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Alkylresorcinol Synthases Expressed in *Sorghum bicolor* Root Hairs Play an Essential Role in the Biosynthesis of the Allelopathic Benzoquinone Sorgoleone


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*Sorghum bicolor* is considered to be an allelopathic crop species, producing phytotoxins such as the lipid benzoquinone sorgoleone, which likely accounts for many of the allelopathic properties of *Sorghum* spp. Current evidence suggests that sorgoleone biosynthesis occurs exclusively in root hair cells and involves the production of an alkylresorcinolic intermediate ([Z,Z]-8,11,14-pentadecatrienyl]resorcinol) derived from an unusual 16:3 fatty acyl-CoA starter unit. This led to the suggestion of the involvement of one or more alkylresorcinol synthases (ARSs), type III polyketide synthases (PKSs) that produce 5-alkylresorcinols using medium to long-chain fatty acyl-CoA starter units via iterative condensations with malonyl-CoA. In an effort to characterize the enzymes responsible for the biosynthesis of the pentadecyl resorcinol intermediate, a previously described expressed sequence tag database prepared from isolated *S. bicolor* (genotype BTx623) root hairs was first mined for all PKS-like sequences. Quantitative real-time RT-PCR analyses revealed that three of these sequences were preferentially expressed in root hairs, two of which (designated ARS1 and ARS2) were found to encode ARS enzymes capable of accepting a variety of fatty acyl-CoA starter units in recombinant enzyme studies. Furthermore, RNA interference experiments directed against ARS1 and ARS2 resulted in the generation of multiple independent transformant events exhibiting dramatically reduced sorgoleone levels. Thus, both ARS1 and ARS2 are likely to participate in the biosynthesis of sorgoleone in planta. The sequences of ARS1 and ARS2 were also used to identify several rice (*Oryza sativa*) genes encoding ARSs, which are likely involved in the production of defense-related alkylresorcinols.

**INTRODUCTION**

Allelopathy, a form of chemical warfare between plants, can be defined as the production and release of chemical substances by one species that inhibit the growth of another species (Inderjit and Duke, 2003; Weston and Duke, 2003). Allelopathic interactions have been proposed to have profound effects on the evolution of plant communities through the loss of susceptible species via chemical interference and by imposing selective pressure favoring individuals resistant to inhibition from a given allelochemical (e.g., Schulz and Wieland, 1999). Furthermore, allelopathic compounds released by grain crop species are thought to play a significant role in the utility of cover crops and intercropping systems where they act as weed suppressants. Allelopathic compounds have been characterized in a number of plants, such as black walnut (*Juglans nigra*), wheat (*Triticum aestivum*), rice (*Oryza sativa*), and sorghum (*Sorghum bicolor*; Bertin et al., 2003; Inderjit and Duke, 2003; Duke et al., 2005).

Despite the ecological and agronomic importance of allelochemicals, relatively few of the corresponding biosynthetic pathways have been characterized in detail at the molecular level. One notable exception is the identification and characterization of all the genes encoding the enzymes responsible for the biosynthesis of the benzoxazinoid 2,4-dihydroxy-7-methoxy-2H-1,4-benzoxazin-3(4H)-one in *Zea mays* (Frey et al., 1997). Benzoxazinoids are thought to act as allelopathic chemicals in the rhizosphere, in addition to being defense compounds against...
Sorgoleone, an allelochemical of particular interest to plant chemical ecology as well as agriculture, has only been found to be produced by members of the genus Sorghum (Czarnota et al., 2003b; Baerson et al., 2008b). The term sorgoleone is most frequently used to describe the compound corresponding to the predominant congener identified in sorghum root exudates (Netzly et al., 1988; Kagan et al., 2003), 2-hydroxy-5-methoxy-3-[[Z,Z]-8',11',14'-pentadecatrieny]resorcinol (Figure 1), which has been estimated to account for between ~40 and 90% of the exudate material (w/w) in various accessions (e.g., Nimbal et al., 1996; Czarnota et al., 2001; Baerson et al., 2008a; Dayan et al., 2009). The remaining exudate consists primarily of 4,6-dimethoxy-2-[[Z,Z]-8',11',14'-pentadecatrieny]resorcinol (methoxy-dihydrosorgoleone) and sorgoleone congeners differing in the length or degree of saturation of the aliphatic side chain and in the substitution pattern of the quinone ring (Erickson et al., 2003; Kagan et al., 2003; Ramíndio et al., 2003; Dayan et al., 2009). The fact that sorgoleone acts as a potent broad-spectrum inhibitor active against many agronomically important monocotyledonous and dicotyledonous weed species and appears to affect multiple targets in vivo (e.g., Netzly and Butler, 1986; Einhellig and Souza, 1992; Nimbal et al., 1996; Ramíndio et al., 1998; Czarnota et al., 2001; Bertin et al., 2003; Duke, 2003) may make it promising for development as a natural product alternative to synthetic herbicides (Duke, 2003).

The biosynthesis of sorgoleone is thought to occur exclusively in root hairs, which appear as cytoplasmically dense cells in sorghum, containing large osmiophilic globules presumably associated with sorgoleone rhizosecretion (Czarnota et al., 2001, 2003a). Prior labeling studies have indicated a polyketide origin for the quinone ring of sorgoleone (Fate and Lynn, 1996; Dayan et al., 2003), thus lending support for the initial steps in the proposed biosynthetic pathway shown in Figure 1, in which 5-pentadecatrienyl resorcinol (5-[[Z,Z]-8',11',14'-pentadecatrieny]resorcinol) is produced by a polyketide synthase enzyme accepting a 16:3Δ9,12,15 fatty acyl-CoA starter unit. A specific subclass of type III polyketide synthases, referred to as alkylresorcinol synthases (first described in microorganisms by Kozubek and Tyman, 1999; Kozubek et al., 2001; 1999; Kozubek et al., 2001). Plant-derived phenolic lipids have also been used by industry, for example, in manufacturing formaldehyde-based polymers and in lacquering processes (Kozubek and Tyman, 1999).

Prior studies on type III PKS-like sequences from S. bicolor have involved the characterization of eight sequences (designated CHS1-8) obtained from genomic library screens and analysis of ESTs (Lo et al., 2002; Yu et al., 2005). Recombinant enzyme studies have identified CHS8 as an STS and CHS2 as a typical CHS-type enzyme, and it is has been proposed that CHS1, 3, 4, 5, 6, and 7 also represent CHS-type enzymes given their high degree of sequence identity (>97.5%) with CHS2 (Yu et al., 2005). In this work, we report the cloning and functional characterization of two paralogous alkylresorcinol synthases (ARSs) from S. bicolor (genotype B73x23), designated ARS1 and ARS2, proposed for the biosynthesis of the 5-pentadecatrienyl resorcinol precursor to sorgoleone. At present, relatively little functional data exists concerning the genes and corresponding enzymes involved in the biosynthesis of alkylresorcinols in higher plants; thus, this study provides new tools for...
Figure 1. Proposed Biosynthetic Pathway of the Allelochemical Sorgoleone.

Dihydrosorgoleone, the hydroquinone produced in vivo, is thought to undergo autoxidation once secreted into the rhizosphere to yield sorgoleone, a more stable benzoquinone. The involvement of one or more cytochrome P450s in the formation of dihydrosorgoleone from 5-pentadecatrienyl resorcinol-3-methyl ether is at present speculative, as indicated by a question mark. DES, fatty acid desaturase; OMT, O-methyltransferase; P450, cytochrome P450.
exploring related pathways, particularly in the Poaceae family where the occurrence of presumed phytoanticipin alkylresorcinols is widespread. In demonstration of the latter, the sequences of ARS1 and ARS2 were also used in this work to identify several genes from rice (ssp japonica cv Nipponbare) likely involved in the biosynthesis of alkylresorcinols in this species.

RESULTS AND DISCUSSION

Tissue-Specific Accumulation of the 5-[(Z,Z)-8',11',14'\-Pentadecatrienyl] Resorcinol Intermediate

Given the evidence obtained from prior studies (e.g., Czarnota et al., 2001, 2003a), it is reasonable to speculate that the ARSs participating in the biosynthesis of sorgoleone are predominantly or exclusively expressed in root hairs and that the expected product, 5-pentadecatrienyl resorcinol, should also predominantly accumulate within this cell type. To explore this further, methanol extracts prepared from root hairs, root systems, developing panicles, stems, immature and fully expanded leaves, and shoot apices were analyzed by gas chromatography-mass spectrometry (GC-MS) for the presence of 5-[(Z,Z)-8',11',14'-pentadecatrienyl]resorcinol. As shown in Figure 2A, 5-pentadecatrienyl resorcinol was identified from total ion chromatograms of extracts prepared from the total root and isolated root hairs (retention time 14.8 min) by extracted ion monitoring at a mass-to-charge ratio (m/z) of 314, and the corresponding mass spectra for the peaks revealed characteristic fragment ions supporting this identification (Figure 2A). Significantly, 5-pentadecatrienyl resorcinol was not detectable in any of the other tissues; more importantly, this identification (Figure 2A). Significantly, 5-pentadecatrienyl resorcinol was identified from total ion chromatograms of extracts prepared from the total root and isolated root hairs (retention time 14.8 min) by extracted ion monitoring at a mass-to-charge ratio (m/z) of 314, and the corresponding mass spectra for the peaks revealed characteristic fragment ions supporting this identification (Figure 2A). Significantly, 5-pentadecatrienyl resorcinol was not detectable in any of the other tissues; moreover, signal levels obtained were consistently far higher in isolated root hairs than in total root systems. Given that the total root samples used in this study were harvested from 8-d-old seedlings (grown under conditions identical to those used for root hair isolations) and contained extensive amounts of root hairs, it is likely that root hairs contributed a large percentage of the total 5-pentadecatrienyl resorcinol detected in these samples. Collectively, these data are consistent with the suggested sorgoleone biosynthetic pathway localization in root hairs, involving a 16:3 fatty acyl-CoA using ARS expressed predominantly in this cell type (Czarnota et al., 2001, 2003a; Dayan et al., 2007).

Identification of PKS-Like Transcripts Preferentially Expressed in Root Hairs

The high levels of sorgoleone produced and exuded by root hair cells of Sorghum spp members (Czarnota et al., 2001) suggests that mRNAs encoding enzymes associated with sorgoleone biosynthesis could be among the most abundant in this cell type. Therefore, in an effort to identify and functionally characterize these enzymes, we previously initiated a functional genomics strategy involving the analysis of a data set comprising 5468 ESTs derived from isolated root hair cells of S. bicolor genotype BTx623 (Pan et al., 2007; Baerson et al., 2008a). Importantly, BTx623 is also the genotype used to generate the recently completed sorghum genome sequence (Patterson et al., 2009); thus, additional information, such as predicted gene structures and chromosomal organization, can be readily obtained for all contigs identified within the root hair EST data set.

To identify potential ARS-encoding transcripts expressed in root hairs, the root hair ESTs were mined for candidate PKSs using both the MAGIC Gene Discovery software (Cordonnier-Pratt et al., 2004) and analyzed by BLASTN and TBLASTN searches (Altschul et al., 1997) using functionally characterized plant type III PKS sequences as queries. From these analyses, nine PKS-like ESTs were identified, which assembled into five unique sequences by cluster analysis, three of which were represented by a single EST (i.e., singletons). Two of the PKS-like assemblies (ID numbers 2_126 and 2_127) are each composed of three ESTs and collectively represent ~0.11% of the total 5468 expressed sequences identified in root hairs (Baerson et al., 2008a). Additionally, two of the sequences (ID numbers 0_164 and 0_1821) correspond to CHS5 and CHS1, respectively, previously identified by Lo et al. (2002).

