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Establishment of strigolactone-producing bacterium-yeast consortium

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Strigolactones (SLs) are a class of phytohormones playing diverse roles in plant growth and development, yet the limited access to SLs is largely impeding SL-based foundational investigations and applications. Here, we developed Escherichia coli—Saccharomyces cerevisiae consortia to establish a microbial biosynthetic platform for the synthesis of various SLs, including carlactone, carlactonoic acid, 5-deoxystrigol (5DS; 6.65 ± 1.71 µg/liter), 4-deoxyorobanchol (4DO; 19.36 ± 5.20 µg/liter), and orobanchol (OB; 19.36 ± 5.20 µg/liter). The SL-producing platform enabled us to conduct functional identification of CYP722Cs from various plants as either OB or 5DS synthesize. It also allowed us to quantitatively compare known variants of plant SL biosynthetic enzymes in the microbial system. The titer of BDS was further enhanced through pathway engineering to 47.3 µg/liter. This work provides a unique platform for investigating SL biosynthesis and evolution and lays the foundation for developing SL microbial production process.

INTRODUCTION

Strigolactones (SLs) were initially characterized as signaling molecules, which are released from plant roots, induce germination of root parasitic weed, regulate the hyphae branching of arbuscular mycorrhiza fungi (AMF), and promote the symbiotic relationship between plants and fungi (1, 2). Later, they were also identified as a previously unknown class of plant hormones that control shoot branching, leaf growth, and senescence and promote the formation of lateral root and growth of primary root (3). SLs thus have been considered as promising agrochemicals, such as biopesticides that enhance the nutrient uptake efficiency through modulating plant-AMF symbiotic association (4–6). To date, more than 30 natural SLs have been isolated (fig. S1) (7, 8). SLs generally consist of a conserved butenolide ring (D ring) connected to a less conserved tricyclic lactone ring via an enol-ether bond (Fig. 1) (9). They can be classified into canonical and noncanonical SLs: The canonical SLs contained the tricyclic lactone-ring (ABC ring), while the noncanonical SLs lack the tricyclic ring scaffold with one (C ring) or two rings (B ring and C ring) missing (10). The canonical SLs can be further subdivided into orobanchol (O)- and strigol (S)-type SLs according to the stereochemistry in the C ring, which are represented by 4-deoxyorobanchol (4DO) and 5-deoxystrigol (5DS), respectively (9). Some of the better-known noncanonical SLs include methylcarlactonoate (MeCLA) (11), heliolactone (12), avenaol (13), zealactone (14), and lotuslactone (15).

SLs are derived from all-trans-β-carotene (ATβC), which is converted to carlactone (CL), the key branching point in SL biosynthesis (16), by the functions of three plastid enzymes: the isomerase DWARF27 (D27), carotenoid cleavage dioxygenases and 7 and 8 (CCD7 and CCD8) (Fig. 1) (16). DWARF27, a [2Fe-2S]–containing polypeptide, catalyzes the isomerization of ATβC to 9-cis-β-carotene (9CβC) (16, 17), followed by CCD7, a non–heme iron–dependent enzyme that catalyzes the C9′-C10′ double bond cleavage of 9CβC to yield 9-cis-β-apo-10′-carotenal (9CβA10′CAL) and β-ionone (βI) (16, 18, 19). Subsequently, CCD8 further catalyzes the oxidative cleavage of 9CβA10′CAL to synthesize CL, with the reaction mechanism remaining elusive (16, 20). CL is then exported into cytoplasm and further oxidized by cytochrome P450s (CYPs) and other oxidases to afford various SL structures (21). The first oxidation step has been characterized to be the C19 oxidation of CL to synthesize carlactonoic acid (CLA), which can be catalyzed by the MORE AXILLARY GROWTH 1 (MAX1), a member of the CYP711A subfamily (11).

MAX1 is conserved in a number of plant species (22) and has many homologs. Rice, known to produce O-type SL such as 4DO and orobanchol (OB) (23), encodes five MAX1 homologs. One MAX1 homolog, OsCYP711A2 encoded by Os900, was identified to catalyze the conversion of CL to 4DO likely via CLA (22, 24). The B ring of 4DO can be further oxidized by MAX1 analogs (OsCYP711A3 encoded by Os1400 from rice and ZmMAX1b from maize) to afford OB (22). However, 4DO is not produced in many OB-producing plants (e.g., cowpea), which hints another synthetic route of OB without involving 4DO (25, 26). This direct synthesis of OB from CL was later identified to be catalyzed by CYP722C in cowpea (27). Recently, a CYP722C from cotton was characterized to catalyze the synthesis of 5DS from CLA (28).

