two apocarotenoids are exported from the plastid to the cytosol by unknown mechanisms. Once in the cytosol, the C27 apocarotenoid can be further cleaved by abundant cytosolic CCD1 to a second C13 molecule and a C14 molecule. An export of C27 compounds into the cytosol is suggested by the finding that a glycosylated form of the C27 apocarotenoid carboxylic acid has been isolated (Fig. 7). Glycosylation is a common modification carried out by cytosolic glycosyltransferases. In ABA biosynthesis, the primary carotenoid cleavage of C40 cis-carotenoids catalyzed by NCEDs occurs inside the plastid in an analogous manner to that now proposed for C13 and C14 apocarotenoid biosynthesis. The C15 aldehyde ABA precursor (xanthoxin) then leaves the plastid through unknown mechanisms, and subsequent modification steps are extraplastidial (Nambara and Marion-Poll, 2005). Taken together, our new model for in planta cleavage of C40 carotenoids to C14 and C13 products is compatible with the data obtained in this study and with those from other laboratories.

What then could be the elusive CCD enzyme to perform the first cleavage step and generate $C_{27}$ from $C_{40}$ inside the plastid? One certainly cannot fully exclude the existence of an unknown novel CCD, but this CCD should then be completely unrelated in amino acid sequence to the existing nine CCD classes. This is considered rather unlikely. From the known CCDs, a prime candidate is CCD7, with the required 9,10 double bond cleavage specificity and a reported plastidial location. So far, only recombinant AtCCD7 has been shown to cleave C40 to C27 in E. coli (Schwartz et al., 2004). Native CCD7 (MAX3) is involved in one of two consecutive cleavage reactions necessary for the generation of a novel carotenoid-derived hormone. This hormone acting as an inhibitor of shoot branching is generated by the genes of the max pathway of Arabidopsis and by orthologous genes in many other plants, exemplified by rms5 in pea (Pisum sativum) and htd1 in rice (Oryza sativa; Booker et al., 2004, 2005; Snowden et al., 2005; Johnson et al., 2006; Zou et al., 2006; Mouchel and Leyser, 2007). The second cleavage activity required for the biosynthesis of the branching inhibitor is contributed by CCD8. Schwartz et al. (2004) showed cleavage activity of a recombinant AtCCD8 on a C27 substrate, resulting in C18 and C9 cleavage products. On the other hand, Auldridge et al. (2006a) reported cleavage of C40 carotenoid substrates to unknown products by recombinant AtCCD8.

More recently, the strict specificity of CCD8 for certain C27 compounds (β-apo-10′-carotenal and its alcohol), the conservation of this specificity in monocot and dicot species, and the nature of the C18 product (β-apo-13-carotenone) have been further elucidated (Alder et al., 2008). AtCCD7, however, cleaves multi-
Figure 9. Proposed organization of C_{40} carotenoid formation and cleavage to apocarotenoids in a compartmentalized mycorrhizal root cell. C_{40} carotenoids (exemplified by luteoxanthine as the tentatively proposed precursor for AM-induced apocarotenoids) synthesized inside plastids are proposed to be cleaved by plastidial CCD7 to one C_{27} aldehyde and one C_{13} ketone intermediate, which both can be exported into the cytosol in an unmodified or modified state (white arrows). The C_{27} intermediate in the cytosol is further cleaved by CCD1 to a C_{14} and a second molecule of C_{13}, which are, together with the first C_{13} molecule, modified to yield the oxidized C_{13} and C_{14} apocarotenoids (cyclohexenone and mycorradicin derivatives) found in mycorrhizal roots. For abbreviations, see Figure 1.

ple C_{40} carotenoids, including lycopene, β-carotene, and zeaxanthin (Booker et al., 2004), leading to the prospect that its biological function is not limited to the formation of the branching inhibitor (Alder et al., 2008). In addition, while double mutants of Arabidopsis ccd7 (max3-11) and ccd8 (max4-6) were phenotypically indistinguishable from either single mutant, a quantitative dosage effect of ccd8 heterozygotes on the branching phenotype argues for CCD8 activity as a limiting step in the pathway (Auldridge et al., 2006a).