To determine if the identified root hair PKS-like sequences exhibit expression patterns correlating with the accumulation of 5-pentadecatrienyl resorcinol (Figure 2A), all five unique root hair PKS-like sequences were subjected to quantitative real-time RT-PCR analysis. Gene-specific primers were designed for monitoring mRNA steady state accumulation levels in assays using cDNAs prepared from root hairs, root systems, developing panicles, stems, immature and fully expanded leaves, and shoot apices (Figure 2B). All cDNAs used in these experiments were derived from the identical tissue samples used for GC-MS analysis of 5-pentadecatrienyl resorcinol.

As shown in Figure 2B, three of the five PKS-like candidates (2_127, 2_126, and 0_1848) exhibited root hair preferential expression patterns, whereas sequences 0_164 (CHS5) and 0_1821 (CHS1) were maximally expressed in developing leaves and total roots, respectively. The expression patterns of the five PKS-like sequences were further analyzed in silico by monitoring EST counts within the 18 different S. bicolor (genotype BTx623) EST libraries developed by the University of Georgia Laboratory for Genomics and Bioinformatics (http://www.fugen.org/Sorghum.htm). Interestingly, ESTs corresponding to PKS-like sequences 0_1848, 0_164 (CHS5), and 0_1821 (CHS1) were found in several different EST libraries, whereas sequences 2_127 and 2_126, exhibiting the most highly root hair preferential expression patterns in quantitative RT-PCR experiments (Figure 2B), could be identified only in root hair ESTs.

Complete open reading frames (ORFs) were determined for the three candidate sequences exhibiting root hair preferential expression (2_127, 2_126, and 0_1848) by 5’-rapid amplification of cDNA ends (RACE). The predicted full-length ORFs for all three sequences exhibited extensive sequence similarity at the amino acid level to previously characterized plant type III PKSs (Austin and Noel, 2003) and contained conserved residues and motifs putatively associated with catalysis and substrate binding, based on the crystal structures determined for Medicago sativa (CHS2) and Gerbera hybrida (2-pyrene synthase [2-PS]) type III enzymes (Ferrer et al., 1999; Jez et al., 2000). The predicted ORFs derived from all three PKS-like sequences encode ~43-kD proteins with predicted isoelectric points of 6.27 (2_127), 5.67 (2_126), and 5.78 (0_1848). Additionally, the ORFs predicted for
Figure 2. Comparison of 5-Pentadecatrienyl Resorcinol and PKS-Like Transcript Accumulation in Various S. bicolor Tissues.

(A) The 5-Pentadecatrienyl resorcinol levels were determined by GC-MS analysis of methanol extracts prepared from isolated root hairs (top panel) and total roots (bottom panel) of 8-d-old etiolated seedlings of S. bicolor genotype BTx623. Extracted ion chromatograms are shown defined at m/z 314, and 5-pentadecatrienyl resorcinol peaks (retention time 14.8 min) are indicated by arrows. The corresponding mass spectrum for 5-pentadecatrienyl resorcinol is shown as an inset in the bottom panel.

(B) The relative expression levels of five PKS-like contig sequences identified in root hair ESTs were determined by quantitative real-time RT-PCR using gene-specific primers. Data were normalized to an internal control (18S rRNA), and the ΔΔCt method was used to obtain the relative expression levels for each sequence, expressed as mean ± SD from assays performed in triplicate.
2_126 and 2_127 share ~91% identity at the amino acid level, and both share ~60% amino acid identity with the predicted ORF for 0_1848 (further details of these sequences will be shown and discussed below). BLASTN queries (Altschul et al., 1997) against the *S. bicolor* genomic sequence revealed that 2_127 (encoded by LOC:Sb05g022500 and LOC: Sb05g022510) separated by ~10 kb in opposite orientation on chromosome 5 (http://www.phytozome.net). These analyses also confirmed the presence of a 198-bp intron within the 5′ coding sequence corresponding to 0_1848, whereas the coding sequences corresponding to 2_126 and 2_127 were not found to contain introns.

**Activity of Recombinant Type III PKSs**

Recombinant enzyme studies were next performed to examine substrate preferences for the three putative type III PKS enzymes maximally expressed in root hairs. The full-length ORFs determined for the sequences 2_127, 2_126, and 0_1848 were over-expressed in *Escherichia coli* as N-terminal polyhistidine fusions, which were then purified by Ni²⁺ affinity chromatography. Acyl-CoAs varying in length and degree of saturation were tested in enzymatic assays with all three recombinant enzymes, as well as the nonlinear starter units benzyol-CoA, isovaleryl-CoA, and isobutyryl-CoA used by several other plant type III PKS enzymes (Austin and Noel, 2003). Trimethylsilyl (TMS) derivatized products were detected by GC-MS selected ion monitoring at m/z 268 for the quantification of 5-alkyresorcinols as previously described (Suzuki et al., 2003).

The results of the recombinant enzyme assays with the enzymes encoded by 2_126 and 2_127 are shown in Figure 3. Both enzymes were able to catalyze the formation of 5-alkyresorcinols using various fatty acyl-CoA starter units with malonyl-CoA as the extender unit and will therefore be hereafter referred to as ARS1 and ARS2, respectively (Figure 3). Overall, the activity profiles exhibited by ARS1 and ARS2 were quite similar; for example, nearly overlapping profiles were obtained for fatty acyl-CoA starter units possessing saturated chains ranging in length from C6 (hexanoyl-CoA) to C20 (arachidoyl-CoA; Figure 3). Furthermore, similar preferences were observed for both enzymes among the unsaturated acyl-CoA substrates palmitoyl-CoA (C16:1Δ9), hexadecatrienoyl-CoA (C16:3Δ9,12,15), oleoyl-CoA (C18:1Δ9), linoleoyl-CoA (C18:2Δ9,12), and arachidonoyl-CoA (C20:4Δ5,8,11,14), with maximal activities observed with palmitoyl-CoA (Figure 3). Among all the acyl-CoAs evaluated, maximal 5-alkyresorcinol-forming activity was observed with myristoyl-CoA (C14) and palmitoleoyl-CoA (C16:1Δ9) for ARS1 and ARS2, respectively, and these starter units represented the two most highly preferred substrates for both enzymes. No activity was observed for either ARS1 or ARS2 when benzyol-CoA, isovaleryl-CoA, or isobutyryl-CoA was provided as a starter unit.

The recombinant PKS-like protein encoded by root hair contig 0_1848 did not exhibit activity with any of the substrates analyzed, despite associating with the soluble fraction of the *E. coli* protein extracts and migrating at the expected position in SDS-PAGE analysis. Presumably, 0_1848 encodes a nonfunctional protein, or the enzyme does not act on any of the substrates used in this work. By contrast, given that the highly expressed ARS1 and ARS2 sequences encode functional PKS enzymes, it is likely that the corresponding gene products for both are active in planta. As discussed further below, a comparison of the predicted 0_1848 ORF sequence with other type III plant PKS enzymes did indeed reveal differences in several highly conserved positions that could potentially account for its observed inactivity.

Evaluation of steady state kinetic parameters for reaction of ARS1 and ARS2 with the starter units palmitoyl-CoA (C16) and palmitoleoyl-CoA (C16:1), as well as the malonyl-CoA extender...
hairs, each accounting for (see Methods). For ARS1, the triketide pyrone constituted identification of these derailment products as triketide pyrones revealed characteristic parent and fragment ions supporting the mass spectra from the corresponding enzymatic assays re-

longer than C14. Analysis of the total ion chromatograms and myristoyl-CoA (C14), which was not detectable for substrates using saturated acyl-CoA substrates from hexanoyl-CoA (C6) to product was observed for ARS1 and ARS2 in enzyme assays Katsuyama et al., 2009; Taura et al., 2009). A single derailment (e.g., Jez et al., 2000; Liu et al., 2003; Abe et al., 2005b; Katsuyama et al., 2009; Taura et al., 2009). A single derailment product was observed for ARS1 and ARS2 in enzyme assays using saturated acyl-CoA substrates from hexanoyl-CoA (C6) to myristoyl-CoA (C14), which was not detectable for substrates longer than C14. Analysis of the total ion chromatograms and mass spectra from the corresponding enzymatic assays re-

Importantly, both recombinant ARS1 and ARS2 enzymes used hexadecatrienyl-CoA (C16:3,9,12,15), the physiological substrate proposed for ARSs participating in sorgoleone biosynthesis (Figure 1); however, it was somewhat unexpected that higher activities were obtained for both enzymes using substrates such as palmitoyl-CoA (C16) and palmitoleoyl-CoA (C16:1) (Figure 3). While alternative explanations cannot be excluded, several points would suggest that the preferred substrates identified for the recombinant enzymes using in vitro assays may not necessarily reflect the primary activities of ARS1 and ARS2 in planta. For example, the C16 and C16:1 starter units would be predicted to generate congeners of sorgoleone possessing C15:0 and C15:1\textsuperscript{a} alkyl side chains, respectively, in vivo. While these congeners have in fact been identified in sorghum root exudates, they represent only minor constituents (Kagan et al., 2003), despite the fact that palmitate (C16:0) and palmitoleate (C16:1) are far more abundant than C16:3 fatty acids in this cell type. Given that the ARS1 and ARS2 transcripts represent the two most highly expressed PKS-like sequences in S. bicolor root hairs, each accounting for \textasciitilde 0.055\% of all transcripts based on EST counts (Baerson et al., 2008a), and no other PKS-like sequences were identified among the 5468 root hair ESTs analyzed, it would seem unlikely that these enzymes are dedicated solely to the purpose of synthesizing minor exudate constituents. Moreover, both recombinant enzymes also exhibited higher activities in vitro with substrates such as lauroyl-CoA (C12) and myristoyl-CoA (C14), yet C12 and C14 fatty acids are not present in either roots or isolated root hairs of S. bicolor (Pan et al., 2007). A similar nonphysiological preference for short-chain substrates was found with recombinant O-methyltransferase (OMT3) catalyzing the subsequent step in sorgoleone biosynthesis (Baerson et al., 2008a). As was proposed for the latter case, intracellular compartmentation mechanisms, such as metabolon formation, could facilitate selective substrate utilization and the channeling of biosynthetic intermediates, thereby directing pathway flux toward the production of pentadecatrienyl resorcinol (Achnine et al., 2004; Jørgensen et al., 2005). Such a scenario would seem likely given that several cellular compartments are involved in sorgoleone biosynthesis: the hexadecatrienyl-moiety is synthesized by plastidic and microsomal enzymes, while the ARS proteins are cytoplasmic, like all other known type III PKS enzymes (Fate and Lynn, 1996; Austin and Noel, 2003; Dayan et al., 2003; Pan et al., 2007).

