A Single Amino Acid Substitution Converts Benzophenone Synthase into Phenylpyrone Synthase

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Benzophenone metabolism provides a number of plant natural products with fascinating chemical structures and intriguing pharmacological activities. Formation of the carbon skeleton of benzophenone derivatives from benzoyl-CoA and three molecules of malonyl-CoA is catalyzed by benzophenone synthase (BPS), a member of the superfamily of type III polyketide synthases. A point mutation in the active site cavity (T135L) transformed BPS into a functional phenylpyrone synthase (PPS). The dramatic change in both substrate and product specificities of BPS was rationalized by homology modeling. The mutation may open a new pocket that accommodates the phenyl moiety of the triketide intermediate but limits polyketide elongation to two reactions, resulting in phenylpyrone formation. 3-Hydroxybenzoyl-CoA is the second best starter molecule for BPS but a poor substrate for PPS. The ary moiety of the triketide intermediate may be trapped in the new pocket by hydrogen bond formation with the backbone, thereby acting as an inhibitor. PPS is a promising biotechnological tool for manipulating benzoate-primed biosynthetic pathways to produce novel compounds.

Benzophenone derivatives constitute a class of plant secondary metabolites, which includes a number of chemically complex and pharmacologically active constituents. Bridged polycyclic compounds result from the stepwise prenylation of the benzophenone nucleus and concomitant intramolecular cyclization reactions of the attached C5 and C10 isoprenoid units (1). To date, no chemical synthesis has been reported for any of these caged molecules. A well studied compound is garcinol (2), where it serves as floral UV pigments and defense compounds against herbivores (3). Benzophenone derivatives also includes xanthones, which are the products of regioselective oxidative phenol coupling reactions catalyzed by cytochrome P450 enzymes (4). An example of a bridged polycyclic xanthone is gambogenic acid (Fig. 1), which induces apoptosis independently of the cell cycle by a novel mechanism of caspase activation (6). Active benzoylphloroglucinol derivatives may be potential lead compounds for designing new drugs.

The carbon skeleton of benzophenone derivatives is formed by benzophenone synthase (BPS).4 The enzyme from Hypericum androsaemum (Clusiaceae) catalyzes the iterative condensation of benzoyl-CoA with three molecules of malonyl-CoA to give a linear tetraketide intermediate, which is subsequently cyclized into 2,4,6-trihydroxybenzophenone via intramolecular Claisen condensation (7). In contrast, BPS from Centaurium erythraea (Gentianaceae) prefers 3-hydroxybenzoyl-CoA as a starter substrate and catalyzes the formation of 2,3’,4,6-tetrahydroxybenzophenone (8). These two BPS products are the precursors of all prenylated benzoylphloroglucinol and xanthone compounds.

BPS is a type III polyketide synthase (PKS) (7). This superfamily of enzymes generates an array of secondary metabolites in various plants and microorganisms (9). The most common type III PKS in higher plants is chalcone synthase (CHS), which initiates flavonoid biosynthesis (10). A number of functionally divergent plant PKSs arose via gene duplication and diversification (11). They differ from CHS in their preference for starter substrates (aliphatic or aromatic units), the number of acetyl additions catalyzed (one to seven), and the mechanism of ring formation used to cyclize linear polyketide intermediates (Claisen condensation, aldol condensation, or heterocyclic lactone formation) (9). Conversely, type III PKS share a homodimeric architecture, a similar subunit molecular mass, and a conserved polyketide extension mechanism. Furthermore, each type III PKS monomer utilizes a Cys-His-Asn catalytic triad with a buried active site cavity, as first revealed in the crystal structure of alfalfa CHS (12). A narrow CoA-binding tunnel provides access to this large bi-lobe internal cavity termed the initiation/elongation cavity. One lobe of this catalytic center forms the starter unit binding pocket and the other accommodates the growing polyketide chain (13).

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The on-line version of this article (available at http://www.jbc.org) contains supplemental Table S1.

The abbreviations used are: BPS, benzophenone synthase; CHS, chalcone synthase; GC-EI-MS, gas chromatography-electron ionization-mass spectrometry; HPLC-DAD, high performance liquid chromatography-diode array detection; PCS, pentaketide chromone synthase; PKS, polyketide synthase; PPS, phenylpyrone synthase; 2PS, 2-pyrone synthase; STS, stilbene synthase; RI, retention index.

