Expanding the chemical space of polyketides through structure-guided mutagenesis of Vitis vinifera stilbene synthase

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Several natural polyketides (PKs) have been associated with important pharmaceutical properties. Type III polyketide synthases (PKSs) that generate aromatic PK polyketides have been studied extensively for their substrate promiscuity and product diversity. Stilbene synthase-like (STS) enzymes are unique in the type III PKS class as they possess a hydrogen bonding network, furnishing them with thioesterase-like properties, resulting in aldol condensation of the polyketide intermediates formed. Chalcone synthases (CHS) in contrast, lack this hydrogen-bonding network, resulting primarily in the Claisen condensation of the polyketide intermediates formed. We have attempted to expand the chemical space of this interesting class of compounds generated by creating structure-guided mutants of Vitis vinifera STS. Further, we have utilized a previously established workflow to quickly compare the wild-type reaction products to those generated by the mutants and identify novel PKs formed by using XCMS analysis of LC-MS and LC-MS/MS data. Based on this approach, we were able to generate 15 previously unreported PK molecules by exploring the substrate promiscuity of the wild-type enzyme and all mutants using unnatural substrates. These structures were specific to STSs and cannot be formed by their closely related CHS-like counterparts.

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1. Introduction

Polyketides (PKs) are a chemically important class of compounds with several beneficial pharmaceutical properties [1,2]. Natural PKs generated by type III polyketide synthases (PKSs) have been associated with the slowing of the aging process in model organisms [3,4], anti-inflammatory and anti-cancer properties [5–10] and have shown potential to ameliorate diabetes and nervous system disorders related complications [11,12]. Type III PKSs are found in several plants, bacteria and fungi [13–15]. They are homodimeric enzymes that catalyze iterative condensation of repeating units to a CoA-tethered starter substrate through a conserved Cys-His-Asn catalytic triad. Functionally diverse type III PKSs arise due to variable substitutions in non-catalytic residues present in the three catalytically important cavities: the substrate binding pocket, composed of the important residues S133, Q192, T194, T197, S338; the CoA binding tunnel, composed of L55, R58, L62; and the cyclization pocket, composed of T132, M137, F215, I254, G256, F265, P375 (residues numbered according to the Vitis vinifera stilbene synthase (VvSTS)). Altering these residues results in diversity of preference for starter substrates, number of extender substrates incorporated through iterative condensations and mechanism of cyclization of the poly-β-keto intermediate formed through Claisen condensation, aldol condensation or lactonization. Moreover, type III PKSs possess unusually broad substrate promiscuity and can accept several non-natural substrates to form novel non-natural PKs [2,16]. Apart from the naturally occurring type III PKSs, several intuitive and structure-guided mutations have also been carried out to alter their enzymatic activity, so as to expand the chemical diversity of PKs [17,18]. Some of the novel non-natural PKs formed through these processes has also been demonstrated to possess important biological activities [16].

Abbreviations: PKs, polyketide synthase; PK, polyketide; Vv, Vitis vinifera; STS, stilbene synthase; CHS, chalcone synthase; WT, wild-type; HBN, hydrogen-bonding network; BNY, bisnoryangonin; CATL, p-coumarylauric acid lactone.

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Stilbene synthase (STS) belongs to the type III PKS superfamily and is unique due to the presence of a hydrogen-bonding networks (HBN) [19], which is absent in the closely related chalcone synthase-like (CHS) enzymes (70–90% sequence similarity to STS). Due to the presence of this HBN, STS-like type III PKSs cyclize their polyketide intermediate through an aldol condensation reaction. CHS-like type III PKSs, in contrast, cyclize their intermediate primarily through a Claisen condensation (Fig. 1). While both STS and CHS form a tetraketide intermediate with $\text{p}$-coumaroyl-CoA and three molecules of malonyl-CoA, STS cyclizes the intermediate into resveratrol through a C2–C7 aldol condensation in contrast to CHS, which cyclizes the intermediate into naringenin chalcone through a C1–C6 Claisen condensation. Both enzymes form bisnoryangonin (BNY) and $\text{p}$-coumaroyltriacetic acid lactone (CTAL) as derailment products during the reaction [20].

We attempted to exploit the novel thioesterase-like property of STSs to further diversify the chemical space of PKs. Along these lines we created 6 mutants of Vitis vinifera stilbene synthase (VvSTS) and challenged these with non-natural substrates (Table 1).