RNA Interference–Mediated Repression of ARS1 and ARS2

To evaluate further the potential role for ARS1 and ARS2 in sorgoleone biosynthesis, hairpin RNA-forming binary vectors were assembled for targeted downregulation of ARS1 and ARS2 (see Supplemental Figure 1 online; Wesley et al., 2001). For these constructs, 602- and 556-bp target regions spanning portions of the 3’ coding regions and untranslated regions of ARS1 and ARS2, respectively, were cloned in both sense and antisense orientation, separated by a 1.13-kb intron spacer derived from the FAD2 gene of Arabidopsis (Okuley et al., 1994; see Supplemental Figure 1 online). To minimize the possibility of off-target silencing of related sequences, all publicly available genomic and EST sequence data for S. bicolor were analyzed to avoid target sequences containing regions possessing \textasciitilde 21 nucleotides of contiguous identity with other gene coding sequences (Xu et al., 2006). It is important to take into account that the sequences for ARS1 and ARS2 are closely related, sharing 93\% overall nucleotide sequence identity within coding regions, and within the chosen RNA interference (RNAi) target regions share \textasciitilde 80\% identity, including numerous contiguous stretches \textasciitilde 21 nucleotides in length of 100\% identity. Thus, in principle, hairpin RNA (hpRNA) generated from either vector would be inhibitory to the expression of both ARS1 and ARS2; however, given the paucity of information concerning RNAi-mediated inhibition in sorghum, both vectors were tested in stable transformation experiments using S. bicolor genotype Tx430 (Howe et al., 2006).

Initially, eight independently transformed S. bicolor events (four pARS1-RNAi transformants and four pARS2-RNAi transformants) were selected to evaluate the effects of downregulation of ARS1/2 on sorgoleone biosynthesis in planta, using segregating T1 seedlings. Kanamycin-resistant individuals wherein expression of the RNAi cassette failed to occur would be expected to accumulate sorgoleone at normal levels, and their inclusion in correlative studies could complicate the interpretation of any data obtained. We therefore prescreened individual 10-d-old seedlings grown in perlite culture (see Supplemental Methods online) using real-time PCR to directly

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<th>Table 1. Kinetic Parameters for Recombinant Sorghum ARS1 and ARS2 with C16:0 and C16:1 Starter Units and Malonyl-CoA Extender</th>
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Data are expressed as mean ± SD from assays performed in triplicate.
confirm expression of the hpRNA-generating transgene in root tissues, and then based on these results, pooled the root tissue samples into hpRNA “+” or hpRNA “−” pools for each event. The same “+” and “−” samples from each event were subsequently used for RNA extraction and analysis of sorgoleone accumulation levels by GC-MS. To distinguish the ARS1/2-targeting hpRNA from endogenous ARS1 and ARS2 transcripts, the reverse primer used for real-time hpRNA detection was complementary to sequences within the octopine synthase polyadenylation region immediately adjacent to the RNAi target sequences in pARS1-RNAi and pARS2-RNAi (see Supplemental Figure 1 online). For two of the events analyzed, hpRNA expression was not detected in any individuals; therefore, the remaining six events were used for detailed follow-up studies.

The results for the analysis of the remaining transgenic events in which hpRNA expression was detected are shown in Figure 4. The expression levels for endogenous ARS1 and ARS2 transcripts in “+” and “−” individuals were independently assayed for each event by quantitative real-time RT-PCR using ARS1- and ARS2-specific primers as shown in Figure 4A. In all events, ARS1 and ARS2 expression levels were substantially reduced in hpRNA “+” individuals relative to hpRNA “−”, reflecting the successful downregulation of ARS1 and ARS2 in hpRNA-expressing individuals. ARS1 silencing appeared somewhat more effective overall than ARS2; however, sufficient sequence identity ostensibly existed between the 3’ coding regions of the two genes to trigger silencing of both, as this was observed irrespective of the vector used (Figure 4A). Reduction of ARS1 transcript accumulation in hpRNA “+” individuals ranged from 66 to 96% relative to corresponding hpRNA “−” individuals for each event, and ARS2 transcript levels were reduced from ~55 to 86% (Figure 4A). Complete loss of ARS1/2 expression was not observed for any event; however, relatively few studies have employed real-time RT-PCR to quantify target inhibition in plant RNAi studies; thus, it is somewhat difficult to draw direct comparisons at present. Additionally, DNA gel blot analyses, performed to estimate the number of T-DNA loci in transformants, indicated approximately one to two T-DNAs per event, with three of the six events (events 3, 4, and 6) harboring a single T-DNA locus (Figure 4B).

Sorgoleone levels were also determined for hpRNA “+” and “−” individuals by GC-MS from all events, as shown in Figure 4C. Overall, a complete correlation was observed between hpRNA expression and a dramatic reduction in sorgoleone accumulation. In all cases where hpRNA expression was detected (“+” samples, Figure 4C), sorgoleone levels were reduced to amounts below the limit of quantitation of the GC-MS analysis employed (~0.003 μg/mg fresh weight). Importantly, this trend was observed in six independently transformed events (Figure 4B), thus establishing that the observed reduction in sorgoleone accumulation was dependent on the expression of the hpRNA-generating transgene and was not transformation event-specific. Taken together, the results obtained from ARS1/2-targeting RNAi experiments (Figures 4A to 4C), enzymatic assays using recombinant ARS1 and ARS2 (Figure 3), and the tissue-specific expression pattern determined for ARS1 and ARS2 (Figure 2B) strongly suggest that ARS1 and ARS2 represent the ARS enzymes proposed for the biosynthesis of sorgoleone.

**Phylogenetic Analysis and Functional Identification of Rice Homologs**

A phylogenetic tree was constructed from 72 representative type III PKSs from various plant families, including functionally characterized enzymes accepting a diversity of CoA thioester starter units (Figure 5). The stilbene carboxylate synthase 2 (STCS2) enzyme from liverwort (*Marchantia polymorpha*) was chosen to represent the outgroup for this analysis. Overall, these results indicated separate clustering of CHS and non-CHS type III PKSs among the diverse angiosperm taxa represented, indicating that the divergence of these enzyme families predates the emergence of angiosperms, as previously suggested by Jiang et al. (2008). Additionally, the close relationships between specific CHS- and non-CHS-type enzymes in genera such as *Pinus*, *Arachis*, and *Sorghum* strongly suggest that the non-CHS-type enzymes are likely polyphyletic in origin, as has also been previously proposed (e.g., Tropf et al., 1994; Huang et al., 2004; Jiang et al., 2008).

Interestingly, ARS1 and ARS2 fall within a clade of monocotyledonous type III PKSs that includes predicted sequences from rice whose functions are presently obscure and that is clearly separated from a second monocotyledonous clade containing predominantly CHS-type enzymes (Figure 5). Given that rice is known to synthesize alkyresorcinols thought to function as antimicrobial defense compounds (e.g., Suzuki et al., 1998, 2003), it is tempting to speculate that at least a subset of the predicted rice PKS-like sequences closely related to ARS1 and ARS2 could also possess alkyresorcinol synthase activity. To explore this possibility, the predicted ORFs encoded by *O. sativa* (cv Nipponbare) LOC_Os05g12180, LOC_Os10g08620, and LOC_Os10g07040 (corresponding to XP_476153, NP_920020, and NP_001064197, respectively, in Figure 5) were also expressed in *E. coli* as N-terminal polyhistidine fusions and tested for ARS activity with the same panel of fatty acyl-CoA substrates used for recombinant ARS1 and ARS2 enzymatic studies (Figure 3).

As shown in Figure 6, all three recombinant rice enzymes accepted various saturated and unsaturated fatty acyl-CoA starter units to produce the corresponding 5-alkylresorcinols and, as was observed for recombinant ARS1 (Figure 3), exhibited maximal 5-alkylresorcinol-forming activity with myristoyl-CoA (C14). Among the unsaturated acyl-CoA starters tested, maximal activity was observed with palmitoleoyl-CoA (C16:1Δ9) for all three rice enzymes, as was also seen for both ARS1 and ARS2. No activity was detected for Os05g12180, Os10g08620, and Os10g07040 with the starters hexanoyl-CoA (C6) and arachidonoyl-CoA (C20:4Δ5,8,11,14), potentially indicating that a wider range of fatty acyl-CoA starters can be accepted by the *S. bicolor* alkyresorcinol synthase enzymes. The maximal activities obtained for recombinant Os05g12180 and Os10g08620 using the preferred myristoyl-CoA substrate (66.6 and 48.6 pkat mg⁻¹, respectively) were comparable to those observed for ARS1 with myristoyl-CoA and ARS2 with palmitoleoyl-CoA (81.0 and 60.1 pkat mg⁻¹, respectively); however, lower activity levels were observed for recombinant Os10g07040 with myristoyl-CoA (13.2 pkat mg⁻¹).

As was the case for ARS1 and ARS2, recombinant Os05g12180, Os10g08620, and Os10g07040 each generated a
single derailment product from specific saturated fatty acyl-CoA starters, which were also identified as triketide pyrones by analysis of the total ion chromatograms and corresponding mass spectra (see Methods). For Os05g12180, triketide pyrone derailment products were produced in assays using C8, C10, and C12 fatty acyl-CoAs, constituting ~7, 11, and 8% of the total moles of product derived from these starters, respectively. For Os10g08620, the triketide pyrone derailment products were produced in assays using C8 and C10 fatty acyl-CoA starters, constituting ~8 and 17% of the total moles of product formed, respectively. For Os10g07040, the derailment products were produced in assays containing C8, C10, C12, and C14 fatty acyl-CoA starters and constituted ~8, 20, 6, and <5% of the total moles product formed, respectively (see Methods). Taken together, the results summarized in Figure 6 strongly suggest a role for Os05g12180, Os10g08620, and Os10g07040 in the biosynthesis of rice alkylresorcinol phytoanticipins; furthermore, these enzymes likely serve analogous functions given the overall similarity of their substrate utilization profiles.

Molecular Modeling Suggests a Basis for the Observed In Vitro Substrate Preferences

To examine the potential structural basis for the observed substrate specificity of ARS1 and ARS2, the crystal structures of M. sativa CHS2 (Ferrer et al., 1999) and G. hybrida 2-PS (Jez et al., 2000) were used as templates to create a model of their hypothetical structures. The overall three-dimensional structures of ARS1 and ARS2 were very similar to CHS2 and 2-PS; however, more detailed analyses directed toward the enzyme active sites revealed several differences that could account for the observed substrate preference of these enzymes. Figure 7 shows a simplified model of the proposed active site pointing out these residues, which are also indicated in the sequence alignment (Figure 8) that includes the S. bicolor and O. sativa ARS sequences characterized in this work.

One potentially critical difference is the presence of a Tyr residue in the S. bicolor and O. sativa ARS enzymes, at the position corresponding to Thr-132 in M. sativa CHS2 and Thr-137 in G. hybrida 2-PS. This residue is in direct contact with the active site cavity. A Thr in this position is highly conserved among plant type III PKS enzymes, and the presence of a bulky Tyr in the S. bicolor and O. sativa ARSs is unique in all functionally characterized enzymes in this family. The substitution of a much larger residue suggests a constriction or narrowing of the active site cavity within this region.

An additional substitution that could potentially influence the substrate preferences of the S. bicolor and O. sativa ARSs is the presence of an Ala residue in their sequences at the position

![Figure 4](image-url)

**Figure 4.** Evaluation of S. bicolor RNAi Transformant Events.

(A) Relative ARS1 and ARS2 endogenous transcript levels in 10-d-old S. bicolor hprNA “+” and hprNA “−” seedlings (representing six independent transformant events) were determined by quantitative real-time RT-PCR using gene-specific primers. Data were normalized to an internal control (18S rRNA), and the ΔΔCT method was used to obtain the relative expression levels for each sequence, expressed as mean ± SD from assays performed in triplicate. Events numbered 1, 3, and 4 were generated using the vector pARS1-RNAi, and events numbered 2, 5, and 6 were generated using pARS2-RNAi (see Supplemental Figure 1 online).

(B) Genomic DNAs isolated from the six S. bicolor RNAi transformant events, and control seedlings (genotype Tx430) were digested with either BamHI or SphI and then probed using radiolabeled Arabidopsis thaliana FAD2 gene intronic sequences present in the RNAi cassette. C, control; B, BamHI; S, SphI.

