Characterization of gossypol biosynthetic pathway

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Gossypol and related sesquiterpene aldehydes in cotton function as defense compounds but are antinutritional in cottonseed products. By transcriptome comparison and coexpression analyses, we identified 146 candidates linked to gossypol biosynthesis. Analysis of metabolites accumulated in plants subjected to virus-induced gene silencing (VIGS) led to the identification of four enzymes and their supposed substrates. In vitro enzymatic assay and reconstitution in tobacco leaves elucidated a series of oxidative reactions of the gossypol biosynthesis pathway. The four functionally characterized enzymes, together with (+)-δ-cadinene synthase and the P450 involved in 7-hydroxy(+)-δ-cadinene formation, convert farnesyl diphosphate (FPF) to hemigossypol, with two gaps left that each involves aromatization. Of six intermediates identified from the VIGS-treated leaves, 8-hydroxy-7-keto-δ-cadinene exerted a deleterious effect in dampening plant disease resistance if accumulated. Notably, CYP718E79, the enzyme responsible for converting this phytoxic intermediate, exhibited the highest catalytic activity among the five enzymes of the pathway assayed. In addition, despite their dispersed distribution in the cotton genome, all of the enzyme genes identified show a tight correlation of expression. Our data suggest that the enzymatic steps in the gossypol pathway are highly coordinated to ensure efficient substrate conversion.

Significance

Cotton is an important crop, and terpenoids form the largest group of natural products. Gossypol and related sesquiterpene aldehydes in cotton function as phytoalexins against pathogens and pests but pose human health concerns, as cotton oil is still widely used as vegetable oil. We report the isolation and identification of four enzymes and the recharacterization of one previously reported P450. We are now close to the completion of the gossypol pathway, an important progress in agricultural and plant sciences, and the data are beneficial to improving food safety. Among the six compounds (intermediates) isolated following gene silencing, one affected plant desaturation and cyclic ether formation steps in the pathway. However, until now, neither the enzymes nor the reactions downstream of (+)-δ-cadinene have been characterized, except a tentative identification of CYP706B1, and even the biosynthetic intermediates remain largely unknown.

All cotton species bear the lysigenous glands located in the subepidermal layer of aerial organs, in which sesquiterpene aldehydes (such as gossypol and hemigossypolone) are stored. There are also glandless cultivars which do not produce these phytoalexins in aerial parts (17, 24, 25) (Fig. 1A and B). Recently, the gene responsible for gland formation, GoPGF, was cloned, which encodes a basic helix–loop–helix transcription factor (25). By transcriptome-based comparison of the glandular and the glandless cultivars and coexpression analyses, in combination with virus-induced gene silencing (VIGS) and partial reconstitutions of the pathway in heterologous system, we isolated four enzymes and identified five steps of the pathway, covering the

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first four consecutive steps and most of the hydroxylation reactions of gossypol biosynthesis.

Results

Isolation of Gossypol Pathway Genes. Upland cotton, *Gossypium hirsutum*, is an allotetraploid species widely cultivated around the world (26). Analyses by HPLC detected a high level of sesquiterpene aldehydes in the leaf, seed (cotyledon), and floral organs of *G. hirsutum* cv. CCR112, but not the glandless mutant CCR112gl (SI Appendix, Fig. S1A). Although the sesquiterpenes are widely distributed throughout the glandular cotton plant, their level and composition in different organs vary: while gossypol is predominant in seed and root, hemigossypolone is abundant in leaf (SI Appendix, Fig. S1A).

In cotton CDN, a sesquiterpene cyclase and the cytochrome P450 monooxygenase CYP706B1 catalyze the first two steps of gossypol biosynthesis (17, 19). To further characterize the pathway, we adopted an integrative approach combining two-stage transcriptome analyses and VIGS to isolate genes encoding the downstream enzymes. Comparison of the transcript abundances in the leaves of glandular and glandless cotton uncovered 902 genes significantly down-regulated in the latter (Fig. 1C). Next, correlation analysis using the correlation value ≥0.5 grouped 5,912 transcripts with the bait CDN of the CDN family (Fig. 1C). Combination of these two datasets disclosed 146 genes in total that were potentially linked to gossypol biosynthesis, among which 82 encode enzymes, including the previously reported CDN and CYP706B1, and the mevalonate (MVA) pathway genes (Fig. 1D). Subsequent analysis of spatial expression patterns using the R pheatmap package identified seven enzymes that form the most likely gene expression cluster related to gossypol biosynthesis (Fig. 1E and SI Appendix, Table S1), of which four have not been investigated before.