### 2. Materials and methods

#### 2.1. Site-directed mutagenesis

The plasmids expressing the his-tagged VvSTS and VvSTS T197G mutant were obtained from our previous studies [21].

![Fig. 1. General reaction catalyzed by wild type STS and CHS. STS and CHS react with their natural substrates, $\text{p}$-coumaroyl-CoA and malonyl-CoA to form primarily resveratrol and naringenin chalcone respectively. Bisnoryangonin (BNY) and $\text{p}$-coumaroyltriacetic acid lactone are natural derailment products of the reaction.](image)

**Table 1**

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Cavity 1 ($Å^3$)</th>
<th>Cavity 2 ($Å^3$)</th>
<th>Propionyl-CoA</th>
<th>Myristoyl-CoA</th>
<th>Octanoyl-CoA</th>
<th>Methylmalonyl-CoA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wt</td>
<td>721.4</td>
<td>265.2</td>
<td>1.2,3</td>
<td>4</td>
<td>5</td>
<td>6.7</td>
</tr>
<tr>
<td>L214I</td>
<td>705.7</td>
<td>291.4</td>
<td>1.2,3</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>T197A</td>
<td>743.1</td>
<td>293</td>
<td>2.8,9</td>
<td>10,11</td>
<td>5.12</td>
<td>6,13</td>
</tr>
<tr>
<td>T197G</td>
<td>1134.3*</td>
<td></td>
<td>1.14</td>
<td>4</td>
<td>15,16</td>
<td>6.7</td>
</tr>
<tr>
<td>T197G265L</td>
<td>612.4</td>
<td>265.2</td>
<td>1.2,8,9</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>T197M</td>
<td>635</td>
<td>265.2</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>7</td>
</tr>
<tr>
<td>T197G265L</td>
<td>1016.6*</td>
<td></td>
<td>–</td>
<td>–</td>
<td>5,12,16</td>
<td>–</td>
</tr>
</tbody>
</table>

* The T197G and the T197G265L mutations result in merging of the two cavities, we thus mention the size of the entire cavity in these cases.

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The rest of the VvSTS mutants were constructed for this study with a QuickChange Site-Directed Mutagenesis Kit (Stratagene) according to the manufacturer's protocol and using the primers in Supplementary Table S1. Each construct was sequence analyzed to confirm the point mutation. The double mutants were created sequentially.

2.2. Protein expression and purification

After confirmation of the sequence, the plasmid was transformed into *Escherichia coli* BL21*®* (DE3). The cells harboring the plasmid were cultured to an OD<sub>600</sub> of 0.6 in LB medium containing chloramphenicol (30 µg/ml) at 37 °C. Subsequently, isopropyl-β-D-galactopyranoside (IPTG) (1.0 mM) was added to induce protein expression, and the cells were further cultured at 30 °C for 4 h. All of the following procedures were performed at 4 °C. *E. coli* cells were harvested by centrifugation at 4000 × g and frozen at –80 °C until further processing. The cells were disrupted by incubating with lysis buffer (50 mM NaH<sub>2</sub>PO<sub>4</sub>, 300 mM NaCl, 10 mM imidazole, pH 8.0), lysozyme (10 mg/ml) and 1 culture volume of Benzonase<sup>®</sup> nuclease (3 units/ml) at 4 °C for 30 min. The lysate was then centrifuged at 12,000 × g for 30 min. The supernatant was loaded onto a Ni-NTA spin column (Qiagen) pre-equilibrated with lysis buffer. The column was then washed with 50 mM NaH<sub>2</sub>PO<sub>4</sub>, 300 mM NaCl, 20 mM imidazole, pH 8.0. The protein was eluted from the column using elution buffer (50 mM NaH<sub>2</sub>PO<sub>4</sub>, 300 mM NaCl, 300 mM imidazole, pH 8.0). The protein concentration was determined by the Bradford method (Protein Assay, Bio-Rad) with bovine serum albumin as the standard.