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Mutagenic Conversion of BPS into PPS

The aim of this work was to diversify BPS activity by introducing mutations into the residues lining the active site cavity of the enzyme from *H. androsaemum*. A point mutation was found to alter both substrate and product specificities of BPS without decreasing the catalytic efficiency. This dramatic functional variation was rationalized by homology modeling. The BPS mutant, which was termed phenylpyrone synthase (PPS), has so far not been found as a naturally occurring type III PKS.

Interestingly, derivatives of the PPS product are found to alter both substrate and product specificities of BPS without decreasing the catalytic efficiency. This dramatic functional variation was rationalized by homology modeling. The BPS mutant, which was termed phenylpyrone synthase (PPS), has so far not been found as a naturally occurring type III PKS.

**EXPERIMENTAL PROCEDURES**

**Chemicals**—2,4,6-Trihydroxybenzophenone and 6-phenyl-4-hydroxy-2H-pyran-2-one were purchased from ICN (Mecklenburg, Germany) and Aurora Fine Chemicals (Graz, Austria). 2,3′,4,6-Tetrahydroxybenzophenone was synthesized as described previously (5). Benzoyl-CoA, malonyl-CoA, and acetyl-CoA were obtained from Sigma. 2-Hydroxybenzoyl-CoA, 3-hydroxybenzoyl-CoA, cinnamoyl-CoA, 2-coumaroyl-CoA, 3-coumaroyl-CoA, and 4-coumaroyl-CoA were synthesized as described previously (8, 15).

**Enzyme Assays, Determination of Kinetic Constants, and Inhibition Experiments**—Enzyme assays (250 μl) contained 15 μM starter CoA ester, 65 μM malonyl-CoA, 0.1 M potassium phosphate buffer, pH 7.0, and 2–5 μg of protein. After incubation at 35 °C for 20 min, the reaction mixture was acidified by addition of acetic acid (5%) and extracted twice with 250 μl of ethyl acetate. The combined organic phase was dried under vacuum, and the residue was dissolved in 50 μl of methanol (50%). Kinetic data were determined, as described previously (7). Inhibition experiments included a 5-min preincubation with varying concentrations of 3-hydroxybenzoyl-CoA and acetyl-CoA (2.7–21.6 μM) in the presence of malonyl-CoA (21.7 μM). Control assays contained only protein or protein and malonyl-CoA. After preincubation, benzoyl-CoA (7.8 μM) was added, and the formation of phenylpyrone was determined over a 20-min incubation period.

**Mutagenesis**—Site-directed mutants were generated using the QuikChange (Stratagene) protocol and verified by DNA sequencing.

**Analytical Methods**—HPLC-DAD analysis of the products from enzyme assays was carried out as described previously (16), except for the following modifications: gradient from 50 to 90% (v/v) methanol in water over 20 min; column Hypersil Gold (5 μm, 4.6 × 150 mm; Thermo). GC-MS was performed using two configurations. Configuration I has been described previously (16). For configuration II, a Hewlett-Packard 6890N gas chromatograph was equipped with a 30-m × 0.32-mm (film thickness 0.25 μm) analytical column (ZB-5, Agilent, Waldbronn, Germany). The capillary column was directly coupled to a GCMate II mass spectrometer (Jeol, Japan). The conditions applied were as follows: injection and transfer line were set at 260 and 280 °C, respectively; the GC oven program used was 70 °C for 3 min, increased by 10 °C/min to a final temperature of 310 °C for 3 min; the injection volume was 1 μl; the split ratio was varied from 1:2 to 1:20, depending on the concentration of the analytes; the constant carrier gas flow was 1.5 ml min⁻¹; the mass spectra were recorded at 70 eV.