The reported strict specificity of CCD8 for the conversion of C_{27} β-apo-10’-carotenal and its inability to convert 3-hydroxy-β-apo-10’-carotenal (Alder et al., 2008) appear to preclude a common C_{27} precursor for both the branching inhibitor and mycorrhizal apocarotenoids. In support of this preliminary conclusion, we have never observed an obvious branching phenotype on strongly mycorrhized plants, which might possibly arise from depleted C_{27} common precursor pools. The data available at present are thus in agreement with CCD8 catalyzing a specific step in a side route from C_{27} to the branching inhibitor, while CCD7 might be required for both branching inhibitor and mycorrhizal apocarotenoid biosynthesis (Fig. 9). Finally, while this article was under revision, two reports became available online, suggesting strigolactones as the long sought after carotenoid-derived branching inhibitor hormone and confirming the specific role of CCD8 in its synthesis (Gomez-Roldan et al., 2008; Umehara et al., 2008).

Based on other very recent data, CCD4 must now also be considered a potential player for the in planta generation of C_{27} apocarotenoids. Earlier investigations have attributed 7,8 (7’,8’) cleavage activity to a CsZCD of saffron, representing an incomplete CsCCD4 sequence (Bouvier et al., 2003), but the aforementioned recent work now reports β-ionone formation from recombinant full-length CsCCD4 acting on β-carotene as a substrate in E. coli, which implies 9,10 and/or 9,10 (9’,10’) cleavage activity (Rubio et al., 2008). CCD4 also fulfills the condition of having a proven localization within plastids in plastoglobules (Ytterberg et al., 2006), but it has not been shown to produce C_{27} apocarotenoids in vitro. As for the in planta action of CCD4, an RNAi-silencing approach on CCD4 expression in white-petal cultivars of Chrysanthemum morifolium resulted in the generation of yellow petals, which indicates that the white-petal phenotype is brought about by carotenoid degradation (Ohmiya et al., 2006). Loss of function of CCD4 thus strongly increased carotenoid content in Chrysanthemum petals as opposed to no or minor effects upon silencing or mutating CCD1 genes (Simkin et al., 2004a, 2004b; Auldridge et al., 2006a).

While one cannot make a safe case for either CCD7 or CCD4 as the missing player for the C_{40} → C_{27} cleavage at this time, there are a couple of arguments favoring CCD7 at least for the mycorrhizal apocarotenoids. Mining of the extensive M. truncatula EST databases does not indicate the presence of MtCCD4-related transcripts in roots or mycorrhizal roots, whereas 18 ESTs can be identified from leaf, stem, and pod libraries. Lack of root expression and strong expression in flower or reproductive tissues has also been reported for Chrysanthemum and saffron CCD4 genes (Ohmiya et al., 2006; Rubio et al., 2008). On the contrary, CCD7 and CCD8 are preferentially expressed in roots (Booker et al., 2004; Snowden et al., 2005; Auldridge et al., 2006a).
cyclohexenone and C14 mycorradicin derivatives compared with mycorrhizal wild-type plants (M.H. Walter, unpublished data).

Besides clarifying the role of CCD1 in the biosynthesis of cyclohexenone and mycorradicin derivatives, our RNAi approach was also meant to elucidate potential functions of either of these apocarotenoids in the AM symbiosis. Such functions might be revealed by the reduction or lack of these compounds in roots of RNAi plants. To proceed toward this goal, we determined the transcript levels of several AM molecular markers. However, neither MtPT4 nor any of the other markers showed a strong decrease (Table II) comparable to the strong down-regulation recently obtained by strong silencing of MtDXS2 (Floss et al., 2008). Then again, the differential reduction of cyclohexenone and mycorradicin derivatives constitutes an opportunity to enter into a discussion on which of the two apocarotenoids might be a major player in the AM symbiosis. Despite strong reduction of mycorradicin derivative levels in MtCCD1 RNAi roots (Fig. 3), the AM molecular markers were largely unaltered (Table II), arguing against a contribution of mycorradicin to the effect on these markers. On the contrary, cyclohexenone derivative levels were reduced to only 30% to 50% of those of EV controls (Fig. 3), which might simply not be sufficient to strongly affect AM marker gene expression. Extensive reduction of cyclohexenone derivative levels as accomplished by strong MtDXS2 silencing reproducibly led to a dramatic abolishment of AM-induced plant marker gene expression (Floss et al., 2008). In the MtCCD1-silenced roots used here, one of the AM markers (MtLEC7) exhibited a minor, but significant, down-regulation (Table II). Interestingly, this marker was also negatively affected in roots moderately silenced for MtDXS2 expression, while all other markers were unaltered (Floss et al., 2008). The outcome of strong repression of MtCCD1 thus appears to be similar to the moderate repression of MtDXS2, which both share moderately but not strongly reduced levels of cyclohexenone derivatives (Table II; Floss et al., 2008).