2.3. Enzymatic reaction

The reaction mixture contained starter substrate [4-coumaroyl-CoA (54 µM), octanoyl-CoA (54 µM), methylmalonyl-CoA (54 µM), or myristoyl-CoA (54 µM)], extender substrate [malonyl-CoA (108 mM) or 13C<sub>3</sub>-malonyl-CoA (108 mM)], and the purified enzyme (20 µg) in potassium phosphate buffer (500 µl, 100 mM, pH 7.0). The purified enzyme was added to the reaction mixture last. Incubations were performed at 30 °C for 30 min and were stopped by the addition of ethyl acetate (500 µl) with 1% HCl. The extracted products (ethyl acetate extracts) were then concentrated in a speed vacuum and re-suspended in ethyl acetate (10 µl), and centrifuged at 14000 rpm for 10 min before running on a column. The products were separated by reverse-phase HPLC (Agilent 1260) on a Zorbax C18 column (4.6 × 150 mm, 5 µm, at a flow rate of 0.7 ml/min). Gradient elution was performed with H<sub>2</sub>O and acetonitrile (ACN), both containing 0.2% trifluoroacetic acid: 0–7 min, 20% ACN; 7–15 min, linear gradient from 20% to 60% ACN; 15–30 min, linear gradient from 60% to 70% ACN; 30–36 min, linear gradient from 70% to 30% ACN. Three reactions (technical replicates) were pooled into ethyl acetate (10 µl) for LC-MS analysis. Online HR-ESI-LCMS spectra were measured with an Agilent Technologies HPLC 1200 series HPLC coupled to a Thermo Scientific LTQ Orbitrap XLITMS mass spectrometer fitted with an electrospray ionization (ESI) source. The ESI capillary temperature and the capillary voltage were 320 °C and 4.0 V, respectively. The tube lens offset was set at 20.0 V. All spectra were obtained in the negative and positive mode, over a mass range of m/z 150–700, and at a range of one scan every 0.2 s. The collision gas was helium, and the relative collision energy scale was set at 30.0% (1.5 eV). Dependent MS/MS scans were acquired for the first four most abundant parent ions.

2.4. Substrates

4-Coumaroyl-CoA was chemically synthesized as previously described [22]. 13C<sub>3</sub>-malonyl-CoA, malonyl-CoA, propionyl-CoA, octanoyl-CoA, methylmalonyl-CoA, myristoyl-CoA were purchased from Sigma.

2.5. XCMS workflow for analyzing online LCMS data

The workflow established included creating the structure-based mutants and carrying out the *in vitro* enzymatic re-actions. The *in vitro* reactions were run with and without the 13C<sub>3</sub>-malonyl-CoA for easier identification of novel PK structure from the HR-LC-MS/MS analysis. The reaction extracts were then run on the LC-MS to obtain online HR-LCMS (±5 ppm accuracy) which was finally compared using XCMS to directly calculate the fold change in production of PKs in the mutants compared to the wild-type and to identify peaks unique to the mutant. Raw data was converted to XML format using R processor (Script S1, obtained from http://www.metabolomics.strath.ac.uk/showPage.php?page=processingscripts), and analyzed using XCMS online [23]. XCMS was utilized for calculating fold changes in CTAL and BNY and all other peaks common to the wild type and mutant proteins (Fig. 6). All calculations were based on triplicates. This workflow obviated thin-layered chromatography (TLC) analysis of the reaction products and utilization of radioactive substrates for identification of products unique to the mutants.

2.6. Homology modeling and cavity size analysis

The models of the Wt VvSTS and all the mutants were generated by the SWISS-MODEL package (http://expasy.ch/spdbv/) provided by the Swiss-PDB-Viewer program [24] based on the crystal structure of wild-type STS from *Acharis hypogea* (PDB code: 1Z1E). The model quality was assessed using PROCHECK [25]. In the Ramachandran plot calculated for the model, most of the amino acid residues were present in the energetically allowed regions with only a few exceptions, primarily Gly residues that can adopt phi/psi angles in all four quadrants. The cavity volume was calculated by the program CASTP (http://cast.engr.uic.edu/cast/) [26].

3. Results

3.1. Substrate promiscuity of wild-type VvSTS

Several non-natural substrates have been supplied to wild-type *Acharis hypogea* STSs, resulting in the formation of mainly CATL and BYN-type pyrones [27,28]. We first investigated the substrate promiscuity of wild-type VvSTS (Wt VvSTS) with non-natural substrates that have not been tested previously. Specifically, we supplied Wt VvSTS with starter substrates of varying size: propionyl-CoA, myristoyl-CoA, octanoyl-CoA and methylmalonyl-CoA; malonyl-CoA was used primarily as the extender substrate for all of the reactions (Fig. 2).

Malonyl-CoA and methylmalonyl-CoA or propionyl-CoA acted as extender substrates in case of compounds 2, 3 & 7, as is clearly indicated in the color-coding in Fig. 3. The Wt VvSTS accepted all of these aliphatic compounds as substrates to afford non-natural PKs (Fig. 3).