The identity of individual compounds was confirmed by comparison of the MS data with spectra in the NIST mass spectral library version 2.0a and with reference compounds. The retention index (RI) on ZB-1 and ZB-5 columns was calculated with the aid of a set of hydrocarbons (even numbered C₈ to C₂₈) by linear interpolation for 2,4,6-tri-(trimethylsiloxy)-benzophenone as follows: RI (ZB-1) 2204, RI (ZB-5) 2250; GC-EI-MS (70 eV), m/z (%), 446 (4) [M⁺], 433 (16), 432 (35), 431 (100), 369 (6), 359 (4), 269 (4), 208 (5), 147 (7), 135 (4), 105 (29), 77 (9), 74 (4), 73 (55), and 45 (9); 2,3′,4,6-tetra-(trimethylsiloxy)-benzophenone as follows: RI (ZB-5) 2436; GC-EI-MS (70 eV)/ m/z (%)/ 534 (9) [M⁺], 523 (3), 522 (8), 521 (26), 520 (48), 519 (100), 369 (7), 357 (3), 193 (19), 165 (3), 147 (4), 74 (3), 73 (43), and 45 (4); 6-phenyl-4-(trimethylsiloxy)-2H-pyran-2-one as follows: RI (ZB-1) 2065, RI (ZB-5) 2135; GC-EI-MS (70 eV), m/z (%), 260 (91) [M⁺], 245 (24), 233 (18), 232 (100), 217 (4), 203 (7), 189 (12), 158 (10), 145 (4), 115 (6), 105 (33), 99 (5), 77 (21), 75 (8), 73 (30), and 45 (6); and 6-(3′-trimethylsiloxylphenyl)-4-(trimethylsiloxy)-2H-pyran-2-one as follows: RI (ZB-5) 2453; GC-EI-MS (70 eV), m/z (%), 348 (77) [M⁺], 334 (7), 333 (28), 322 (11), 321 (29), 320 (100), 305 (8), 246 (5), 193 (18), 159 (10), 99 (4), 75 (9), 74 (7), 73 (79), and 45 (8).

**Molecular Modeling**—A homology model of *H. androsaemum* BPS was constructed based on the crystal structure of alfalfa (Medicago sativa) CHS2 (12) complexed with resveratrol (Protein Data Bank code 1CGZ) using the Swiss-Model server (17, 18). The conserved CoA-binding sites and catalytic residues were correctly aligned, which indicates that the generated model is reliable. Modeling of protein-substrate complexes was performed using the program MOLOC and the implemented united atom force field MAB (19). The CoA molecule was placed according to the reference structure of alfalfa CHS2 (12) complexed with malonyl-CoA (Protein Data Bank code 1CML). Models for the full elongation cycle, including the hemithioketal covalently bound to Cys-167, were constructed based on the known catalytic mechanism of type III PKSs. All generated models were initially minimized and subsequently relaxed by molecular dynamics simulations for 1 ns at 300 K. We used the MAB force field (20) with implicit solvation and a hydrogen bond weight of 1.786. Modeling of point mutants was...
performed on the basis of the previously constructed model of wild-type BPS to investigate the perturbation introduced by the mutations in the active site cavity.

RESULTS

Generation of BPS Mutants—Based on the crystal structure of *M. sativa* CHS2 complexed with resveratrol (12), a homology model of BPS was generated using the Swiss-Model server. CHS and BPS share 59.1% amino acid sequence identity (7). The sequential polyketide intermediates of the phlorbenzophenone biosynthesis reaction from monoketide through tetraketide were modeled into the BPS active site cavity as Cys-167 and malonyl-CoA that is used for the next chain elongation reaction. Amino acids of the initiation and elongation pockets are highlighted in pink and green, respectively. The catalytic triad is displayed in yellow (Cys-167) and gray (His-307 and Asn-340).

![Figure 2. CHS-based homology model of the BPS active site cavity complexed with both the benzoyl-primed diketide that is covalently bound to the catalytic cysteine 167 and malonyl-CoA that is used for the next chain elongation reaction. Amino acids of the initiation and elongation pockets are highlighted in pink and green, respectively. The catalytic triad is displayed in yellow (Cys-167) and gray (His-307 and Asn-340).](Image)