Real-time quantitative PCR confirmed the RNA-sequencing data: the four enzyme genes were tightly coexpressed with CDN and CYP706B1, with their transcript levels high in glandular leaves but low or undetectable in glandless leaves (Fig. 2A). During development, young ovules (seeds) do not produce gossypol until 20 d postanthesis (SI Appendix, Fig. S1B), when CDN and CYP706B1 as well as the four candidate genes were coordinately activated, concomitant with gossypol accumulation (Fig. 2B).

Previous investigations demonstrated that biosynthesis of sesquiterpene phytoalexins in cotton cells can be induced by the pathogenic fungus *Verticillium dahliae* (17, 20). HPLC analysis showed that treatment of cotton cotyledons by the *V. dahliae*
elicit VdNEP (27) led to increased production of gossypol and hemigossypolone, whereas in glandless cotyledons, in which the sesquiterpene aldehydes were undetectable before elicitation, hemigossypolone was induced to accumulate (SI Appendix, Fig. S2). Consistently, the six enzyme genes were all up-regulated by elicitation (Fig. 2 C and D).

Selected candidate genes were submitted to VIGS, and silenced genes were then monitored by metabolite analysis of cotton leaves (28). Silencing of CDNC decreased hemigossypolone and gossypol levels by 95.1% and 96.2%, respectively, and silencing of CYP706B1 decreased the sesquiterpene levels by 59.4% and 61.2%, respectively, compared with empty vector controls (Fig. 2 E). An extended assay showed that silencing of four enzymes, including two cytochromes P450 (CYP82D113 and CYP71BE79), one alcohol dehydrogenase (DH1), and one 2-oxoglutarate/Fe(II)-dependent dioxygenase (2-ODD-1), each reduced the level of gossypol and hemigossypolone by more than 50% (Fig. 2 E). These data strongly suggested the involvement of the candidate genes in gossypol biosynthesis.

Identification of Biosynthetic Intermediates. As silencing of CYP706B1 resulted in an accumulation of its substrate (+)-δ-cadinene in cotton leaves (Fig. 3 A), we further analyzed the leaf extracts of the VIGS-treated plant by GC-MS and LC-MS to explore clues to the enzyme activity. We found that the compound as 7-hydroxy-(+)-δ-cadinene (Fig. 4). Subsequent incubation revealed that DH1 converted the CYP706B1 product into a compound of m/z of 228 (Fig. 3 E).

We also noted that the VIGS-CYP71BE79 plants grown in the greenhouse frequently developed disease phenotypes (brown sunken lesions covering the hypocotyl–root junction) (SI Appendix, Fig. S3 A and B), similar to the symptoms caused by the soilborne necrotrophic fungus Rhizoctonia solani (29), whereas the control and other VIGS-treated plants did not. As PGF silencing blocked the whole gossypol biosynthesis pathway (25), the decreased amount of sesquiterpene phytoalexins in VIGS-CYP71BE79 plants was unlikely responsible for the enhanced susceptibility. Determination by LC-MS revealed that the substrate of CYP71BE79 accumulated in the hypocotyl–root junction after the gene silencing (SI Appendix, Fig. S3 C).

Functional Characterization of Enzymes. To obtain intermediate standards for structure elucidation and to perform enzyme assays in vitro, we expressed the three cytochromes P450 in Saccharomyces cerevisiae and other enzymes in Escherichia coli. As determined by GC-MS, incubation of the starting substrate FPP with CDNC produced (+)-δ-cadinene, and further reaction with CYP706B1 gave rise to a hydroxylated product (Fig. 4) that was previously proposed to be 8-hydroxy-(+)-δ-cadinene (19). Subsequent incubation revealed that DH1 converted the CYP706B1 product into a compound of m/z 218 (Fig. 4), suggesting a dehydrogenation reaction. NMR spectroscopy detected a ketonic group at the C-7 position; thus, the product is 7-keto-δ-cadinene (Fig. 4).