Propionyl-CoA and malonyl-CoA resulted in the formation of 3 major products. Compound 1 with the parent ion peak [M + FA–H<sup>-</sup>] at m/z 241.0700, a resorcinol derivative. Compound 2 with parent ion peak [M–H<sup>-</sup>] at m/z 263.1290 and 3 with m/z 323.1392, a
Myristoyl-CoA and malonyl-CoA formed carboxylic acid 4, which displayed a parent ion peak \([\text{M} + \text{FA-H}]^-\) at \(m/z\) 293.1756 (Fig. 3, Supplementary Table S2). Octanoyl-CoA and malonyl-CoA resulted in the formation of primarily a phlorophenone (5) with the parent ion \([\text{M} - \text{H}]^-\) at \(m/z\) 209.1185 (Fig. 3, Supplementary Table S2). Methylmalonyl-CoA and malonyl-CoA resulted in the formation of two non-natural PKs, a pentanone 6 at \(m/z\) 263.1274 \([\text{M} - \text{H}]^-\) and a pyranone 7 at \(m/z\) 307.1292 \([\text{M} + \text{COO}^-]\) (Fig. 3, Supplementary Table S2).
3.2. Substrate promiscuity of structure-based mutants of VvSTS

Next we tested the above substrates with the structure-based mutants of VvSTS. As hypothesized, perturbation of residues in the catalytically relevant pockets of VvSTS altered its substrate promiscuity. We supplied relevant substrates to each mutant based on predicted change in the properties of the catalytically relevant pockets; so all substrates were not supplied to all mutants. The mutants created led to the formation of novel, non-natural PKs that have not been reported previously (Figs. 4 and 5).

The leucine at position 214 is important for the structure of the cyclization pocket. Not surprisingly, replacing it with isoleucine (L214I) did not significantly alter the product profile. Compared to Wt VvSTS, mutant L214I led to a 14.4-fold, 27-fold, and 9.4-fold increase in production of compounds 1, 2, and 3, respectively, when propionyl-CoA was used as a starter substrate (Fig. 3, Supplementary Fig. S1). This could be explained by the increase in the starter substrate binding pocket from 265.2 Å³ for Wt VvSTS to 291.4 Å³ in the L214I mutant, making it easier for the mutant to utilize propionyl-CoA as the substrate. The decrease in the cyclization pocket from 721.4 Å³ to 705.7 Å³ explains the absence of formation of any longer PKs.

Upon increasing the size of the cyclization pocket from 721.4 Å³ to 743.1 Å³ and the substrate-binding pocket from 265.2 Å³ to 293 Å³ by replacing the tyrosine at position 197 with an alanine, VvSTS T197A catalyzed formation of 5 new PKs unique to the mutant. Propionyl-CoA and malonyl-CoA resulted in the formation of two novel compounds, a benzoic acid with [M−H]⁻ m/z of 279.0858 (8) and a phlorophenone with m/z 233.0811 [M−2H]⁻ (9) (Fig. 4). In both 8 & 9 propionyl-CoA was utilized as an extender substrate as well. A 1.2-fold increase in the production of the 2263.1290 with m/z 331.2014 [M + CH₃COO]⁻ was observed (Supplementary Fig. S2). Upon supplying myristoyl-CoA and malonyl-CoA as the starter substrate and the extender substrate, respectively, two unique, novel phlorophenones were formed: 10 with a parent ion peak [M−H]⁻ at 335.20, and 11 with a parent ion peak of [M + FA−H]⁻ at 337.2016 (Fig. 4). Octanoyl-CoA and malonyl-CoA were also utilized by T197A to form 5 but 30-fold lower than the amount formed by the wild-type (Supplementary Fig. S2). Instead, the novel phlorophenone 12 with m/z [M−H]⁻ of

![Chemical structures](image_url)

**Fig. 4. Non-natural PKs (8–13) formed by mutant VvSTS.** Novel PKs formed by the mutants when supplied with propionyl-CoA (compounds 8 and 9), myristoyl-CoA (compounds 10 and 11), octanoyl-CoA (compound 12) and methylmalonyl-CoA (compound 13) as starter substrates, coupled with malonyl-CoA primarily as the extender substrate.
Next we created a double mutant, replacing the threonine at position 197 to an isoleucine and the glycine at position 256 to a leucine (T197I256L). This resulted in a decrease of the cyclization pocket from 721.4 Å³ to 635 Å³ without significant change in the substrate binding pocket size. We did not carry out the in vitro reaction with myristoyl-CoA or octanoyl-CoA as we did not expect the mutant to form any novel PKs with these bulkier substrates. Propionyl-CoA and malonyl-CoA were utilized as substrates by T197I256L to form 1 and 2 at 6-fold and 2-fold lower levels, respectively, than the wild-type (Supplementary Fig. S4). Moreover, compounds 8 and 9, formed by T197A, were also formed by T197I256L. The mutant did not form any new PKs with methylmalonyl-CoA and malonyl-CoA as substrates.