![Diagram 1](Image)

| TABLE 1 | Corresponding active site cavity residues of crystallized alfalfa CHS2 (12) and *H. androsaemum* BPS |
|---------|---------------------------------------------------------------------------------------------------------------------------------|
| CHS *M. sativa* | BPS *H. androsaemum* |
| Initiation pocket | Methionine 217 | Aspartate 217 |
| Leucine 214 | Isoleucine 254 | Isoleucine 258 |
| Isoleucine 254 | Glycine 256 | Alanine 260 |
| Glycine 256 | Phenylalanine 265 | Tyrosine 269 |
| Elongation pocket | Methionine 200 | Serine 219 |
| Threonine 132 | Threonine 194 | Threonine 197 |
| Serine 133 | Valine 196 | Methionine 199 |
| Threonine 194 | Threonine 197 | Threonine 200 |
| Valine 196 | Glycine 216 | Serine 219 |
| Threonine 197 | Leucine 263 | Methionine 267 |
| Glycine 216 | Serine 338 | Glycine 342 |
| Leucine 263 | Serine 338 | Glycine 342 |
| Catalytic triad | Aspartate 307 | Aspartate 336 |
| Cysteine 164 | Histidine 303 | Asparagine 336 |
| Histidine 303 | Asparagine 336 |
| Asparagine 336 | Asparagine 336 |
| | | |

idues. Seven amino acids are different (Table 1). None of the seven single mutants generated accepted CoA thioesters of cinnamic acids, such as 4-coumaroyl-CoA and cinnamoyl-CoA. The majority of the subsequently created double to 5-fold mutants were either inactive or inhibited the growth of their host *E. coli* cells. The remaining active mutants functionally resembled the wild-type enzyme. Thus, the functional conversion of BPS into CHS failed. Previously, *H. androsaemum* CHS was transformed by triple mutation into an enzyme with a preference for benzoyl-CoA over 4-coumaroyl-CoA. However, the catalytic efficiency of the triple mutant was 10-fold lower than that of wild-type BPS (7).

In further rounds of mutations, BPS active site residues were replaced with amino acids present at the corresponding positions in type III PKSs other than CHS, such as acridone synthase and 2-pyrene synthase (2PS), which prefer bulkier (N-methylanthraniloyl) and smaller (acetyl) starter units, respectively (21, 22). In addition, a number of mutations were created on the basis of the CHS-based homology model of the BPS active site cavity complexed with the sequential reaction intermediates. However, the resulting enzyme mutants were either inactive, did not produce any soluble protein, or functionally resembled the wild-type enzyme, albeit with reduced catalytic activity. The only dramatic change in the functional behavior of BPS was observed with the substitution of the Thr-135 residue for a leucine. A threonine residue is also present in CHS and 2PS at the corresponding position, whereas acridone synthase has a serine.

Functional Characterization of the T135L Mutant—Wild-type BPS formed 2,4,6-trihydroxybenzophenone as a major product when incubated with benzoyl-CoA and malonyl-CoA, along with small amounts of 6-phenyl-4-hydroxy-2-pyrene (phenylpyrone) and traces of benzoyltriacetic acid lactone as derailment products (Fig. 3a). In contrast, the T135L mutant formed phenylpyrone as a major product and only traces of 2,4,6-trihydroxybenzophenone (Fig. 3b). The products were identified by GC-MS after silylation in comparison with authentic reference compounds.

Kinetic characterization of the wild-type enzyme and the T135L mutant demonstrated that the catalytic efficiencies

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*Corresponding active site cavity residues of crystallized alfalfa CHS2 (12) and *H. androsaemum* BPS*
The catalytic efficiency for benzoyl-CoA ($k_{cat}/K_m$) was 28,345 M$^{-1}$ s$^{-1}$, $k_{cat}$ = 9.66 min$^{-1}$, $K_m$ = 5.7 μM compared with the His$_6$-tagged protein used in this study (7). This remarkable discrepancy, mainly in turnover rate, was confirmed by careful reinvestigation of the kinetic properties of glutathione S-transferase-BPS and might be attributed to the effect of the different affinity tags on either protein stability or protein determination.

Wild-type BPS also accepted 3-hydroxybenzoyl-CoA as a starter molecule and formed 2,3',4,6-tetrahydroxybenzophenone as a major product and 6-(3'-hydroxyphenyl)-4-hydroxy-2-pyrone (hydroxyphenylpyrone) as a side product in the presence of malonyl-CoA (Fig. 3c). In contrast, the T135L mutant was almost inactive with 3-hydroxybenzoyl-CoA (Fig. 3d). It formed only trace amounts of hydroxyphenylpyrone, and 2,3',4,6-tetrahydroxybenzophenone was not observed. The wild-type enzyme and the T135L mutant were both inactive with 2- and 4-hydroxybenzoyl-CoA, acetyl-CoA, and the CoA thioesters of cinnamic acids (Table 3).