Formation of 7-keto-δ-cadinene cast doubt on the previous identification of the CYP706B1 product as 8-hydroxy-(+)-δ-cadinene based on 1H-NMR spectroscopy (19). Indeed, both 13C NMR and heteronuclear multiple-bond correlation spectra revealed the compound as 7-hydroxy-(+)-δ-cadinene (SI Appendix, Figs. S4–S6). Thus, CYP706B1 is reassigned as (+)-δ-cadinene-7-hydroxylase, and DH1 is 7-keto-δ-cadinene synthase (Fig. 4).
The compound 7-keto-δ-cadinene was first identified from *G. hirsutum* plants engineered to express an RNAi construct targeting CYP82D109, which was named (4αR, 5β)-δ-cadinene-2-one (24), but the activity of CYP82D109 has remained unknown. CYP82D113 is 92% identical to CYP82D109. To determine the enzyme activity of CYP82D113, yeast microsomes enriched with CYP82D113 were incubated with 7-keto-δ-cadinene. LC-MS analysis identified an expected peak of the product having an m/z of (+) 257. MS and NMR analyses indicated that, in the presence of NADPH, CYP82D113 transferred a hydroxyl group to C-8 of 7-keto-δ-cadinene, generating 8-hydroxy-7-keto-δ-cadinene (Fig. 3D). To test whether CYP71BE79 is involved in further decoration of the δ-cadinene backbone, we incubated it with 8-hydroxy-7-keto-δ-cadinene, which was then efficiently converted into a product with an m/z of (+) 273 [M + Na]+ (Fig. 4). NMR analysis identified that CYP71BE79 transferred a new hydroxyl group to C-11 to form 8,11-dihydroxy-7-keto-δ-cadinene (SI Appendix, Figs. S10–S12).

Lastly, the metabolite accumulated in the 2-ODD-1–silenced leaves (Fig. 3E) was identified to be furcalalen-2-one (SI Appendix, Figs. S13–S14). As expected, incubation with 2-ODD-1 converted it to a new compound, 3-hydroxy-furocalamen-2-one (Fig. 4 and SI Appendix, Figs. S15–S16).

We next measured the kinetic parameters of the five enzymes (Table 1). Notably, CYP71BE79 exhibited a much higher maximum activity (Vmax) than other enzymes tested, including two upstream cytochromes P450 (CYP706B1 and CYP82D113), and its catalytic efficiency (Vmax/Km) was also clearly higher. To test substrate specificity, the five enzymes were assayed with available intermediates possessing similar structures. Most enzymes showed little activity toward alternative substrates under identical assay conditions (SI Appendix, Fig. S17). However, in addition to 7-hydroxy-(+)-δ-cadinene, DH1 also accepted 8-hydroxy-7-keto-δ-cadinene and 8,11-dihydroxy-7-keto-δ-cadinene as substrates, although with lower efficiency (SI Appendix, Fig. S17). Thus, DH1 is,
to some extent, promiscuous in dehydrogenation of the hydroxyl group-containing metabolites.

**Partial Reconstitution of Gossypol Pathway in Tobacco Leaf.** Along with in vitro assays of enzyme activities, we utilized the Agrobacterium-mediated transient expression system to reconstitute the gossypol pathway reactions in *Nicotiana benthamiana* leaves. The 3SS promoter was used to express each of the six enzymes, including an FPP synthase (AtFPS2) from *Arabidopsis thaliana* (AT4G17190), as well as CDNC, CYP706B1, DH1, CYP82D113, and CYP71BE79 from cotton, which catalyze the six consecutive steps of gossypol biosynthesis starting from isopentenyl diphasphate/dimethylallyl diphosphate. Four metabolic intermediates, (+)-δ-cadinene, 7-hydroxy-(+)-δ-cadinene, 7-keto-δ-cadinene, and 8-hydroxy-7-keto-δ-cadinene, were detected by GC-MS, and metabolite profiles were monitored as total-ion chromatograms (TIC), whereas 8-hydroxy-7-keto-δ-cadinene, 8,11-dihydroxy-7-keto-δ-cadinene, furocalamen-2-one, and 3-hydroxy-furocalamen-2-one were detected by LC-MS with UV, as indicated. The sample without the relevant protein served as negative control. Structures of all compounds, except (+)-δ-cadinene, were further determined by MS/MS and NMR spectroscopy (SI Appendix, Figs. S4–S16 and Tables S2 and S3). The purified recombinant proteins of DH1 and 2-ODD-1 expressed in *E. coli* and the microsomes of yeast cells expressing the respective cytochromes P450 were assayed.