Replacing the threonine at position 197 with the bulkier methionine (T197M) resulted in a reduction of the substrate-binding pocket from 721.4 Å³ to 635 Å³. When we supplied methylmalonyl-CoA and malonyl-CoA as substrates, compound 7 was formed, however a 2.3-fold decrease in comparison to the Wt VvSTS was observed (Supplementary Fig. S4).

Upon creating the double mutant by replacing threonine at position 197 with a glycine and the glycine at position 256 with a leucine (T197G256L), the homology model again predicted the merging of the cyclization and substrate binding pockets, except this time with a decreased volume of 1016.6 Å³ as compared to 1029.2 Å³ for Wt VvSTS. Surprisingly supplementation with propionyl-CoA and malonyl-CoA did not result in the formation of any significant products. Octanoyl-CoA and malonyl-CoA resulted in the formation of 12 and 5, and 16, however 15 was not formed in this case. Both 12 and 15 are formed from the same 18-carbon intermediate. Compound 12 is a derailment product, thus it is a pyrone, while formation of 15 requires cyclization by the enzyme. The absence of 15 can be explained by the possible incapability of the T197I256L mutant to successfully cyclize the longer PK intermediate (18 carbons). This is further supported by the significant increase in the amount of the derailment product 12 formed.

3.3. XCMS work-flow for analyzing the in vitro enzymatic reactions

We utilized a previously established workflow [21] for quick identification of non-natural PKs by the 6 mutants of VvSTS created in this study (Fig. 6). Briefly, enzymatic reactions were analyzed by high resolution LC-MS (HR-LCMS), and novel reaction products were automatically tabulated by comparison against the product profile of a control reaction using an online LC-MS data comparison analysis software known as XCMS [23,29]. Structural elucidation of novel reaction products was further aided by performing paired reactions, where one reaction used unlabeled malonyl-CoA and the other utilized stable-isotope labeled 13C3-malonyl-CoA extender units. Novel parent ions were subjected to HR-LCMS/MS, and comparison of fragmentation patterns between paired reactions enabled identification of novel products by tracking the additional mass conferred by incorporation of 13C3-malonyl-CoA extender units.

4. Discussion

We have successfully diversified the PK space by creating structure-guided mutants of VvSTS, and supplying it with non-natural substrates. Specifically we altered the size of the substrate binding pocket (T197), the cyclization pocket (L214) and both (T197 & G256).

The data obtained demonstrated that wild-type VvSTS could utilize methylmalonyl-CoA, propionyl-CoA, octanoyl-CoA and myristoyl-CoA as starter substrates coupled with malonyl-CoA primarily as the extender substrate; of the 7 non-natural PKs generated by wild-type VvSTS, 6 have not previously been reported.
Several other possible combinations of substrates, with a range of different sizes and polarities, could be tested, to afford a comprehensive expansion of the PK space, although testing all of these possible combinations is beyond the scope of this work. Upon feeding non-natural substrates to the different mutants we were able to further diversify the PK space by generating 9 additional non-natural PKs that have not been previously reported. All of the PKs characterized in this study are unique to STS-like enzymes, establishing STS as a candidate enzyme for future protein engineering efforts and as a tool for generation of libraries of novel PKs with potential therapeutic value. Finally, XCMS analysis was utilized for quick identification of PKs that were formed only by the mutants, and the established workflow aided in simple and rapid identification of all 15 novel, non-natural PKs directly from the online HR-LCMS data.

We propose that this quick identification workflow can be utilized for generation of many more novel PKs. In the future, supplementation with nitrogen-containing substrates could help generate novel non-natural alkaloid analogs. Further optimization of the biosynthetic production of these novel PKs can be carried out, as has been previously reported [30–35].

**Conflict of interest**

None.

**Acknowledgments**

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**Appendix A. Supplementary data**

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.biochi.2015.05.019.

**References**