Biosynthesis of most other SLs have not been fully elucidated. Studying these pathways could be sped up if total biosynthesis of key SLs and pathway intermediates (e.g., CL, OB, 5DS) from sugar can be established in fast-growing microbial hosts. In situ–produced SLs and pathway intermediates in the microbial system can serve as substrates to characterize recombinant SL biosynthetic enzyme candidates, eliminating the need of acquiring these molecules from plant materials or chemical synthesis. On the other hand, much remains to be investigated to understand the evolutionary history of SL biosynthesis and signaling in plants, especially nonvascular plants. The SL biosynthetic genes are either missing from the nonvascular plants or quite distinct from the corresponding variants encoded by the vascular plants (29). For example, although Charales (green algae) have been reported to produce SLs (e.g., sorgolactone) (30), no close CCD7 and CCD8 analogs have been found from either expression profiles or genomes, whether Charales can synthesize SL or not remains in debate (29).
An efficient functional identification strategy aided by a microbial SL-producing platform can also advance understanding of SL origin and evolution.

In this study, we first attempted to establish the SL biosynthetic pathway in \textit{Saccharomyces cerevisiae} but failed, possibly due to the unsuccessful functional reconstitution of D27 and CCD7. We then tested the use of \textit{Escherichia coli} as the host, since the active form of D27, CCD7, and CCD8 have been heterologously expressed and isolated from \textit{E. coli} before (16, 18–20, 31). An engineered \textit{E. coli} strain successfully produced CL but failed to further functionalize it. The challenge was later overcome by using mixed cultures of \textit{E. coli} and \textit{S. cerevisiae}. The total biosynthesis of OB, 4DO, and 5DS was achieved from xylose. We used this microbial biosynthetic platform to identify eight CYP722Cs that produced either OB or 5DS from CLA and established a sequence-function correlation that could be used to predict the function of unknown CYP722Cs. Next, we chose 5DS as a model SL and improved the titer in the microbial consortium by 220\% to 47.3 g/liter (shake flask culture) through metabolic engineering. The improved titer could be useful to isolate adequate pathway intermediates in future studies. Last, we quantitatively compared variants of D27, CCD7, and CCD8 from different plant species in the optimized system.

**RESULTS**

**It is challenging to establish CL production in \textit{S. cerevisiae}**

To synthesize CL, we first attempted to functionally reconstitute D27 and CCD7 in a AT\textbeta{}C-producing \textit{S. cerevisiae} strain. We reconstructed the AT\textbeta{}C-producing yeast strain (YYL23; table S2) as previously described (32). Trace amount of 9C\textbeta{}C naturally existed in the established AT\textbeta{}C-producing yeast strain with a ratio to the AT\textbeta{}C approximately 1:25 (fig. S2A). Expressing D27 from \textit{Arabidopsis thaliana} (ATD27) or rice (OsD27) did not increase the ratio (fig. S2A). We extensively explored commonly used strategies for functionally reconstituting D27 (protein N-terminal engineering, protein localization to mitochondria, improving [2Fe-2S] cluster biogenesis, and reducing oxidative stress; fig. S2, A to C), but without success. We also tried to reconstitute the function of CCD7 from \textit{A. thaliana} due to the presence of the substrate of CCD7, 9C\textbeta{}C, although at a low titer in YYL23. Although CCD1 has been functionally expressed in yeast to cleave \textbeta{}-carotene to afford the synthesis of 13C, we did not detect any activity of CCD7 possibly due to the fact that the CCD1 is natively localized in cytosol, while the other CCDs are in plastid (fig. S2D) (34). Truncating N terminus of CCD7 and/or targeting it to yeast mitochondria, which resemble plastids to certain extent, did not lead to detection of CCD7 activity in the yeast (fig. S2D).