**Fig. 4.** Functional characterization of enzymes by in vitro assays and determination of the products. (+)-δ-Cadinene, 7-hydroxy-(+)-δ-cadinene, and 7-keto-δ-cadinene were detected by GC-MS, and metabolite profiles were monitored as total-ion chromatograms (TIC), whereas 8-hydroxy-7-keto-δ-cadinene, 8,11-dihydroxy-7-keto-δ-cadinene, furocalamen-2-one, and 3-hydroxy-furocalamen-2-one were detected by LC-MS with UV, as indicated. The sample without the relevant protein served as negative control. Structures of all compounds, except (+)-δ-cadinene, were further determined by MS/MS and NMR spectroscopy (SI Appendix, Figs. S4–S16 and Tables S2 and S3). The purified recombinant proteins of DH1 and 2-ODD-1 expressed in *E. coli* and the microsomes of yeast cells expressing the respective cytochromes P450 were assayed.
8,11-dihydroxy-7-keto-δ-cadinene itself, was formed (SI Appendix, Fig. S18 E–G).

Together, data from VIGS and in vitro and tobacco leaf transient expression assays suggest that CYP706B1, DH1, CYP82D113, and CYP71BE79 catalyze four consecutive oxidative reactions on (+)-δ-cadinene, and 2-ODD-1 is responsible for a later hydroxylation step in the biosynthetic pathway leading to sesquiterpene aldehydes (Fig. 5).

Gossypol Pathway Genes Are Dispersed in the Cotton Genome. Several examples exist where genes encoding biosynthetic pathway enzymes of specialized metabolites, including terpenoids and alkaloids, tend to be clustered together in the plant genome (3, 6, 30, 31). In cotton, however, the gossypol pathway genes are dispersed among different chromosomes (Fig. 5 and SI Appendix, Fig. S19). On the other hand, the gene families of the gossypol as well as the core MVA pathways are often extensively expanded with tandem duplications (Fig. 5 and SI Appendix, Fig. S19).

Most of the gossypol pathway enzymes identified, including CDN, DH1, CYP82D113, and 2-ODD-1, appear to have arisen from local duplications in the cotton genome. For example, in the allotetraploid genome of *G. hirsutum*, there are 11 genes encoding the alcohol dehydrogenase DH1 and homologs, all of which are tandemly arranged, with four genes (Gh_A01G1736, Gh_A01G1737, Gh_A01G1739, and Gh_A01G1740) on chromosome A1 (chromosome 1 of A subgenome) and seven (Gh_D01G1983 to Gh_D01G1989) on chromosome D1 (Fig. 5 and SI Appendix, Fig. S19).

Among the five enzymes catalyzing oxidative steps in the gossypol biosynthetic pathway, three are cytochromes P450 of different families. Members of CYP71 and CYP82 families are commonly involved in biosynthesis of specialized metabolites such as noscapine (32), podophyllotoxin (33), and artemisinin (34). As cotton CYP71BE79 is distinct in its high activity (Table 1), we analyzed it further.

Using CYP71BE79 as query, we performed a bioinformatic blast search of CYP71 family proteins from publicly available genomes of nine plant species, including three species from the family Malvaceae: *G. hirsutum*, *Durio zibethinus*, and *Theobroma cacao*. In total, 312 CYP71 proteins were retrieved (SI Appendix, Fig. S20). We found that the CYP71BE proteins form a Malvaceae-specific subfamily (green in Fig. 6A), which contained 37 members clustered into five clades. Clade II was composed of six CYP71BES, including the two CYP71BE79 homologs of *G. hirsutum* (Gh_A13G1133 and Gh_D13G1407). Notably, CYP71BE genes have been maintained as a truly single copy in diploid genomes or subgenomes (Fig. 6B).

The nonsynonymous (Ks) and synonymous substitution rates (Ka) of three gossypol pathway cytochromes P450 (CYP706B1, CYP82D113, and CYP71BE79) in *G. hirsutum* were compared with their homologs in *D. zibethinus* (Table 2). The higher Ks values and the lower Ka/Ks ratios of CYP71BE79 indicate that this P450 has undergone less relaxed selection. Moreover, CYP71BE79 has a high V_max value compared with other, identified cytochromes P450 of the gossypol pathway (Table 1), which supports an efficient transformation of its substrate (8-hydroxy-7-keto-δ-cadinene) that affects plant resistance to pathogens if accumulated (SI Appendix, Fig. S3). We propose that CYP71BE79 is functionally more conserved in *Gossypium* and in closely related genera in order to catalyze a highly controlled step to prevent the accumulation of the phytotoxic metabolite, along with gossypol pathway evolution.