(C) Sorgoleone levels were determined by GC-MS analysis of root exudates prepared from 10-d-old hprNA “+” and hprNA “−” seedlings representing the six RNAi transformant events. Data are expressed as mean ± SD from four measurements. The limit of quantitation (LOQ), determined to be ~0.003 μg/mg fresh weight, is also indicated by a dashed line.
Figure 5. Phylogenetic Relationships of ARS1, ARS2 Relatives.

Strongly supported nodes (posterior probability >0.95) are indicated by shaded circles. Shaded boxes are included to highlight the placement of S. bicolor CHS and non-CHS-type sequences, and the bar at bottom represents the distance corresponding to 0.2 substitutions per amino acid. The M. polymorpha stilbene carboxylate synthase 2 (STCS2) sequence served as the outgroup for this analysis. ACS, acridone synthase; ALS, aloesone synthase; BAS, benzalacetone synthase; BBS, bibenzyl synthase; BPS, benzophenone synthase; CHS-LK, chalcone synthase-like (unknown function); CURS, curcumin synthase; DCS, diketide CoA synthase; OKS, octaketide synthase; OLS, olivetol synthase; PCS, pentaketide chromone synthase; PSS, pinosylvin synthase; STCS, stilbene carboxylate synthase; VPS, valerophenone synthase.
corresponding to Met-137 in *M. sativa* CHS2 and Met-142 in *G. hybrida* 2-PS. The majority of type III PKS enzymes contain a Met in this position, which provides the sole contribution of the second subunit to the opposing subunit’s active site cavity within the PKS homodimer (reviewed in Austin and Noel, 2003). The Ala found in the corresponding position of the *S. bicolor* and *O. sativa* ARS enzymes is rare among type III PKSs. As discussed below within the context of aldol condensation-based cyclization (Figure 9), it is of interest that bacterial proteins using long-chain fatty acyl-CoA as substrates also contain small residues in this position, either Ala or Ser. PKS18, a pyrone synthase from *Mycobacterium tuberculosis*, also contains an Ala in this position, and the simultaneous substitution of Met-137 → Ala and Gly-256 → Leu (see also discussion below) appears to be an important factor in defining the tunnel necessary for accepting fatty acyl-CoA substrates (Sankaranarayanan et al., 2004). Taken together, the substitution of this Met for the much smaller Ala residue in the *S. bicolor* and *O. sativa* ARS enzymes appears likely to play a significant role in determining their preferences for long- to medium-chain acyl-CoA substrates.

Of further interest is the presence of an Ala residue in the *S. bicolor* ARSs and a Cys residue in the *O. sativa* ARSs, at the position corresponding to Thr-197 in *M. sativa* CHS2 and Leu-202 in *G. hybrida* 2-PS. Prior studies indicate that this position appears to be of particular significance for determining the accepted substrate size and extent of the active site cavity in type III PKSs. Several important examples where Thr-197 is replaced in the corresponding position by other residues in non-CHS-type PKSs include aloesone synthase from *Rheum palmatum* (Thr → Ala; Abe et al., 2004, 2006) and several enzymes from *Aloe arborescens*, including pentaketide chromone synthase (Thr → Ala; Abe et al., 2005b, 2007; Morita et al., 2007), octaketide synthase (Thr → Gly; Abe et al., 2005a), and a second aloesone synthase (Thr → Ala; Mizuuchi et al., 2009). A large number of in vitro studies performed with these enzymes have shown that single amino acid substitutions at this position drastically alter the size of the active site cavity and resulting products formed. Specifically, an inverse relationship has been demonstrated between residue side chain bulk and the number of condensation reactions performed, thus determining the size of the resulting products (Abe et al., 2004, 2005b, 2006, 2007; Morita et al., 2007; Mizuuchi et al., 2009). It therefore seems likely that the corresponding Ala in the *S. bicolor* ARSs and Cys in *O. sativa* similarly contribute to the size of their respective active site cavities.

The Met residue present in the *S. bicolor* and *O. sativa* ARSs, at the position corresponding to Gly-256 in *M. sativa* CHS2 and Leu-261 in *G. hybrida* 2-PS, is also likely to influence the types of starter units accepted by these enzymes. The importance of this amino acid position for determining the shape and size of the *M. sativa* CHS2 and *G. hybrida* 2-PS active sites has long been recognized, and site-directed mutagenesis studies where CHS2 Gly-256 was replaced with residues containing bulkier side chains resulted in altered substrate utilization and a reduction in the number of condensation reactions performed (reviewed in Austin and Noel, 2003). In fact, in studies performed by Jez et al. (2000), it was demonstrated that a Thr197Leu/Gly256Leu/Ser338Ile triple mutant of *M. sativa* CHS2 was sufficient to convert the CHS into a pyrone synthase that would no longer accept bulky phenylpropanoid starter units. An additional noteworthy example is the *M. tuberculosis* PKS18 enzyme (discussed above within the context of CHS2 Met-137 substitutions), which contains the much bulkier Leu residue at this position. The substitution of the bulkier Met residue for Gly-256 in all of monocotyledonous ARS enzymes characterized in this work would be anticipated to play a role similar to the corresponding Leu substitution in the *M. tuberculosis* enzyme.

Taken together, these analyses predict that the active sites of the *S. bicolor* and *O. sativa* ARS enzymes are narrower near the upper region adjacent to the catalytically important conserved

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**Figure 6.** Enzymatic Activities of ARSs Encoded by *O. sativa* LOC_Os05g12180, LOC_Os10g08620, and LOC_Os10g07040. Relative 5-alkylresorcinol–forming activities were determined for all three recombinant enzymes in assays using acyl-CoA starter units varying in chain length and degree of saturation. Triketide pyrone derailment products were also produced by Os05g12180 in assays containing C8 to C12 acyl-CoA starter units, by Os10g08620 in assays containing C8 and C10 starter units, and by Os10g07040 in assays containing C8 to C14 acyl-CoA starter units. The corresponding arylresorcinol products represented between ~80 and 95% of the total moles product formed in these cases. Data are expressed as relative mean ± SD from assays performed in triplicate. The 100% relative activity corresponds to 66.6 pkat mg⁻¹ for Os05g12180, 48.6 pkat mg⁻¹ for Os10g08620, and 13.2 pkat mg⁻¹ for Os10g07040, defined as pmol of alkyresorcinol formed/s per mg of protein.
identified for the S. bicolor critical positions (see above discussion) from the consensus starters units analyzed, 0_1848 deviates in several potentially substrates tested in this work. With respect to the fatty acyl-CoA potential reasons for the lack of activity observed with the various ORF with other type III plant PKS enzymes (Figure 8) did suggest (Goyal et al., 2008). starter fatty acyl-CoA chain length specificity for this enzyme similar hydrophobic active site tunnel is involved in determining the various resorcinolic metabolites, have demonstrated that a sim-

Figure 7. Molecular Modeling of ARS1 and ARS2 Active Sites. The three-dimensional active site structures of M. sativa CHS2 and G. hybrida 2-PS, as well as homology models of the ARS1 and ARS2 active site structures, are shown as ribbon diagrams. Selected residues contributing to the shape/size of the active site architecture are shown in space-filling representation.

(A) M. sativa (Ms CHS2), (B) G. hybrida 2-PS (Gh 2-PS), (C) and (D) Models for ARS1 and ARS2, respectively.

Cys residue (Cys-164 in M. sativa CHS2) and also are of increased overall length when compared with M. sativa CHS2 and G. hybrida 2-PS (Jez et al., 2000). This is likely to produce a cavity more adequately suited to accommodating long- to medium-chain fatty acyl-CoA starter units (Figure 7). Interestingly, the sharp drop in activity observed for both ARS1 and ARS2 with fatty acyl-CoAs longer than C16 (Figure 3) could indicate that this cavity length becomes limiting for the type of substrates that can be accepted. Additionally, the lack of activity with acyl-CoAs less than C6 (Figure 3) could indicate that substrate filling of this elongated active site cavity is critical to enzyme activity. Consistent with this notion, structural studies recently performed with ORAS, a type III PKS from Neurospora crassa using long-chain fatty acyl-CoAs for the production of various resorcinolic metabolites, have demonstrated that a similar hydrophobic active site tunnel is involved in determining the starter fatty acyl-CoA chain length specificity for this enzyme (Goyal et al., 2008).

Additionally, a comparison of the S. bicolor 0_1848 predicted ORF with other type III plant PKS enzymes (Figure 8) did suggest potential reasons for the lack of activity observed with the various substrates tested in this work. With respect to the fatty acyl-CoA starters units analyzed, 0_1848 deviates in several potentially critical positions (see above discussion) from the consensus identified for the S. bicolor and O. sativa ARS enzymes, for example (numbering based on Ms CHS2), Tyr-132 → Asn, Ala-137 → Ile, and Cys (O. sativa)/Ala (S. bicolor) 197 → Leu. The interpretation for other systematic differences is unclear at present (e.g., His-162 → Asn, Gly-211 → Asn, and Phe (O. sativa)/Tyr (S. bicolor) 265 → Gly); however, the exchange of the highly conserved Ser-338 for Phe in 0_1848 could be highly significant, given that Ser-338 represents a key component of the aldol switch mechanism identified from studies with the STS1 enzyme from Pinus sylvestris (Austin et al., 2004a; discussed in the following section). Additionally, the lack of activity seen with the substrates benzoyl-CoA, isobutyryl-CoA, and isovaleryl-CoA suggests that 0_1848 does not represent a CHS-, STS-, 2-PS-, or VPS-type PKS, since these enzymes typically do exhibit significant levels of activity with these nonphysiological substrates.

Aldol Condensation Mechanisms of ARSs and Other Type III PKS Enzymes

For ARS proteins, cyclization is performed via a C2→C7 aldol condensation similar to that used by STS-type PKSs, in contrast with the C6→C1 Claisen condensation associated with CHS-type enzymes (Tropf et al., 1994; Funahashi et al., 2006, 2007; Goyal et al., 2008). The mechanisms underlying the different cyclization reactions performed by type III PKS enzymes have been of considerable interest, and a detailed structural/mutagenesis study performed with P. sylvestris STS1 and M. sativa CHS2 by Austin et al. (2004a) succeeded in generating a functional STS-type enzyme from CHS2 by site-directed mutagenesis. The results of Austin et al. further revealed that the primary underlying structural determinants for the different cyclization functions associated with CHS2 and STS1 reside within a small displaced loop within STS1 (designated as the area 2 loop), corresponding to positions Thr-131 to Met-137 in the CHS2 sequence. In STS1, this region facilitates the formation of a hydrogen bond network between Thr-132, Glu-192, and Ser-338 (numbering according to M. sativa CHS2) not present in CHS2. This results in the activation of a cryptic thioesterase activity leading to the release of the tetraketide intermediate, thus initiating the decarboxylation/cyclization characteristic of the STS reaction. This mechanism has been referred to as the aldol switch and was successfully transferred to the M. sativa CHS2 enzyme through the exchange of just eight amino acids, resulting in the generation of a fully functional STS-type enzyme (Austin et al., 2004a). The five most critical amino acid exchanges were made within the area 2 loop, while the remaining three exchanges were compensatory and located in protein domains referred to as region 1 and region 3. A comparison of the corresponding positions for area 2, as well as other key residues associated with the aldol switch mechanism, are shown for various microbial and plant type III PKS enzymes in Figure 9. Whenever possible, a CHS- and a STS-type sequence from the same species were included in the alignment to facilitate these comparisons. For simplification, all amino acid residues will be referred to in the discussion below based on the equivalent M. sativa CHS2 positions.