While any consequences of the strong silencing of MtCCD1 on AM molecular markers were hardly detectable, a commonality to MtDXS2 silencing was the significant increase in older, degenerating arbuscules (Fig. 8D; Floss et al., 2008). Again, these alterations follow quantitatively the moderate alterations of cyclohexenone derivatives in MtCCD1-silenced roots more closely than the strongly decreased levels of mycorradicin derivatives. Only a further reduction of cyclohexenone derivatives, as achieved in the MtDXS2-silencing approach, led to a more pronounced effect on the numbers of degenerating arbuscules and added an increase in dead arbuscules (Floss et al., 2008). Additional experiments recently indicated that with the MtDXS2 target, moderate levels of silencing and cyclohexenone derivative reduction can also bring about an increase in the number of degenerating arbuscules independent of the strong drop in AM marker transcript levels (D.S. Floss and M.H. Walter, unpublished data). These results lend further support to a possible link between the moderate reduction of cyclohexenone derivative levels (Fig. 3) and the moderate increase in arbuscules at the degenerating stage (Fig. 8D). Both of the observations of changes of AM marker transcript levels and of the number of degenerating arbuscules are thus compatible with a role of cyclohexenones in these two instances, which is in agreement with previous arguments (Walter et al., 2007). Mycorradicin derivatives accumulate massively next to degenerating arbuscules within osmiophilic droplets finally located to vacuoles (Fester et al., 2002a). These derivatives might have other functions in the decay and reemergence of arbuscules, but they seem to be not directly involved in regulating the number of physiologically active arbuscules.

CONCLUSION

While the definite roles of cyclohexenone and mycorradicin derivatives in the AM symbiosis still need to be clarified, this report sheds new light on the mechanism of carotenoid cleavage and on its compartmentation in plant cells. It is now most interesting to search for C27 apocarotenoids accumulating in other CCD1-repressed plants or respective mutants. Moreover, a potential contribution of CCD7 and/or CCD4 to the generation of flower scents, fruit aromas, or the flavor of wines complementing CCD1 activities (Simkin et al., 2004a; Mathieu et al., 2005) will now be a promising area to study. For mycorrhizal research, the CCD7 gene will constitute a new target to modify the levels of mycorrhizal apocarotenoids and to finally elucidate their function in the AM symbiosis.

MATERIALS AND METHODS

Plant Cultivation and Fungal Inoculation

Medicago truncatula var. Jemalong was cultivated in pots on expanded clay substrate (Lecaton; Fibo Exclay) in a greenhouse during the months of June and July (RNAi experiments) or December and January (wild-type cultivation) with a 16-h daily light period. Plants were supplied with deionized water three times per week and with a phosphate-reduced fertilizer (10 ml of half-strength Long Ashton medium; 20% of regular phosphate) once a week. Seven-day-old wild-type plantlets or 6-week-old plants from hairy root transformation with transgenic root systems and wild-type shoots were inoculated with the AM fungus Glomus intraradices isolate 49 provided by H. von Alten (University of Hannover; Maier et al., 1995). This was done by transferring to clay substrate containing 30% (v/v) fungal inoculum enriched
in propagules by previous co-cultivation with Allium porrum 'Elefant' as described (Floss et al., 2008).

**MCCDI1 cDNA Cloning and Expression in Escherichia coli**

An 825-bp fragment of a MCCDI cDNA was amplified from mycorrhizal root RNA of *M. truncatula* using the primers MCCDI1-fw (5'-ACGCTT-GATTATGGAGAAG-3') and MCCDI1-re (5'-ATCACGTCTTTATCTTTAC-3') deduced from TC100912 of the DFCI *M. truncatula* gene index (version 8; http://compbio.dfci.harvard.edu/cgi-bin/tgi/tgi/gimain.pl?gdb=medicago). This fragment was used as a probe to screen a cDNA library from *M. truncatula* mycorrhizal roots as described (Walter et al., 2002). Several full-length clones with sequence identity to TC100912 were obtained.