**Establishment of CL production in \textit{E. coli}**

Previous investigations indicate that D27, CCD7, and CCD8 can be expressed and isolated in soluble form from \textit{E. coli} for the in vitro biochemical investigations (16, 18–20, 31). Thus, we shifted the CL production from yeast to \textit{E. coli}. First, OsD27 was expressed from a medium copy number plasmid \textit{pCDFDuet-1} (table S1) in \textit{E. coli}, under the control of \textit{T7} promoter, in the presence of the well-documented AT\textbeta{}C-producing plasmid \textit{pAC-BETAipi} (35) (resulting strain ECL-2, \textit{E. coli} harboring \textit{pAC-BETAipi} to produce AT\textbeta{}C is designated as ECL-1; table S2). Upon the introduction of OsD27, the ratio between 9C\textbeta{}C to AT\textbeta{}C was increased from 1:4.1 to 1:1.3, which indicates the functional reconstitution of OsD27 in \textit{E. coli} (fig. 2A and fig. S3, A and B).
In the presence of OsD27, the titer of 9CβC was 1.41 mg/liter. The activity of D27 from A. thaliana (AtD27) was also examined in ECL-1 (ECL-2′; table S2), which exhibited a lower activity than OsD27 (fig. S4). We truncated the putative plastid transit peptide (first 40 amino acids) from OsD27 but did not detect obvious enhancement in the conversion toward 9CβC (fig. S4). Previous investigation on D27 indicated that this β-carotene isomerase catalyzes reversible conversion between ATβC and 9CβC (31), and thus, the ratio between 9CβC to ATβC in ECL-2 might have reached the equilibrium.

Subsequently, the N terminus of CCD7 from A. thaliana (AtCCD7) was truncated [first 31 residues (31), trAtCCD7] and introduced to 9CβC-producing ECL-2 from the same plasmid pCDFDuet-1—expressing OsD27, under the control of T7 promoter (ECL-3; table S2). Liquid chromatography–mass spectrometry (LC-MS) analysis indicated the synthesis of a compound with a mass/charge ratio (m/z) positive = 379.3 that agrees with 9-cis-β-apo-10′-carotenol (9CβA10′COL) (Fig. 2, B and C, and fig. S3C) (16, 18, 19). Although the product of CCD7 in vitro is 9CβA10′COL (36), the aldehyde can be reduced to the corresponding alcohol, 9CβA10′COL, in E. coli (18, 19). The synthesis of βI, the other product of CCD7, was confirmed through comparison with the authentic standard (figs. S3D and S5). No 9CβA10′COL (Fig. 2C) but a trace amount of βI was detected in ECL-2 (fig. S5), and βI is believed to be produced by the autoxidation of ATβC (36).
The N-terminal truncated (first 56 residues) AtCCD8 (trAtCCD8) was then expressed in the 9CβA10′-COL-producing ECL-3 from a medium copy number plasmid pET21a under the control of T7 promoter (ECL-4; table S2). Although a marked decrease in the synthesis of 9CβA10′-COL was detected upon the introduction of trAtCCD8, we were unable to detect the synthesis of CL (fig. S6, A and B). Previous reports imply that CL was unstable (31, 37) and that pH, solvent composition, and temperature can affect its chemical stability (37, 38). We added 200 mM Hepes buffer (pH 7.0) to the growth medium upon isopropyl β-D-1-thiogalactopyranoside (IPTG) induction (buffer volume/medium volume = 0.5), which led to the detection of a tiny peak in chromatograms of both the cell pellets and the culture medium at RT = 20.58 min with m/z = 303.2, which agrees with that of CL (figs. S3E and S6, C and D). However, the yield of CL was low (fig. S6, C and D) and was not proportional to the decrease in the synthesis of 9CβA10′.CL. Thus, we tested different medium conditions for the detection of CL. Under the optimal fermentation conditions (XY medium; fig. S6, G), a distinguished peak with the maximum absorption at 269 nm and m/z consistent with those of CL at the same retention time as the putative CL was observed (Fig. 2, D and E, and fig. S3E (16, 31).

**Synthesis of CLA in E. coli–S. cerevisiae coculture**

According to the pioneering in planta study, most of the canonical SLs are branched from CLA, which is synthesized from CL with the function of MAX1 (11). Although E. coli does not contain membrane-bound organelles, many plant cytochrome P450s have been functionally reconstitute in E. coli (39). However, further introduction of truncated MAX1 and cytochrome P450 reductase I from A. thaliana (40) (AtMAX1 and ATR1, respectively) in the CL-producing ECL-4 did not convert CL toward CLA (ECL-5; table S2 and fig. S8). Considering AtMAX1 is an endoplasmic reticulum (ER)-localized enzyme, the eukaryotic model species S. cerevisiae may be a more suitable organism to reconstitute the activity of the ER-localized CYP than E. coli (39). Since CL can be detected in both cell pellets and culture medium, it is possible to establish the synthesis of downstream SLs using an E. coli–S. cerevisiae coculture, with which CL is expected to be translocated from E. coli to yeast for further functionalization.