Interestingly, the replacement of the Thr-135 residue with a glycine, alanine, valine, isoleucine, asparagine, and tyrosine, instead of a leucine, gave catalytically inactive proteins. SDS-PAGE demonstrated that all of these mutant proteins were successfully expressed in E. coli and were highly purified by affinity chromatography. The mutants T135S and T135F functionally resembled the wild-type enzyme, albeit with reduced catalytic activities.

**Rationalization of the Functional Behavior of the T135L Mutant by Homology Modeling**—Wild-type BPS catalyzes the iterative condensation of benzoyl-CoA with three malonyl-CoAs to give a linear triketide that undergoes intramolecular C-6 → C-1 Claisen condensation to yield phlorbenzophenone (Fig. 4). In contrast, the T135L mutant catalyzes the addition of only two acetyl groups to the benzoyl starter unit. The triketide is the final linear intermediate and cyclizes into phenylpyrone via C-5 keto-enol oxygen → C-1 lactonization. These alternative reaction mecha-
TABLE 2  
Steady-state kinetic parameters for wild-type BPS and the T135L mutant

| Substrate     |  \( k_{\text{cat}} \) | \( K_m \) | \( k_{\text{cat}}/K_m \) | \( K_m \) |
|---------------|----------------------|---------|---------------------|--------|
| Benzoyl-CoA   | 3.31 \( \mu \) \text{m} | 1.4 \( \mu \) \text{m} | 4.8 \text{min} \^{-1} \text{m} \text{M} \^{-1} | 1.4 \( \mu \) \text{m} |
| Malonyl-CoA   | 0.41 \( \mu \) \text{m} | 0.7 \( \mu \) \text{m} | 0.58 \text{min} \^{-1} \text{m} \text{M} \^{-1} | 0.7 \( \mu \) \text{m} |

TABLE 3  
Substrate specificities of wild-type BPS and the T135L mutant

| Substrate       | Wild-type BPS | T135L mutant | % of maximum each |
|-----------------|---------------|--------------|------------------|
| Benzoyl-CoA     | 100           | 100          |                  |
| 3-Hydroxybenzoyl-CoA | 57            | 4            |                  |
| 2-Hydroxybenzoyl-CoA | 0             | 0            |                  |
| 4-Hydroxybenzoyl-CoA | 0             | 0            |                  |
| Cinnamoyl-CoA   | 0             | 0            |                  |
| 2-Coumaroyl-CoA | 0             | 0            |                  |
| 3-Coumaroyl-CoA | 0             | 0            |                  |
| 4-Coumaroyl-CoA | 0             | 0            |                  |
| Acetyl-CoA      | 0             | 0            |                  |

![Mutagenic Conversion of BPS into PPS](image)

**Mutagenic Conversion of BPS into PPS**

When 3-hydroxybenzoyl-CoA serves as a starter substrate, the wild-type enzyme catalyzes the stepwise addition of three acetyl groups to the starter unit inside the large initiation/elongation cavity to give a tetraketide, which cyclizes into 2,3',4,6-tetrahydroxybenzophenone (Fig. 5c). In the active site cavity of the T135L mutant, however, no tetraketide is formed. The arylo moiety of the triketide intermediate may enter the new pocket whose small size does not allow for a third chain extension reaction (Fig. 5d). Interestingly, the 3-hydroxyl group of the arylo moiety of the triketide intermediate resides in the position that is occupied in the wild-type enzyme by the hydroxyl group of the Thr-135 side chain. As a consequence, the 3-hydroxyl group of the triketide now forms hydrogen bonds with the backbone carbonyl and amide groups of Gly-166. Thus, the 3-hydroxybenzoyl-primed triketide may be trapped in the new pocket of the active site cavity of the T135L mutant. It acts as an inhibitor, explaining why the mutant enzyme is almost inactive with 3-hydroxybenzoyl-CoA as a starter substrate.