Analysis of the STS- and CHS-type PKSs from Pinus spp revealed identical residues for area 2 and positions 192 and 338

Figure 8. Examples of exchange mutations designed to determine the critical residues associated with cyclization by ARSs, CHSs, and STSs. For each enzyme, the sequence was altered by the exchange of the residues determined to be critical for cyclization (shown in black). One representative sequence alignment is shown for each enzyme (numbering based on Ms CHS2).
Figure 8. Alignment of S. bicolor and O. sativa Sequences Exhibiting ARS Activity.

Alkylresorcinol Synthases from Sorghum
for the respective STS- or CHS-type enzymes (with the exception of a Ser → Thr exchange in position 131 of the *P. strobos* STS1), suggesting a conservation of the aldol switch and similar mechanistic differences occurring between STS- and CHS-type enzymes in gymnosperms (Figure 9). By contrast, the signature characteristic for the aldol switch-type mechanism is not recognizable in the *Arachis hypogaea* STS3 and *Vitis vinifera* STS sequences; however, a preliminary analysis of their three-dimensional structures did suggest that other residues are responsible for carrying out a similar mechanism in these cases (Austin et al., 2004a). Additionally, the bibenzyl synthase—type PKS area 2 sequences from *Bromheadia finlaysonia* and *Phalaeonopsis* spp show similarities to the A. hypogaea STS3 and *V. vinifera* STS sequences in this region, suggesting a similar mechanism for these enzymes.

The *Sorbus aucuparia* biphenyl synthase (BIS) and the *Cannabis sativa* olivetol synthase (OLS) sequences both contain an Ala instead of the aldol switch-type Thr-132 within area 2. Additionally, the BIS sequence has a Gly in place of Ser-338, and OLS has an Asp residue in place of Glu-192. As discussed above, all three positions play an essential role in the aldol switch hydrogen-bonding network, suggesting that these STS-type enzymes use an alternative aldol condensation cyclization mechanism (this is also likely true in the case of the *Rheum tataricum* STS1 enzyme, which contains an Ile at position 132). The stilbene carboxylate synthases (STCSs) from *Hydrangea macrophylla* and from the liverwort *M. polymorpha* are unique with respect to the other type III PKS enzymes shown in Figure 9, in that they use an aldol condensation—type cyclization mechanism but retain the terminal carbonyl group in the end product (Eckermann et al., 2003; http://www.biologie.uni-freiburg.de/data/bio2/schroeder/STCS_Marchantiaceae_polymorpha.html). Their sequences within area 2 are identical to that of typical CHS-type PKSs, and the three-dimensional structure of the *M. polymorpha* STCS2 enzyme (PDB: 20UA) revealed no significant differences when compared with that of *M. sativa* CHS2 within the area 2 loop or other regions associated with aldol switch function. The *M. polymorpha* cyclization mechanism must therefore differ from the aldol switch type, and it has been proposed that the aldol condensation in resorcylic acid synthases, such as STCS, is a nonenzymatic reaction from the released tetraketide pyrone (Austin et al., 2004a). For the two STS-type sequences from the primitive fern *Psilotum nudum*, it is difficult to predict the underlying mechanism for the aldol cyclization performed solely based on primary sequence data (STS PnL contains a gap in the area 2 sequence, and both PnL and PnL contain a unique Pro134 residue), and this is also the case for a putative ARS/apyrone synthase identified from the moss *Physcomitrella patens* (Jiang et al., 2008).

The single fungal type III PKS representative shown in Figure 9 from *N. crassa* has been alternatively described as an oxoalkylresorcylic acid synthase and an ARS by two groups working independently. Funa et al. (2007) first identified 2-oxoalkylresorcylic acid synthases (derived from four condensations) as the major in vitro products and thus named the enzyme ORAS, whereas Goyal et al. (2008) identified 5-alkylresorcinols (derived from three condensations) as the major products formed. Crystal structures were independently established by the two groups, and although their interpretations differ in some details, both studies concluded that an aldol switch-type cyclization mechanism is not involved due to the presence of a Cys residue in the position equivalent to Thr-132 in the *M. sativa* CHS2 enzyme (Figure 9), which would preclude the formation of a thioesterase-like hydrogen bonding network (Goyal et al., 2008; Rubin-Pitel et al., 2008). The significance of this conclusion, however, could depend on the products identified: alkylresorcinolic products would suggest an alternative mechanism for aldol condensation, whereas alkylresorcylic acid products would not require that. According to the model previously proposed for resorcylic acid biosynthesis, the formation of such molecules is likely a nonenzymatic reaction from released pyrones (Austin et al., 2004a).

Of particular interest within the context of cyclization specificity is the enzyme 1,3,6,8-tetrahydroxynaphthalene synthase (THNS), a bacterial type III PKS present in various *Streptomyces* strains. These enzymes perform four condensation reactions resulting in a linear pentaketide intermediate and use both a Claisen and an aldol condensation for the formation of the end product. The THNS from *Streptomyces coelicolor* has been crystallized, and the analysis unexpectedly revealed that position 106, which corresponds to Thr-132 in *M. sativa* CHS2, contains a Cys residue that plays an important catalytic role in facilitating polyketide extension beyond the triketide stage (Austin et al., 2004b). Another important conclusion drawn from that work, based on the enzyme’s three-dimensional structure, is the observation that the *S. coelicolor* THNS active site does not possess an aldol switch, as is also suggested by the replacement of the highly conserved Ser-338 by Ala (Figure 9).

Three bacterial type III PKS enzymes have also been shown to possess ARS activity: SrsA from *Streptomyces griseus* (Funabashi et al., 2008), CHS-LK from *Synechococcus* (Jiang et al., 2008), and ArsB from *Azotobacter vinelandii* (Funa et al., 2006). SrsA contains a Val at the critical 132 position of area 2, and the *Synechococcus* and *A. vinelandii* enzymes both contain a Cys residue at this position, as noted above for the *N. crassa* ORAS enzyme. Additionally, the otherwise strictly conserved Glu-192 is replaced by Asp in ArsB and ArsC, as was also noted above for the *C. sativa* OLS enzyme. All three bacterial enzymes are

Figure 8. (continued).

The sequences for *S. bicolor* ARS1 and ARS2 and the ARSs encoded by *O. sativa* LOC_Os05g12180, LOC_Os10g08820, and LOC_Os10g07040 were aligned with *M. sativa* CHS2, *G. hybrida* 2-PS, *S. bicolor* CHS2, *S. bicolor* g_1848, and *O. sativa* CHS1 using ClustalW. Residues associated with PKS functional diversity, catalysis (catalytic triad), and CoA binding are indicated based on previous crystallography studies (Ferrer et al., 1999; Jez et al., 2000) and by computational homology modeling of ARS1 and ARS2. Numbering shown above the catalytic triad positions, as well as several key residues potentially contributing to active site architecture, is based on the *M. sativa* CHS2 sequence. Also indicated by boxes are atypical residues identified within the g_1848-encoded polypeptide that could account for the lack of enzymatic activity observed in recombinant enzyme studies.
therefore unlikely to possess the classical (STS1-like) aldol switch mechanism. Within the Azotobacter ars operon, a gene encoding a second type III PKS was also identified (ArsC) that shares 71% amino acid identity with ArsB (Funa et al., 2006). Like ArsB, ArsC was also shown to accept various fatty acyl-CoA starter units; however, in contrast with ArsB, the major in vitro products generated by ArsC were found to be tetraketide pyrones. While both enzymes generate an identical tetraketide intermediate from three condensation reactions, cyclization of the ArsC intermediates would be predicted to occur via intramolecular C5...

Figure 9. Alignment of Key Sequence Motifs Underlying the Aldol Switch Cyclization Mechanism.

A partial alignment was generated using functionally diverse type III PKS enzymes from various sources, highlighting regions critical to the aldol switch (STS-type) cyclization mechanism first described by Austin et al. (2004a). For simplicity, numbering is shown according to the *M. sativa* CHS2 sequence. Positions 131 to 137 correspond to the displaced area 2 loop (outlined by rectangle) critical for the formation of the hydrogen bond network formed by Thr-132, Glu-192, and Ser-338 (highlighted in blue) in *P. sylvestris* STS1. Only three residues in area 2 (positions 132, 133, and 137; indicated by closed circles) are in direct contact with the active site cavity. The five area 2 positions mutated in *M. sativa* CHS2 to generate a functional STS-type enzyme (Austin et al., 2004a) are highlighted in yellow. Deviations from the consensus in positions Thr-132, Glu-192, and Ser-338 are boxed.

Therefore, in contrast with ArsB, the major in vitro products generated by ArsC were found to be tetraketide pyrones. While both enzymes generate an identical tetraketide intermediate from three condensation reactions, cyclization of the ArsC intermediates would be predicted to occur via intramolecular C5...
oxygen → C1 lactonization rather than the aldol condensation used by ArsB and other ARS enzymes. The diagnostic area 2 sequences found in ArsB and ArsC are identical with the exception of position 137 (Ser in ArsB; Ala in ArsC), which represents one of the three residues contacting the second subunit within the PKS homodimer (Figure 9). Thus, it is unlikely that this region is responsible for differences in the products produced by these enzymes (i.e., alklyresorcinol versus tetraketide alky/lyprone).

The S. bicolor and O. sativa ARSds described in this work are unique with respect to their area 2 sequence composition; for example, in the position equivalent to Thr-132 in M. sativa CHS2, a Tyr is found in all five enzymes (Figure 9). This residue was mentioned above within the context of enzyme substrate preference; however, its presence in this position also raises questions concerning whether these ARSs possess a STS1-like aldol switch mechanism. Both the position and size of this residue, as well as the pK_a value of its hydroxyl group (pK_a ∼ 10 versus 15 for Thr), would suggest that the STS1-type cyclization model is not applicable to these enzymes. However, in the absence of crystal structures, it is not possible to draw definitive conclusions in this regard, in view of the minor conformational differences observed between CHS- and STS-type enzymes in those cases analyzed in detail (e.g., Austin et al., 2004a, 2004b; Goyal et al., 2008; Rubin-Pitgel et al., 2008). Interestingly, for the two additional functionally tested PKS enzymes from S. bicolor, one of which is a typical CHS enzyme (CHS2) and the second a STS (STS1; Yu et al., 2005), their sequences within the critical area 2 are identical; thus, an alternative aldol condensation mechanism is likely used by S. bicolor STS1 as well (Figure 9).

The results described in this work provide evidence arguing in favor of a role for ARS1 and ARS2 in the sorgoleone biosynthetic pathway as well as roles for the ARS enzymes identified from O. sativa in the biosynthesis of defense-related alklyresorcinols in this species. The ARS sequences will undoubtedly serve as valuable tools for the detailed analysis of alklyresorcinol biosynthetic pathways from other plants, particularly those identified in other cereals such as rye (Secale cereale), wheat, and barley (Hordeum vulgare; reviewed in Kozubek and Tyman, 1999; Kozubek et al., 2001) where the alklyresorcinol synthases involved would be anticipated to share a high degree of sequence identity with the enzymes identified in this work.