| Enzyme          | Substrate                  | K_m μM   | V_max nmol·min⁻¹·mg⁻¹ |
|-----------------|----------------------------|----------|-----------------------|
| CYP706B1        | (+)-δ-Cadinene             | 7.57 ± 1.14 | 31.26 ± 1.56          |
| DH1             | 7-Hydroxy-(+)-δ-cadinene   | 0.48 ± 0.04 | 10.42 ± 0.21          |
| CYP82D113       | 7-Keto-δ-cadinene          | 1.02 ± 0.13 | 22.00 ± 0.73          |
| CYP71BE79       | 8-Hydroxy-7-keto-δ-cadinene| 9.67 ± 1.34 | 304.90 ± 10.88        |
| 2-ODD-1         | Furocalamen-2-one          | 1.81 ± 0.21 | 49.54 ± 1.11          |

Each dataset represents means ± SD (n = 3 independent experiments).

Discussion

Recent achievements in sequencing cotton genomes (26, 35–37) have facilitated the isolation and characterization of gossypol pathway enzymes through transcriptome mining. It is striking that the first oxidation reaction of (+)-δ-cadinene catalyzed by CYP706B1 toward gossypol biosynthesis occurs at the C-7 position, instead of C-8 as proposed previously. Besides gossypol and related sesquiterpene aldehydes that have a characteristic 8-hydroxy group, there are other cadinene derivatives featuring oxidation at C-7 in cotton, such as 2-hydroxy-7-methoxycedalene (24). An earlier study showing that the tritiated CYP706B1 product was incorporated into gossypol (38) supported the involvement of this cytochrome P450 in gossypol biosynthesis. Here, we provide evidence that CYP706B1 produces 7-hydroxy-(+)-δ-cadinene, which is an upstream intermediate in the gossypol pathway.

Interestingly, 7-hydroxy-(+)-δ-cadinene is subjected to C-8 oxidation following C-7 carbylation, and the C-7 carbonyl group seems indispensable for C-8 hydroxylation. The cadinene-type sesquiterpenes oxidized at both C-7 and C-8 have not been found before; subsequent oxidation at C-11 by CYP71BE79 presumably to react with a C-8 hydroxyl group to form a C-8–C-11 ether bridge in the structure of gossypol (Fig. 4). The fate of the C-7 carbonyl group awaits determination but could be deduced from structural comparison of 8,11-dihydroxy-7-keto-δ-cadinene and furocalamen-2-one, because the two intermediates leave a biosynthesis gap that may involve isomerization of carbonyl functionality to an enol group and the successive dehydrogenation to form a benzene ring (ring B). Isomerization and dehydration are not uncommon in aromatization, such as the shikimate pathway rearrangement of chorismate to prephenate by chorismate mutase and the dehydrogenation of arogenate to phenylalanine by arogenate dehydrogenase (39). Furthermore, ring B is also aromatized during deoxyxymegogossypol formation from 3-hydroxy-furocalamen-2-one (Fig. 4). The present investigation resolves most of the oxidation reactions involved, leaving two remaining gaps that each involves similar aromatization reactions.

Notably, the reaction steps of gossypol formation are not randomly cascaded but rather accurately cascaded, from an energy point of view. The oxidation always occurs in the position much easier to take place, and the introduced oxidized group reduces the energy barrier of the next oxidation. For example, the first hydroxylation proceeds in the active C-7 allylic position, and then the newly formed carboxylation leaves its α position more active for subsequent hydroxylation; such is also the case of air oxidation reactions in positions 3 and 8, where there are preexisting carbonyl groups. Lastly, aromatization provides the most stable naphtalene ring. Thus, the gossypol pathway has evolved and been optimized through several low-energy intermediates.

Table 1. Kinetic analyses of the enzymes determined in vitro
The clear order and the strict substrate specificity of these biosynthetic reactions imply that the gossypol biosynthetic pathway may have evolved step by step, which might be a reason for discrete distributions of enzyme genes in the genome. We anticipate that in some plants of Malvaceae, such as cacao, okra, and roselle, the biosynthetic pathways of cardenolide-type sesquiterpenes are not necessarily destined to be gossypol; the short-cut or diversified routes may result in a rich array of specialized structures.