This conclusion was supported by inhibition experiments. The 3-hydroxybenzoyl-primed triketide as an inhibitor was formed by preincubation of the T135L mutant with malonyl-CoA and varying concentrations of 3-hydroxybenzoyl-CoA. After addition of benzoyl-CoA following preincubation, decreasing amounts of phenylpyrone correlated with increasing concentrations of 3-hydroxybenzoyl-CoA in preincubation (Fig. 6). In contrast, similar increases in phenylpyrone formation were observed after preincubation of the mutant enzyme with malonyl-CoA and increasing concentrations of acetyl-CoA, which is not accepted as a starter substrate, and malonyl-CoA only.
DISCUSSION

The functional behavior of BPS was dramatically altered by a single amino acid substitution in the active site cavity, which transformed BPS into PPS. The point mutation modulated between the formation of a tetraketide product derived from Claisen condensation and a triketide product derived from lactone formation. Although a number of functionally diverse type III PKSs have been detected and characterized (9), an enzyme accomplishing a complete derailment of the BPS reaction has not yet been identified. A small proportion of the BPS reaction intermediates on the path to benzophenone become permanently derailed, leading to the formation of phenylpyrone as a side product. Increasing amounts of this truncated reaction product were found with increasing dithiothreitol concentrations in enzyme assays (7). Similar lactones as by-products of in vitro reactions are also formed by other type III PKSs (9).

The condensation of benzoyl-CoA, which is a rare starter substrate for PKSs (23), with two acetyl units from malonyl-CoA is also catalyzed by 2PS from Gerbera hybrida (22). However, the physiological starter molecule for this enzyme is acetyl-CoA, resulting in the formation of 6-methyl-4-hydroxy-2-pyrone (methylpyrone), which is converted by downstream enzymes to gerberin and parasorboside. Unlike 2PS, PPS and BPS are inactive with acetyl-CoA, nor do they catalyze malonyl-CoA decarboxylation to generate acetyl-CoA, which can serve as starter substrate for methylpyrone formation (22). Acetyl-CoA is a central intermediate of primary metabolism, whereas benzoyl-CoA is a component of secondary metabolism. Benzoic acid biosynthesis branches from cinnamic acid from the general phenylpropanoid pathway (24).

Replacement of the polar residue Thr-135 of BPS with the apolar amino acid leucine in PPS may open the gate to a buried pocket, into which the triketide intermediate is redirected during the elongation process of the diketide. Compared with the initiation/elongation cavity of BPS, the size of the newly accessible pocket in PPS is smaller and does not allow for a third extension step with malonyl-CoA, thereby preventing the formation of a tetraketide. Kinetic characterization of BPS and PPS demonstrated that the T135L mutation increased the affinity for the starter substrate, benzoyl-CoA, but decreased the turnover number to the same extent, resulting in similar catalytic efficiencies ($k_{cat}/K_m$). In stilbene synthase (STS), the
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Recently, the polyketide elongation tunnel of the M207G mutant of PCS was further widened by the simultaneous replacement of the aromatic residues Phe-80 and Tyr-82 at the bottom of the new pocket with an alanine residue each (29). The resulting triple mutant formed low yields of a novel nonaketide naphthopyrone that is the longest polyketide chain formed by a type III PKS.

Further results that functionally link the size of the active site cavity to chain length determination were obtained with M. sativa CHS. The side chain volume of position 256 influenced both polyketide chain-length determination and intramolecular cyclization of the polyketide intermediates (30). Although the G256A and G256V mutations altered the regiospecific cyclization reaction, leading to increased coumaroyltriacetic acid lactone formation, the G256L and G256F mutations resulted in a derailment of the reaction at the triketide step, with bis-noryangonin and methylpyrone being released as products. When the corresponding residue of BPS (Ala-260) was replaced with valine, the phlorbenzophenone:phenylpyrone ratio was shifted from 1:0.08 (wild-type) to 1:0.39 (A260V). The A260L mutant was inactive with the starter substrates used.