METHODS

Plant Material and Growth Conditions

Seeds of Sorghum bicolor genotype BTx623 were purchased from Crosbyton Seed Company, and seeds of S. bicolor genotype Tx430 were harvested from greenhouse-grown plants maintained at the University of Nebraska–Lincoln greenhouse facilities. Construction and analysis of ARS1 and ARS2 RNAi lines are described in Supplemental Methods online. For real-time RT-PCR experiments and 5-pentadecatrayl resorcinol GC-MS analyses, root tissues were obtained from 8-d-old dark-grown BTx623 seedlings grown under soil-free conditions using a capillary mat system devised by Czarnota et al. (2001), and root hairs were isolated from this material in bulk as previously described (Baerson et al., 2008a). Immature leaves and shoot apices were isolated from BTx623 seedlings maintained in a growth chamber at 28°C for 8 d in standard (~20 × 40 cm) nursery flats using Premier Pro Mix PGX potting media (Hummer Inte-
assemblies (Baerson et al., 2008a), which served as the basis to obtain full-length ORFs by 5′-RACE. For these experiments, a SMART RACE cDNA amplification kit (Clontech Laboratories) was used per the manufacturer’s instructions with total RNA isolated from *S. bicolor* genotype Btx623 root hairs (described above). PCR amplification products containing the complete ORFs for ARS1, ARS2, and PKS-like 0, 1848 flanked by Ndcl (5′ end) and BamHI (3′ end) restriction sites were then generated by PCR amplification to facilitate direct ligation with Ndcl- and BamHI-digested pET15b (EMD Biosciences). Similarly, the predicted full-length ORFs for the *Orzya sativa* PKS-like sequences encoded by LOC_5_1848 for *N. sativa*, digested with *Nde*I and *BamHI*-D-thiogalactopyranoside and allowed to grow an additional 15 min. Temperature and buffer pH optima for ARS1/2-containing assays were controlled to ensure linearity of the assays. Reactions were quenched by addition of 10 μL of 20% HCl, and products were extracted by phase partitioning with 1 mL of ethyl acetate. The organic phase (upper layer) obtained by centrifugation at ~14,000g for 1 min was transferred to a fresh tube, dried under vacuum, and subsequently analyzed by GC-electrospray ionization (EI)-MS as a TMS derivative. Product formation was quantified using selective ion monitoring at m/z 268, a fragment ion common to all alkyresorcinols, generated by benzylic cleavage of the alkyl side chain. The identification of the m/z 268 ([5-methyl-1,3 phenylene]bi(oxy)bi(TMS)) base ion and the parent [M]+ provide confirmation of the 5-alkyresorcinolic structure as well as the length and degree of saturation of the associated side chain (Occolowitz, 1964; Suzuki et al., 1998, 2003). Spectral data obtained for all identified alkyresorcinolic products are provided in Supplemental Table 1 online. Further confirmation for the formation of olivetol (ARS0) and pentadecylresorcinol (ARS1:0) in assays provided with hexanol- and palmitol-CoA starters was obtained by comparison of product retention times and mass spectra relative to authentic standards. GC-EL-MS: ARS0 = R1 to R5, 0.40 min, m/z 324 [M]+, m/z 309 [M + CH3]+, m/z 295 [5-propyl-1,3-phenylene]bi(oxy)bi(TMS)+, m/z 281 [5-ethyl-1,3-phenylene]bi(oxy)bi(TMS)+, m/z 268 [5-methyl-1,3-phenylene]bi(oxy)bi(TMS)3, ARS1:0 = R10 to 17.17 min, m/z 464 [M]+, m/z 449 [M + CH3]+, m/z 361 [449 + 2(CH3)+], m/z 361 [449 + OTMS, + H]+, m/z 323 [5-pentyl-1,3-phenylene]bi(oxy)bi(TMS) + H]+, m/z 310 [5-buty1-1,3-phenylene]bi(oxy)bi(TMS)+, m/z 310 [5-propyl-1,3-phenylene]bi(oxy)bi(TMS)+, m/z 281 [5-ethyl-1,3-phenylene]bi(oxy)bi(TMS) + H]+, m/z 268 [5-methyl-1,3-phenylene]bi(oxy)bi(TMS) + H]+, m/z 258 [268 + CH3]+.

Enzyme assays, performed in triplicate, for starter unit CoA kinetics contained 100 mM potassium phosphate buffer, pH 7.0, 40 μM [2-14C]-malonyl-CoA (50 to 60 μmol/ml and 1.85 to 2.22 GBq/mmol; American Radiolabeled Chemicals), 1.43 to 10 μM starter unit (e.g., palmitol-CoA), and 0.75 μg protein in a 200 μL volume at 30°C for 5 min. Enzyme assays for malonyl-CoA kinetics contained 100 mM potassium phosphate buffer, pH 7.0, 1.66 to 25 μM [2-14C]-malonyl-CoA, 25 μM starter unit (palmitol-CoA), and 0.75 μg protein in a 200 μL volume at 30°C for 5 min. Reactions were quenched by addition of 10 μL of 20% HCl. The products were extracted as described above and were separated by thin layer chromatography (silica gel 60 F254; chloroform:ethanol = 70:30; 15 min). Product detection and formation were quantified through the use of the Cyclone Chromatography System and OptiQuant 3.0 image analysis software (Perkin-Elmer). Data were fit to the Michaelis-Menten equation using the Sigma Plot 9.01 enzyme kinetics module (Systat). Thin layer chromatographs of ARS1, ARS2, Os05g12180, Os10g08620, and Os10g07040 reaction products in assays using saturated acyl-CoA substrates from hexanol-Co (C6) to myristol-CoA (C14) revealed the formation of secondary products, which were subsequently identified by GC-EL-MS as triketide pyrones from the mass spectrum of the peaks appearing in reconstructed ion chromatograms, supported by the appearance of fragment ions characteristic of the specific triketide pyrone. GC-EL-MS (starter substrate, m/z triketide pyrone): hexanol-CoA (C6), m/z 254 [M]+, m/z 239 [M + CH3]+, m/z 224 [M – CO, 2H]+, m/z 195 [M – CO, 2(CH3) + H]+, m/z 211 [3-OTMS-5-methylfuran, – H]+; caprylol-CoA (C8), m/z 253 [M – CO, – H]+, m/z 211 [M – TMS, +2H]+, m/z 166 [282 + CO, - OTMS]+, m/z 139 [166 + CO, + H]+, capryol-CoA (C10), m/z 311 [M + H]+, m/z 281 [M – CO, – H]+, m/z 221 [M – OTMS]+, m/z 191 [281 – OTMS, – H]+; lauroyl-CoA (C12), m/z 337 [M + H]+, m/z 312 [M – CO, + 2H]+, 325 [M – CH3 + 2H]+, m/z 249 [M – OTMS]+, m/z 221 [249 + CO]+, m/z 197 [221 – CO + H]+; myristol-CoA (C14), m/z 366 [M]+, m/z 351 [M – CH3]+, m/z 221 [M – CO3H – OTMS]+, m/z 207 [221 + CH3]+, m/z 170 [5-OTMS-5-methylfuran]+.
Phylogenetic Analyses

Amino acid sequences of putative type III plant PKSs were retrieved from the NCBI nonredundant peptide sequence database by BLASTP searches using default parameters (http://blast.ncbi.nlm.nih.gov). A candidate list was screened for redundancy and errors, and a final data set was assembled containing 72 sequences including the three S. bicolor and three O. sativa sequences biochemically characterized in this work. Multiple sequence alignments were constructed with GENEIOUS version 4.6.2 (Biomatters) employing the BLOSUM62 log-odds probability matrix (Henikoff and Henikoff, 1993) and gap open and extension penalties of 12 and 3, respectively. Terminal amino acids with <50% coverage were trimmed to yield a final alignment of 409 residues (see Supplemental Data Set 1 online).

Assessment of phylogenetic relationships among gene sequences employed the Bayesian Markov chain Monte Carlo simulation technique implemented in MRBAYES v. 3.1 (Ronquist and Huelsenbeck, 2003). This analytical approach takes advantage of probabilistic models of amino acid substitution and has been shown to be robust to among-site rate heterogeneity and branch length differences (Mar et al., 2005). Two separate analyses, each containing two independent searches, were run for 10^6 generations, sampling every 2000. To incorporate the uncertainty in the appropriate amino acid substitution model, we used mixed priors with ë-distributed rate variation; posterior support for the Wagner model (Whelan and Goldman, 2001) was 1.0. From this posterior sample of trees with an analytical approach takes advantage of probabilistic models of amino acid substitution and has been shown to be robust to among-site rate heterogeneity and branch length differences (Mar et al., 2005). Two separate analyses, each containing two independent searches, were run for 10^6 generations, sampling every 2000. To incorporate the uncertainty in the appropriate amino acid substitution model, we used mixed priors with ë-distributed rate variation; posterior support for the Wagner model (Whelan and Goldman, 2001) was 1.0. From this posterior sample of trees

Molecular Modeling

The previously determined crystal structures of Medicago sativa CHS2 (Ferrer et al., 1999) and Gerbera hybrida 2-PS (Jez et al., 2000) were used as templates for modeling the active site structures for ARS1 and ARS2. Structural models were developed using SWISS-MODEL, the automated protein homology-modeling server (http://www.expasy.org/spdbv/). Structural models were developed using SWISS-MODEL, the automated protein homology-modeling server (http://www.expasy.org/spdbv/). Structures were visualized using the Swiss-Pdb Viewer program (Guex and Peitsch, 1997; Schwede et al., 2003).

Accession Numbers

Sequence data from this article can be found in the Arabidopsis Genome Initiative or GenBank/EMBL data libraries under the following accession numbers: FAD2, AT3g12120; O. sativa LOC_Os05g12180, XP_476153; O. sativa LOC_Os10g08620, NP_920020; O. sativa LOC_Os10g07040, NP_001064197; O. sativa CHS1, BAB39764; M. sativa CHS1, P30074; G. hybrida 2-PS, GAA86219; S. bicolor ARS1 (sequence ID number 2_126 in Figure 2B), XM_002449699; S. bicolor ARS2 (sequence ID number 2_127 in Figure 2B), XM_002441794; S. bicolor CHS1 (sequence ID number 0_1821 in Figure 2B), AAD41873; S. bicolor CHS2, AAD41874; S. bicolor CHS5 (sequence ID number 0_164 in Figure 2B), AAD41877; S. bicolor STS1, AAA49965; and S. bicolor 0_1848 (sequence ID number 0_1848 in Figure 2B), XP_002462898. Accession numbers for the sequences used in the phylogenetic analysis are labeled in Figure 5. Accession numbers are also labeled on the alignment in Figure 9. M. sativa CHS2 structural information can be found in the Protein Data Bank (pdb code 1B1S), as can structural information for G. hybrida 2-PS (pdb code 1QLV).
Supplemental Data

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

Supplemental Figure 1. Binary Vectors Developed for RNAi-Mediated Inhibition of ARS1 and ARS2 Expression.

Supplemental Figure 2. Specificity of PCR Primer Pairs Used for the Detection of Endogenous ARS1 and ARS2 Transcripts in pARS1-RNAi and pARS2-RNAi S. bicolor Transforms.

Supplemental Table 1. Mass Spectral Characteristics of 5-Alkylresorcinols produced in ARS1, ARS2, Os05g12180, Os10g08620, and Os10g07040 Enzymatic Assays.


Supplemental References.

Supplemental Data Set 1. Text File of the Alignment Used for the Phylogenetic Analysis in Figure 5.

ACKNOWLEDGMENTS

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Fato, G.D., and Lynn, D.G. (1996). Xenogosin methylation is critical in defining the chemical potential gradient that regulates the spatial


Supplemental Figure 1. Binary Vectors Developed for RNAi-Mediated Inhibition of ARS1 and ARS2 Expression.