For bacterial expression, the coding sequence of *MCCDI1* was amplified from the cDNA using the primers MccID1-fw (5'-TATCTAGGAGTCAGCTGTAGAATATCC-3') and MccID1-re (5'-TATCTGAGGCTTTAATAGGAGG-3') and cloned in the plasmid sites of the plasmid vector carrying pDSRED1 as a fluorescent marker (Limpens et al., 2002). Details of the cloning process and further steps of construct introduction into Agrobacterium rhizogenes Arqaud as well as the hairy root transformation procedure have been described (Floss et al., 2008). Briefly, 7-d-old *M. truncatula* seedlings were injected with small volumes of bacterial culture (50–100 μl) three to five times into their hypocotyls. Seedlings were transferred into humid expanded clay for a 2-d recovery and then placed in a growth chamber (20°C, 16 h of light/8 h of dark) for 2 weeks, after which emerging hairy roots were covered by a vermiculite:sand (1:1) mixture. Four weeks after *A. rhizogenes* infection, a first screen for pDSRED1 fluorescent roots was performed with a fluorescence stereomicroscope (Leica MZ FLIII with a DsRED1 filter), removing from each plant individually all nonfluorescent hairy roots as well as the wild-type roots. A second screen was carried out after a further 2 weeks of cultivation. Composite plates carrying exclusively transgenic roots were then inoculated with *G. intraradices* and further cultivated as described above.

**Metabolite Analyses**

Homogenized root samples (200 mg) from individual plants were each extracted three times with 400 μl of 80% aqueous methanol. After centrifugation, the supernatants were analyzed by reverse-phase HPLC to determine the levels of cyclohexenol derivatives (Schliemann et al., 2008a). For mycorrhizaric derivative quantification, 300 μl of the methanolic extracts was adjusted with potassium hydroxide to a final concentration of 0.5 M and a rapid gradient from 0% to 100% B within 60 min. Alkaline hydrolysis was done as described for mycorradicin, and reaction products were separated by applying a constant gradient from 0% to 100% B in 35 min followed by a 4-min hold and a rapid switch to 0% B within 1 min. For semipreparative HPLC, 1.5% aqeous H3PO4 was replaced by 2% formic acid and by a gradient raising solvent B from 0% to 100% within 35 min. The high-resolution ESI mass spectra (ESI-FTICR-MS) of fractions Apo3 and Apo2 isolated after alkaline hydrolysis were obtained from a Bruker Apex III Fourier transform ion cyclotron resonance mass spectrometer (Bruker Daltonics) equipped with an Infinity cell, a 7.0-tesla superconducting magnet (Bruker), a radiofrequency-only hexapole ion guide, and an Apollo electro-spray ion source (Agilent; off axis spray). Nitrogen was used as a drying gas at 150°C. The sample solutions were introduced continuously via a syringe pump with a flow rate of 120 μl h⁻¹. The m/z value obtained for Apo2 was 733.3948 [M + H⁺], calculated for C₃₉H₅₉O₁₃,M+1. The positive ion ESI mass spectra of Apo2 were obtained from a Finnigan MAT TSQ Quantum Ultra AM system equipped with a hot ESI source (electrospray voltage, 3.0 kV; sheath gas, nitrogen; vaporizer temperature, 50°C; capillary temperature, 250°C). The MS was coupled with a Surveyor Plus Micro-HPLC (Thermo Electron) equipped with a Ultrasphere ES RP18e column (5 μm, 1 × 100 mm; Separex). For HPLC, a gradient system was used starting from solvent C (90% water/10% acetonitrile, each of them containing 0.2% acetic acid) to solvent D (5% water/95% acetonitrile) within 15 min followed by a hold at 5% water/95% acetonitrile for a further 15 min with a flow rate of 50 μl min⁻¹. The CID mass spectra of Apo2 [M + H⁺] ions at m/z 733 (retention time, 19.7 min) were recorded during the HPLC run with collision energies of 15 and 25 eV, respectively (collision gas, argon; collision pressure, 1.5 mTorr).