Fig. 5. Genes of gossypol pathway enzymes and their expressions. Genes of the enzymes catalyzing the defined steps in MVA and gossypol pathways and their homologs are shown. The expressions are indicated by heatmap, estimated using Cuffdiff by computing the FPKM value (fragments per kilobase of transcript per million reads sequenced) for each transcript. Genes encoding the identified enzymes or showing an expression pattern correlated to gossypol biosynthesis are on the top. Dashed arrows indicate unidentified reaction(s). 0Ov, 0-dpa ovule; 25Ov, 25-dpa ovule; 0Sd, 0-h postgermination seed; 5Sd, 5-h postgermination seed; Lf, leaf; PMD, diphosphomevalonate decarboxylase; IPP, isopentenyl diphosphate; IPPI, IPP isomerase; Lf, leaf; MVK, mevalonate kinase; MVP, phosphomevalonate kinase; Pe, petal; Pi, pistil; PMD, diphosphomevalonate decarboxylase; Rt, root; Sm, stamen; St, stem; To, torus. Distributions of the genes in G. hirsutum genome are indicated by their accession numbers and also shown in the genome atlas (SI Appendix, Fig. 519).
metabolites. Comparative analyses of these pathways will enrich our knowledge on evolution of sesquiterpene biosynthetic pathways and provide valuable data for safe use and further exploration of food, oil, and vegetable crops in the Malvaceae and related families.

There are two lines of evidence that support a tight regulation of the gossypol biosynthetic pathway. First, although not clustered in the genome as frequently observed with other specialized pathways (3, 6, 18, 30), genes of all six enzymes characterized show highly similar expression patterns. This raises the possibility that all these genes are regulated by a common transcription factor complex, as seen from the MYB-bHLH-WD40 complex in the anthocyanin biosynthetic pathway (40, 41). Second, products of these gossypol pathway enzymes are mostly undetectable in plant tissues unless the downstream enzyme genes are silenced, suggesting a highly efficient conversion, which could be a result of substrate channeling (42). For example, the monoterpene indole alkaloid pathway in *Catharanthus roseus* involves a complex and highly regulated biosynthesis in which the upstream pathway enzymes are separated in different cellular compartments to prevent inappropriate accumulation of highly reactive strictosidine aglycone (43).

In addition to their function as phytoalexins in plants, gossypol and related sesquiterpene aldehydes also show anticancer (44, 45), antimicrobial (46, 47), and spermicidal (48) activities. We wonder whether the six intermediates identified here have similar or novel biological activities. In particular, the structure of 8-hydroxy-7-keto-δ-cadinene features an α, β-unsaturated ketone

| Table 2. The evolution rates and K_α/Ks values of three homologous P450 gene pairs between G. hirsutum and D. zibethinus |
|-----------------|-----------------|-----------------|------|------|
| Gene name       | Genes in G. hirsutum | Homologs in D. zibethinus | K_α | Ks |
| CYP706B1_D      | Gh_D03G1513      | XM_022882367.1  | 0.1271 | 0.4514 | 0.2816 |
| CYP706B1_A      | Gh_A03G2006      | XM_022882367.1  | 0.1253 | 0.4342 | 0.2886 |
| CYP82D113_D     | Gh_D05G1894      | XM_022910758.1  | 0.1093 | 0.5405 | 0.2022 |
| CYP82D113_A     | Gh_A05G1705      | XM_022910758.1  | 0.105  | 0.5382 | 0.1951 |
| CYP718E79_D     | Gh_D13G1407      | XM_022861030.1  | 0.1201 | 0.9599 | 0.1251 |
| CYP718E79_A     | Gh_A13G1133      | XM_022861030.1  | 0.1165 | 0.9398 | 0.124 |

Tian et al. PNAS | vol. 115 | no. 23 | E5417
and an α-hydroxyl group next to the carbonyl, which may act as a Michael acceptor for biological nucleophiles; the similar enone group has been suggested as a general structural requirement for optimal cytotoxicity of quassinoids, a group of degraded terpenes with promising antitumor and cytotoxic activity (49, 50), suggesting that this intermediate may harbor interesting biological activities. Clonization of the enzymes makes it possible to obtain these hidden natural products in large quantity for drug or agrochemical screening.

**Details**

Details about plant materials and growth conditions are described in SI Appendix, SI Materials and Methods. Gene expression, elicitation, plant transformation, heterologous expression and purification of proteins, pathway reconstitution in N. benthamiana leaves, pathogen infection, enzymes assays, metabolite extraction, detection, and analysis were carried out according to protocols described in SI Appendix, SI Materials and Methods.

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