The T135L mutation not only changed the product specificity but also the substrate specificity of BPS. 3-Hydroxybenzoyl-CoA is the second best starter substrate for wild-type BPS but a poor starter molecule for PCS. In this enzyme mutant, the 3-hydroxybenzoyl-primed triketide may be trapped in the new pocket by hydrogen bonds of the 3-hydroxyl group with the backbone, thereby acting as an inhibitor. Preincubation of the T135L mutant with malonyl-CoA and increasing 3-hydroxybenzoyl-CoA concentrations resulted in decreasing rates of phenylpyrone formation from benzoyl-CoA and malonyl-CoA. So far, changes in starter molecule selectivity of type III PKSs were attributed to alterations in the volume of the initiation/elongation cavity caused by specific amino acid substitutions. A combination of three mutations (T197L/G256L/S338I) converted M. sativa CHS into 2PS by reducing the volume of the active site cavity from 923 to 274 Å³ (13). The constricted catalytic centers of 2PS and the functionally identical CHS triple mutant sterically excluded the bulky 4-coumaroyl starter unit but allowed access of the smaller acetyl unit. When the same amino acid substitutions were introduced at the corresponding positions of BPS (T200L, G342I, and A260L), the resulting triple mutant was inactive with both benzoyl-CoA and acetyl-CoA, nor did the three single mutants use acetyl-CoA to form methylpyrone. Conversely, in M. sativa CHS the mutation of Phe-215, which is a gatekeeper to the active site, into a serine widened the active site entrance, allowing for productive binding of N-methylantraniloyl-CoA, which is too bulky to be accepted as a starter substrate by wild-type CHS (31). Contrary to acridone synthase that naturally uses N-methylantraniloyl-CoA as starter, the CHS F215S mutant did not catalyze the regiospecific Claisen cyclization reaction, resulting in release of N-methylantraniloyltriacetic acid lactone as a novel alkaloid.

Derivatives of the PPS product are known as natural products. A 4′-hydroxylated phenylpyrone named psilotinin was found in the pteridophyte Psilotum nudum, which is the most primitive vascular plant (32). cDNAs encoding CHS, STS, and...
valerophenone synthase were cloned from this species; however, a PPS cDNA was not identified (33). Feeding experiments suggested that psilotinin is formed by condensation of 4-coumaroyl-CoA with one molecule of malonyl-CoA followed by lactonization (34). The same one-step condensation is catalyzed by benzalacetone synthase that, however, decarboxylates the diketide intermediate (35). Recently, phenylpyrones have also been isolated from some Aloe species (36, 37). It will be interesting to isolate a PPS cDNA and to compare the structural and kinetic properties of the naturally occurring enzyme optimized by evolution with the T135L mutant created in vitro. PPS is a promising biotechnological tool for manipulating benzophenone and biphenyl biosyntheses in Clusiaceae and the Maloideae, respectively. The latter rosaceous subfamily includes economically important fruit trees, such as apple and pear. Metabolically engineered plants expressing PPS might produce new natural products as either constitutive metabolites or pathogen-induced phytoalexins.