For Apo2: 15-eV ESI-CID mass spectrum [m/z, relative intensity (%)]: 733 [M + H⁺] (10), 715 (8), 571 [M + H⁺ – C₄H₅O₆ (hexose moiety)] (22), 553 (8), 409 [M + H⁺ – 2 × C₄H₅O₆] (75), 391 [M + H⁺ – 2 × C₂H₄O₂ – H₂O] (100), 373 (8).

For Apo2: 25-eV ESI-CID mass spectrum [m/z, relative intensity (%)]: 733 [M + H⁺] (7), 409 [M + H⁺ – 2 × 162 (hexose moiety)] (43), 391 [M⁺ + H₂O – 2 × 162-H₂O] (29), 291 (17), 269 (100), 210 (47), 209 (29), 201 (61), 187 (41), 183 (35), 145 (12).

**Root Staining Procedures and Microscopic Evaluation of Fungal Structures**

Root fractions from a defined part of the root systems (center) were stained by 5% (w/v) Shaefer skrip jet black; Shaefer Manufacturing) in 2% acetic acid (Vierheilig et al., 1998). Root colonization, fungal structures, and arbuscule abundance were assessed using a stereomicroscope. The results were evaluated using the software Mycoocal according to Trouvelot et al. (1986). For experiment I, additional root aliquots were stained with 0.01% acid-fuchsin in lactoglycerol (lactic acid:glycerol:water, 1:1:1 [v/v/v]). Further details of subsequent visualization and classification of fungal arbuscules by confocal scanning microscopy (LSM 510 Meta; Zeiss) using the 543-nm laser line for excitation have been described elsewhere (Floss et al., 2008).

**Real-Time RT-PCR**

Total RNA was prepared using the Qiagen RNeasy Plant Mini Kit followed by DNase digestion (RNase-free DNase Set; Qiagen). RNA (1 μg) was reverse transcribed into cDNA with M-MLV Reverse Transcriptase RNaSe H Minus, Point Mutant (Promega) using oligo(dT)19 primers. Real-time RT-PCR assays were performed using a SYBR Green-based kit according to the manufacturer’s protocol (Applied Biosystems). Five nanograms of reverse-transcribed RNA was used in a Mx 3000P QPCR system (Stratagene) using conditions and primer sets recently described (Floss et al., 2008). Briefly, the primer information for the four most important genes is given here: *MCCDI1*-qPCR-fw, 5'-TCCTGATCTTACCCGCTAT-3'; *MCCDI1*-qPCR-re, 5'-AGGCCGAGTGTACAAACG-3'; *MtPT4*-qPCR-fw, 5'-CACCTGGATCTAAATCTTTAATGAAGCTC-3' and *MtPT4*-qPCR-re, 5'-CGAA-TTCCTTCTCACAACAAAGA-3'; *MtPT4*-qPCR-fw, 5'-AGAAGGAAGCTGCT-GAGATGAAC-3'. The relative transcription (E(ΔCt)) was calculated by subtracting the Ct values of the target gene from the arithmetic mean cycle threshold (Ct) value of the normalizing translation elongation factor of *M. truncatula* (MEfE) obtained
from three technical replicates. The efficiency of each PCR was determined by import of the fluorescence data into the LinReg PCR software (Ramakers et al., 2003).

Statistical Analysis

Data were evaluated for normal distribution and, if positive, a one-way test was performed, which at homogeneity of variance corresponds to ANOVA. If data did not comply with a normal distribution, the Kruskal-Wallis test was applied. The graphs show mean and so values of individual transgenic root systems transformed by EV or the RNAi constructs of single composite plants.

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession number FM204879 (MtCCD1).

Supplemental Data

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

Supplemental Figure S1. Alignment of amino acid sequences from 14 plant CCD1 proteins, including MtCCD1.

ACKNOWLEDGMENTS

We thank M.J. Harrison (Boycote Thompson Institute) for providing the M. truncatula cDNA library; R. Geurts (Wageningen University) for providing pRedRoot and pRNAi; C. Rameau (INRA) for providing pea mss mutant seeds, and H. Bothe (Cologne University) for providing the mycorrhizocin dimethyler ester standard. The skillful technical assistance of K. Manke and C. Kuhnt (Leibniz-Institut für Pflanzenbiochemie) is gratefully acknowledged.

Received June 20, 2008; accepted September 8, 2008; published September 12, 2008.

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