Construction of pARS1-RNAi and pARS2-RNAi are described in Supplemental Methods online. hpRNA-generating cassettes for both vectors were directionally-cloned using asymmetric SfiI restriction sites within the T-DNA borders of the binary vector pLH9000 (Hausmann and Toepfer, 1999), and include (5' to 3'): the constitutive Zea mays polyubiquitin-1 gene promoter and intron 1 [P-ZmUbi-1; (Christensen et al., 1992)], sense and antisense ARSI and ARS2 target regions separated by intron 1 of the Arabidopsis thaliana FAD2 gene [FAD2 int; (Okuley et al., 1994)], and the Agrobacterium tumefaciens octapine synthase gene terminator [OCS-ter; (De Greve et al., 1982)].
Supplemental Figure 2. Specificity of PCR Primer Pairs Used for the Detection of Endogenous \textit{ARS1} and \textit{ARS2} Transcripts in pARS1-RNAi and pARS2-RNAi \textit{S. bicolor} Transformants.

\textit{ARS1}-specific and \textit{ARS2}-specific primer pairs (described in Supplemental Methods online) were tested in quantitative real-time RT-PCR assays to confirm the specific amplification of their intended targets. For these experiments, approximately 1.5 kb PCR products containing the coding sequences and proximal 5'/3' flanking regions for \textit{ARS1} and \textit{ARS2} were used as templates (1 pg template DNA per 25 µL reaction) in reciprocal tests with both \textit{ARS1}- and \textit{ARS2}-specific primer pairs. The primers used for generation of the two PCR templates from \textit{S. bicolor} (genotype BTx623) genomic DNA were as follows: \textit{ARS1} forward, 5'-AGCTAGCTGCTCTCGATCTGTCC-3' , reverse, 5'-TTGCTCAAGAGGAATCGGATTG-3'; and \textit{ARS2} forward, 5'-CCGTACCACAGCTGCTACACTAGC-3', reverse, 5'-GTTTCTCTTATCAACAAGAGGATCG-3'. The change in fluorescence of SYBR® Green I dye in every cycle was monitored by the GenAmp® 7300 Sequence Detection System software (Applied Biosystems, Foster City, CA), and the threshold cycle (\(C_T\)) above background for each reaction was calculated. The \(C_T\) value obtained for each primer pair / template combination is shown, expressed as mean ± SD from assays performed in triplicate. n.d., amplification signals not detected.
### Supplemental Table 1. Mass Spectral Characteristics of 5-Arylresorcinols produced in ARS1, ARS2, Os05g12180, Os10g08620, and Os10g07040 Enzymatic Assays

<table>
<thead>
<tr>
<th>Acyl-CoA Starter</th>
<th>Product</th>
<th>(m/z)</th>
<th>Difference (amu)&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>C6:0&lt;sup&gt;b&lt;/sup&gt;</td>
<td>5-pentyl resorcinol (AR5:0; C_{11}H_{16}O_{2})</td>
<td>324[M]&lt;sup&gt;+&lt;/sup&gt;</td>
<td>56</td>
</tr>
<tr>
<td>C8:0</td>
<td>5-heptyl resorcinol (AR7:0; C_{13}H_{20}O_{2})</td>
<td>352[M]&lt;sup&gt;+&lt;/sup&gt;</td>
<td>84</td>
</tr>
<tr>
<td>C10:0</td>
<td>5-nonyl resorcinol (AR9:0; C_{15}H_{24}O_{2})</td>
<td>380[M]&lt;sup&gt;+&lt;/sup&gt;</td>
<td>112</td>
</tr>
<tr>
<td>C12:0</td>
<td>5-undecyl resorcinol (AR11:0; C_{17}H_{28}O_{2})</td>
<td>408[M]&lt;sup&gt;+&lt;/sup&gt;</td>
<td>140</td>
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<tr>
<td>C14:0</td>
<td>5-tridecyl resorcinol (AR13:0; C_{19}H_{32}O_{2})</td>
<td>436[M]&lt;sup&gt;+&lt;/sup&gt;</td>
<td>168</td>
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<tr>
<td>C16:0</td>
<td>5-pentadecyl resorcinol (AR15:0; C_{21}H_{36}O_{2})</td>
<td>464[M]&lt;sup&gt;+&lt;/sup&gt;</td>
<td>196</td>
</tr>
<tr>
<td>C16:1</td>
<td>5-pentadecenyl resorcinol (AR15:1; C_{21}H_{34}O_{2})</td>
<td>462[M]&lt;sup&gt;+&lt;/sup&gt;</td>
<td>194</td>
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<tr>
<td>C16:3</td>
<td>5-pentadecatrienyl resorcinol (AR15:3; C_{21}H_{30}O_{2})</td>
<td>458[M]&lt;sup&gt;+&lt;/sup&gt;</td>
<td>190</td>
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<tr>
<td>C18:0</td>
<td>5-heptadecyl resorcinol (AR17:0; C_{23}H_{40}O_{2})</td>
<td>492[M]&lt;sup&gt;+&lt;/sup&gt;</td>
<td>224</td>
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<tr>
<td>C18:1</td>
<td>5-heptadecenyl resorcinol (AR17:1; C_{23}H_{38}O_{2})</td>
<td>490[M]&lt;sup&gt;+&lt;/sup&gt;</td>
<td>222</td>
</tr>
<tr>
<td>C18:2</td>
<td>5-heptadecadienyl resorcinol (AR17:2; C_{23}H_{36}O_{2})</td>
<td>488[M]&lt;sup&gt;+&lt;/sup&gt;</td>
<td>220</td>
</tr>
<tr>
<td>C20:0</td>
<td>5-nonadecyl resorcinol (AR19:0; C_{25}H_{40}O_{2})</td>
<td>520[M]&lt;sup&gt;+&lt;/sup&gt;</td>
<td>252</td>
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<tr>
<td>C20:4&lt;sup&gt;b&lt;/sup&gt;</td>
<td>5-nonadecatetraenyl resorcinol (AR19:4; C_{25}H_{36}O_{2})</td>
<td>512[M]&lt;sup&gt;+&lt;/sup&gt;</td>
<td>244</td>
</tr>
</tbody>
</table>

<sup>a</sup> Difference denotes the mass value inferred by subtraction of the \(m/z\) 268 [(5-methyl-1,3 phenylene)bis(oxy)bis-(trimethylsilane)]<sup>+</sup> base ion from the [M]<sup>+</sup>.

<sup>b</sup> Substrates not utilized by Os05g12180, Os10g08620, or Os10g07040.
Supplemental Methods

Binary Vector Construction and Analysis of pARS1-RNAi and pARS2-RNAi Transgenic Events

Binary vectors were developed for RNAi-mediated repression of ARS1 and ARS2 using target regions of 602 and 556 bp in length, respectively, cloned in both sense and antisense orientation, separated by a 1.13 kb intron sequence derived from the Arabidopsis FAD2 gene (Okuley et al., 1994) and positioned downstream of the constitutive polyubiquitin-1 promoter from Zea mays with its cognate intron (Christensen et al., 1992). The specific target sequences selected for ARS1 correspond to nucleotides 758-1215 (3’ end) of the coding sequence plus an additional 145 bp of contiguous 3’ UTR sequence; the specific target sequences selected for ARS2 correspond to nucleotides 762-1218 (3’ end) of the coding sequence plus an additional 100 bp of contiguous 3’ UTR sequence. ARS1 and ARS2 target regions flanked by EcoRI (5’ end) and BamHI (3’ end) restriction sites were first generated by PCR amplification using S. bicolor (genotype Btx623) genomic DNA as template (described in ‘Methods’), to facilitate direct ligation with EcoRI- and BamHI-digested pUbi-IF2 (DNA Cloning Service, Hamburg, Germany). The primer pairs used for these PCR amplifications were as follows: ARS1 forward, 5’-CCCTGAATTCAACACGATACCGGA-3’, and reverse, 5’-CTCTGGATCCTTACGCACCGCCTTAT-3’; ARS2 forward, 5’-CCCTGAATTCAACACGATACCGGA-3’, and reverse, 5’-CCCGGATCCACCTTATGGTCCCAT-3’. The resulting intermediate constructs were then digested with BsrGI and MluI, and ARS1 and ARS2 target regions flanked by BsrGI (5’ end) and MluI (3’ end) were also generated in a second round of PCR amplifications as above. The primer pairs used for the second round of PCR amplifications were as follows: ARS1 forward, 5’-CCCTTGTACAGACCACGATACCGGA-3’, and reverse, 5’-CTCTACCGTTACGCACCGCCTTAT-3’; ARS2 forward, 5’-CCTCTGTACAGACCACGATACCGGA-3’, and reverse, 5’-CTCTACCGTGTCATCCTTATGGTCCCAT-3’. Following digestion with BsrGI and MluI, the PCR products were ligated with their corresponding intermediate vectors, resulting in the final intermediate vectors, pUbi-ARS1 and pUbi-ARS2 (not shown), containing the complete hpRNA-
generating transgene cassettes for *ARS1* and *ARS2* as confirmed by DNA sequence analysis. Finally, pUbi-ARS1 and pUbi-ARS2 were digested with SfiI, then the approximately 4.0 kb RNAi cassette-containing fragments were gel-purified and ligated with SfiI-digested pH9000 (Hausmann and Toepfer, 1999). The resulting binary vectors contain the hpRNA-generating cassettes arranged in a head-to-tail orientation with respect to the nptII selectable-marker cassette, and were designated pARS1-RNAi and pARS2-RNAi (Supplemental Figure 1 online). All DNA manipulations involved in the construction of pARS1-RNAi and pARS2-RNAi involved standard cloning procedures (Sambrook et al., 1989). For the generation of transgenic *S. bicolor* events, recombinant *A. tumefaciens* strains (NTL4/Chry5) harboring pARS1-RNAi and pARS2-RNAi were prepared, and used to transform immature embryos of *Sorghum bicolor* (genotype Tx430) as previously described (Howe et al., 2006).

To confirm expression of the hpRNA in the various transgenic events, 30 T₁ seeds were sown per event in perlite (Hummert’s, Earth City, MO) saturated with 0.5X Hoagland’s solution (Sigma-Aldrich, St. Louis, MO), and maintained for 10 days at 25°C in a growth chamber under a 16 h photoperiod, and light intensity of approximately 400 μmole m⁻² s⁻¹. On day 10, the perlite was gently removed and root systems from each seedling were individually numbered and harvested into 3.0 mL polypropylene tubes, flash-frozen in liquid nitrogen, then stored at -80°C prior to use. As a control, root systems were also harvested from non-transformed *S. bicolor* genotype Tx430 seedlings grown in an identical manner.

For detection of transgene-derived hpRNA transcripts, individual root systems were first hand-pulverized using a mortar and pestle, then 10 mg tissue aliquots from each root system were used for total RNA extractions. Total RNAs were prepared using an RNeasy Plant Mini-Kit (Qiagen Inc., Valencia, CA) with the inclusion of an additional homogenization step of 30 s at 25,000 rpm using a handheld homogenizer to aid tissue disruption, and an “on column” DNase I treatment using a RNase-Free DNase kit as per manufacturer’s instructions, to remove residual DNA contamination (Qiagen, Inc., Valencia, CA). RNA recovery and purity were determined spectrophotometrically, and sample integrity was assessed by agarose gel electrophoresis. Eight different transgenic events were initially screened for hpRNA expression (4 transformed with pARS1-RNAi and 4 transformed with pARS2-RNAi), thus a total of 240 root systems (30
seedlings per event) were independently analyzed in triplicate assays by quantitative real-time RT-PCR, as described in ‘Methods’. To distinguish the transgene-derived hpRNA transcripts from endogenous ARSI and ARS2 transcripts, the reverse primer used for real-time PCR assays was complementary to sequences within the octopine synthase (OCS) polyadenylation region immediately adjacent to the RNAi target sequences within pARS1-RNAi and pARS2-RNAi (Supplemental Figure 1 online). Since identical OCS polyadenylation regions were present in both vectors, the same reverse primer (5’-CGCATATCTCTATTAAAGCAGGGTC-3’) was used to screen both pARS1-RNAi and pARS2-RNAi transformants. For pARS1-RNAi transformants the forward primer used was 5’-CTCCTTGTGGCTAATTCATGGAC-3’, and for pARS2-RNAi transformants the forward primer used was 5’-CACATATATCGCCAATT CATGGAC-3’.