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REFERENCES

1. Hu, L. H., and Sim, K. Y. (2000) Tetrahedron 56, 1379–1386
2. Yoshida, K., Tanaka, T., Hirose, Y., Yamaguchi, F., Kohno, H., Toida, M., Hara, A., Sugie, S., Shibata, T., and Mori, H. (2005) Cancer Lett. 221, 29–39
3. Hong, J., Sang, S., Park, H. J., Kwon, S. J., Suh, N., Huang, M. T., Ho, C. T., and Yang, C. S. (2006) Carcinogenesis 27, 278–286
4. Gronquist, M., Bezzeraides, A., Attygalle, A., Meinwald, J., Eisner, M., and Eisner, T. (2001) Proc. Natl. Acad. Sci. U.S.A. 98, 13745–13750
5. Peters, S., Schmidt, W., and Beerhues, L. (1998) Planta 204, 64–69
6. Zhang, H. Z., Kasibhatla, S., Wang, Y., Herich, J., Guastella, J., Tseng, B., Drewe, J., and Cai, S. X. (2004) Bioorg. Med. Chem. 12, 309–317
7. Liu, B., Falkenstein-Paul, H., Schmidt, W., and Beerhues, L. (2003) Plant J. 34, 847–855
8. Beerhues, L. (1996) FEBS Lett. 383, 264–266
9. Austin, M. B., and Noel, J. P. (2003) Nat. Prod. Rep. 20, 79–110
10. Schroeder, J. (1999) in Comprehensive Natural Products Chemistry (Sankawa, U., ed) Vol. 1, pp. 749–771, Elsevier Science Publishers B.V., Amsterdam
11. Helariutta, Y., Kotilainen, M., Elomaa, P., Kalkkinen, N., Bremer, K., Terei, T. H., and Albert, V. A. (1996) Proc. Natl. Acad. Sci. U.S.A. 93, 9033–9038
12. Ferrer, J. L., Jez, J. M., Bowman, M. E., Dixon, R. A., and Noel, J. P. (1999) Nat. Struct. Biol. 6, 775–784
13. Jez, J. M., Austin, M. B., Ferrer, J., Bowman, M. E., Schröder, J., and Noel, J. P. (2000) Chem. Biol. 7, 919–930
14. Vara Prasad, J. V. N., Lunney, E. A., Ferguson, D., Tummino, P. J., Rubin, J. R., Reynier, E. L., Stewart, B. H., Guttenrund, R. J., Domagala, J. M., Suvorov, L. I., Gulnik, S. V., Topol, I. A., Bhat, T. N., and Erickson, J. W. (1995) J. Am. Chem. Soc. 117, 11070–11074
15. Abdel-Mawla, A. M., Schmidt, W., and Beerhues, L. (2001) Planta 212, 288–293
16. Liu, B., Raeth, T., Beuerle, T., and Beerhues, L. (2007) Planta 225, 1495–1502
17. Guex, N., and Peitsch, M. C. (1997) Electrophoresis 18, 2714–2723
18. Arnold, K., Bordoli, L., Kopp, I., and Schwede, T. (2006) Bioinformatics 22, 195–201
19. Gerber, P. R., and Müller, K. (1995) J. Comput. Aided Mol. Des. 9, 251–268
20. Gerber, P. R. (1998) J. Comput. Aided Mol. Des. 12, 37–51
21. Junghans, K. T., Kneusel, R. E., Baumert, A., Maier, W., Gröger, D., and Matern, U. (1995) Plant Mol. Biol. 27, 681–692
22. Eckermann, S., Schröder, G., Schmidt, J., Strack, D., Edrada, R. A., Helariutta, Y., Elomaa, P., Kotilainen, M., Kilpelainen, L., Proksch, P., Terei, T. H., and Schröder, J. (1998) Nature 396, 387–390
23. Moore, B. S., and Hertweck, C. (2002) Nat. Prod. Rep. 19, 70–99
24. Abdel-Mawla, A. M., and Beerhues, L. (2002) Planta 214, 727–733
25. Austin, M. B., Bowman, M. E., Ferrer, J. L., Schröder, J., and Noel, J. P. (2004) Chem. Biol. 11, 1179–1194
26. Morita, H., Kondo, S., Oguro, S., Noguchi, H., Sugio, S., Abe, I., and Kohno, T. (2007) J. Am. Chem. Soc. 129, 359–369
27. Abe, I., Oguro, S., Utsumi, Y., Sano, Y., and Noguchi, H. (2005) J. Am. Chem. Soc. 127, 12709–12716
28. Abe, I., Watanabe, T., Lou, W., and Noguchi, H. (2006) FEBS J. 273, 208–218
29. Abe, I., Morita, H., Oguro, S., Noma, H., Wanibuchi, K., Kawahara, N., Goda, Y., Noguchi, H., and Kohno, T. (2007) J. Am. Chem. Soc. 129, 5976–5980
30. Jez, J. M., Bowman, M. E., and Noel, J. P. (2001) Biochemistry 40, 14829–14838
31. Jez, J. M., Bowman, M. E., and Noel, J. P. (2002) Proc. Natl. Acad. Sci. U.S.A. 99, 5319–5324
32. McInnes, A. G., Yoshida, S., and Towers, G. H. N. (1965) Tetrahedron 21, 2939–2946
33. Yamazaki, Y., Suh, D. Y., Sitthithaworn, W., Ishiguro, K., Kobayashi, Y., Shibuya, M., Ebizuka, Y., and Sankawa, U. (2001) Planta 214, 75–84
34. Leete, E., Muir, A., and Towers, G. H. N. (1982) Tetrahedron Lett. 23, 2635–2638
35. Abe, I., Takahashi, Y., Morita, H., and Noguchi, H. (2001) Eur. J. Biochem. 268, 3354–3359
36. Rebecca, W., Kayser, O., Hagem, H., Zessin, K. H., Madundo, M., and Gamba, N. (2003) Phytochem. Anal. 14, 83–86
37. Duri, L., Morelli, C. F., Crippa, S., and Speranza, G. (2004) Fitoterapia 75, 520–522