Based on the real-time PCR analyses described above, individual seedlings were scored as either hpRNA “+” (hpRNA transcript detected) or hpRNA “-” (hpRNA transcript not detected). Within each transgenic event, equal samples from all hpRNA “+” individuals were pooled, and a second pool of tissues was similarly prepared from all hpRNA “-” individuals. To determine whether hpRNA expression in roots correlated with inhibition of sorgoleone biosynthesis, all hpRNA “+” and “-” pooled samples were subjected to GC-MS (Figure 4C), as described in ‘Methods’. Additionally, the same hpRNA “+” and “-” pooled samples were used for follow-up real-time RT-PCR analyses, performed as described above, to examine the extent of ARSI and ARS2 endogenous transcript repression within pARS1-RNAi and pARS2-RNAi transformants (Figure 4A). The extensive sequence identity occurring between the coding and proximal 5’/3’ flanking regions of the two genomic ARSI-like copies (discussed in the text) precludes the design of primer pairs capable of distinguishing between these two sequences, thus the expression levels of total endogenous ARSI- and ARS2-like transcripts were monitored. For these analyses, PCR primers were designed within coding sequences excluded from ARSI and ARS2 RNAi-targetted regions, and were as follows: ARSI - forward, 5’-AGCTCCTTGTGGCTAATTCATGG-3’, reverse, 5’-TATAGGCACAAATAAATATAACACTTG-3’; ARS2 - forward, 5’-ATGGGGTTCCATGGGGAAG-3’, reverse, 5’-GGTGCCGGTACGCTC-3’. The specificity of both primer pairs were confirmed in reciprocal real-time RT-PCR tests using PCR-generated ARS1 and ARS2 templates, shown in Supplemental Figure 2 online.
Substrates and Chromatography Standards

Benzoyl-CoA, malonyl-CoA, butryl-CoA, isobutyryl-CoA, isovaleryl-CoA, hexanoyl-CoA, capryloyl-CoA, caproyl-CoA, lauroyl-CoA, myristoyl-CoA, palmitoyl-CoA, palmitoleoyl-CoA, stearoyl-CoA, oleoyl-CoA, linoleoyl-CoA, arachidoyl-CoA, arachidonoyl-CoA, and olivetol were purchased from Sigma-Aldrich (St. Louis, MO). Pentadecylresorcinol was purchased from Chem Service Inc. (West Chester, PA). 5-Pentadecatrienyl resorcinol was purified from cashew nut shell liquid as previously described (Paramashivappa et al., 2001).

For the preparation of hexadecatrienyl ((9Z, 12Z)-hexadeca-9,12,15-trienyl)-CoA, the identities of all compounds were confirmed using both physical and spectroscopic methods, including $^1$H-NMR, $^{13}$C-NMR, and high-resolution time-of-flight mass spectroscopy (HRTOFMS), as described below. 5-Hexen-2-yn-1-ol was prepared from allyl bromide and propargyl alcohol using the method described by Taber and You (1995), 5-Hexen-2-yn-1-ol was converted to 1-bromohexen-5-en-2-yne by the method described by Tyman and Visani (1997), and 9-decynoic acid was prepared from 8-bromoocanoic acid and lithium acetylide using the method described by Singh and Schnur (1986). 9-Decynoic acid was then converted to its methyl ester using the method described by Itoh et al. (2002). For preparation of methyl hexadeca-15-en-9,12-diynoate, methyl dec-9-ynoate (1.0 g, 5.5 mmol) was added to a stirred mixture containing CsCO$_3$ (1.79 g, 5.5 mmol), NaI (825 mg, 5.5 mmol), and CuI (825 mg, 5.5 mmol) in dry DMF (10 ml) at room temperature under nitrogen, then allowed to stir for an additional 20 min. To this mixture 6-bromohexen-1-en-4-yne (880 mg, 5.5 mmol) in DMF (2 mL) was added drop-wise, and allowed to stir overnight at room temperature. The reaction mixture was then quenched with saturated NH$_4$Cl, extracted with ethyl acetate, and the organic layer was dried over MgSO$_4$ and evaporated to completion. The residue was then chromatographed over silica gel and eluted with hexanes:ethyl acetate to yield methyl hexadeca-15-en-9,12-diynoate (1.3 g). The characterization data for methyl hexadeca-15-en-9,12-diynoate is provided below.
HRTOFMS: m/e (M+1) 261.18534 (calculated for C_{17}H_{25}O_{2}, 261.18545)

H NMR: (δ, CDCl3) 5.79 (1H, m, 15-H), 5.29 (1H, dd, J = 16.8, 1.6 Hz, 16-H), 5.13 (1H, dd, J = 10.0, 1.6 Hz, 16-H), 3.65 (3H, s, OCH3), 3.14 (2H, m, 14-H), 2.93 (2H, m, 11-H), 2.28 (2H, t, J = 7.2 Hz, 2CH2), 2.13 (2H, m, 8-H), 1.60 (2H, m), 1.46 (2H, m), 1.34 (2H, m), 1.23 (4H, m)

13C-NMR: (δ, CDCl3) 174.2 (C=O), 132.7 (C-15), 115.9 (C-14), 80.5, 77.1, 76.8, 74.3 (C-13,12,10,9), 51.4 (OMe), 34.0 29.0, 28.7, 28.6, 24.8, 23.0, 22.7, 18.6, 9.7. (9Z, 12Z)

For the preparation of methyl hexadeca-9,12,15-trienoate, a solution of methyl hexadeca-15-en-9,12-diynoate (800 mg) in ethyl acetate (15 mL) was hydrogenated in the presence of Lindlar catalyst (300 mg) at ambient pressure using a balloon for 20 h. The catalyst was then removed by filtration, and the solvent was evaporated to yield (9Z, 12Z)-methyl hexadeca-9,12,15-trienoate as the major product. A portion of the product was then purified by preparative thin layer chromatography using hexanes:ethyl acetate 98:2 (3 developments) to yield pure (9Z, 12Z)-methyl hexadeca-9,12,15-trienoate. The characterization data for (9Z, 12Z)-methyl hexadeca-9,12,15-trienoate is provided below.

For the preparation of methyl hexadeca-9,12,15-trienoic acid, 10 mL of 0.5 M LiOH was added dropwise while stirring to a solution of methyl hexadeca-9,12,15-trienoate (75 mg) in THF (25 mL) at 0°C. The reaction mixture was then stirred for an additional 30 min, allowed to cool to room temperature, then left stirring for an additional 12 h. The majority of the THF was removed under vacuum, then the remaining solution was adjusted to pH 2.0 with HCl (1 M), and extracted with ether. The organic phase was then dried to completion, yielding pure...
(9Z, 12Z)-hexadeca-9,12,15-trienoic acid. The characterization data for (9Z, 12Z)-hexadeca-
9,12,15-trienoic acid is provided below.

\[ ^1H \text{NMR:} \ (\delta, \text{CDCl}_3) 
5.83 \text{ (1H, m, 15-H)}), 
5.49-5.33 \text{ (4H, m, 13,12,10,9-H)}, 
5.07 \text{ (1H, dd, J = 17.2, 1.6 Hz, 16-H)}, 
5.00 \text{ (1H, dd, J = 10.0, 1.6 Hz, 16-H)}, 
2.82 \text{ (4H, m, 11,14-H)}, 
2.36 \text{ (2H, t, J = 7.2 Hz, 2-H)}, 
2.07 \text{ (2H, m, 8-H)}, 
1.65 \text{ (2H, m, 3-H)}, 
1.34 \text{ (10H, brs, 7,6,5,4,3-H)} \]

\[ ^13C\text{-NMR:} \ (\delta, \text{CDCl}_3) 
180.4 \text{ (C=O)}, 
136.8 \text{ (C-15)}, 
130.3 \text{ (C-9)}, 
129.3 \text{ (C-12)}, 
127.7 \text{ (C-10)}, 
126.8 \text{ (C-13)}, 
114.7 \text{ (C-16)}, 
34.1, 31.5, 29.5, 29.1, 29.0, 29.0, 27.2, 25.6, 24.6. \]

For the preparation of (9Z, 12Z)-Hexadeca-9,12,15-trienyl chloride, (9Z, 12Z)-Hexadeca-
9,12,15-trienoic acid in toluene (3.5 mL) was first treated with oxalyl chloride (0.5 mL) at 35-
40°C for 1 h, then the solvent and excess oxalyl chloride was evaporated under vacuum. The
residue was then re-dissolved in toluene and the solvent was evaporated under vacuum. This
process was repeated twice more, then the product, (9Z, 12Z)-hexadeca-9,12,15-trienyl chloride,
was directly used in the subsequent reaction. (9Z, 12Z)-Hexadeca-9,12,15-trienyl-CoA was then
prepared from (9Z, 12Z)-hexadeca-9,12,15-trienyl chloride and coenzyme A using a
modification of the procedure described by Bishop and Hajra (1980). Briefly, to a stirred
solution of 300 mg (0.38 mmol) coenzyme A in 5 mL of a 1:2.2 mixture of aqueous NaHCO₃
(150 mM, pH adjusted to 8.8 with NaOH) and tetrahydrofuran at 35°C under nitrogen, (9Z,
12Z)-hexadeca-9,12,15-trienoyl chloride (48 mg, 0.19 mmol) was added. After 30 min, the
reaction was quenched by the addition of 80 μL of 10% HClO₄, and then concentrated under a
stream of nitrogen. The precipitate was then recovered by centrifugation at 20,000 x g for 15 min
at 4°C, decanted, then mixed with 8 mL of 1.3% perchloric acid. The mixture was then chilled
using an ice bath, and re-centrifuged at 20,000 x g for 15 min at 4°C. The pellet was then
washed with ice-cold perchloric acid (1.3%, 8 mL), then washed with 12 mL acetone, followed
by two additional washes with 12 mL ether, and finally air-dried to completion and resuspended
in phosphate buffer (pH 7.0) at a final concentration of 1.0 mM. The molecular weight of (9Z,
12Z)-Hexadeca-9,12,15-trienyl-CoA was determined to be m/e (M+1) 1000.7276 by
HRTOFMS.
For the above synthesis, allyl bromide, propargyl alcohol, 8-bromooctanoic acid and lithium acetylide ethylenediamine complex were purchased from Sigma-Aldrich (St. Louis, MO). $^1$H-NMR and $^{13}$C-NMR spectra were recorded using an Avance DPX-400 spectrometer (400 MHz for $^1$H NMR, 100 MHz for $^{13}$C NMR; Bruker Biospin Corp., Billerica, MA) in CDCl$_3$, using tetramethylsilane as an internal standard. HRTOFMS were measured on an Agilent Series 1100 SL mass spectrometer equipped with an ESI source (Agilent Technologies, Santa Clara, CA).
Supplemental References