AtMAX1 and ATR1 were then introduced to S. cerevisiae on two low copy number plasmids and expressed downstream of PGK1 and TEF1 promoter, respectively (resulting strain: YSL-1; table S2). When the CL-producing E. coli strain (ECL-4) was cocultured with YSL-1, the peak of CL substantially decreased, and a new compound was detected in the organic extract of both cell pellets and medium under ultraviolet (UV) detection (Fig. 3A). The UV-visible (UV-VIS) spectrum of the peak had the maximum absorption at 271 nm, with [M-H]− = 331.1 matching those of CL at the same retention time as the putative CL (consistent with those of 18-hydroxy-CLA previously reported (27)). HR-MS analysis confirmed that OB can be generated through two different routes using the authentic OB standard; Fig. 3D and figs. S3I and S11).

**Synthesis of OB using CYP722C in E. coli–S. cerevisiae coculture**

Previous investigations confirmed that VuCYP722C from cowpea can directly convert CLA into OB and its diastereomer ent-2′-epi-OB by in vitro experiments (27). To establish this OB biosynthetic route in the microbial system, we introduced VuCYP722C to YSL-1 on a low copy number plasmid downstream of the GPD promoter (YSL-4; table S2), then cocultured YSL-4 with the CL-producing ECL-4. The ECL-4/YSL-4 coculture produced much less CLA than the ECL-4/YSL-4/5N coculture (YSL-4/5N is equivalent to YSL-1 but harboring an additional empty vector; table S2) coculture, the ECL-4/YSL-3/N coculture (YSL-3/N is a low copy number plasmid downstream of PGK1 promoter; table S2), then cocultured YSL-3/N with the CL-producing ECL-4, which produced CLA at a lower concentration (YSL-3/N; table S2). Despite this, we were able to detect the putative new compound at m/z = 347.1 in the negative ion mode (figs. S3H and S10B) and is putatively 18-hydroxy-CLA (27). HR-MS analysis confirmed that [M-H]− = 347.1521, and the fragmentation patterns are both consistent with those of 18-hydroxy-CLA previously reported (C19H24O6; fig. S7B) (27).

Furthermore, we introduced Os1400 (gene of OsCYP711A3) to YSL-2 on a low-copy plasmid driven by GPD promoter (YSL-3; table S2). As expected, compared to the ECL-4/YSL-3N (YSL-3N is equivalent to YSL2 but harboring an additional empty vector; table S2) coculture, the ECL-4/YSL-3 coculture synthesized substantially less 4DO and OB (0.75 ± 0.01 µg/liter (identified and quantified by using the authentic OB standard; Fig. 3D and figs. S3I and S11)). Different from 4DO detected in both pellets and medium, OB mostly in the medium indicates that most OB synthesized was exported into the medium. The successful reconstitution of OB synthesis from CL in the microbial consortium using OsCYP711A2 and OsCYP711A3 confirmed the previously proposed synthetic pathway of OB in rice (24).

**Synthesis of OB through 4DO using CYP711As in E. coli–S. cerevisiae coculture**

In rice, the two MAX1 homologs, OsCYP711A2 and OsCYP711A3, were identified to catalyze the conversion of CL to 4DO and 4DO to OB, respectively (22, 24). To synthesize 4DO in the microbial system, we expressed Os900 (OsCYP711A2 gene) using a low copy number plasmid downstream of PGK1 promoter in a yeast strain harboring ATR1 (YSL-2; table S2). When ECL-4 was cocultured with YSL-2, a peak with m/z+ at 331.1 (consistent with either 5DS or 4DO) was detected (Fig. 3C), which was further confirmed to be 4DO through comparison with the authentic 4DO and 5DS standards (Fig. 3C and figs. S3G and S10, A to C). The titer of 4DO in the consortium was 3.46 ± 0.28 µg/liter. In addition to 4DO, we were also able to detect the synthesis of another new compound in comparison to the negative control (without OsCYP711A2) in the organic extract of the medium. The putative new compound showed m/z at 275.1 in the negative ion mode (fig. S3I and S10B and is putatively 18-hydroxy-CLA (27). HR-MS analysis confirmed that [M-H]− = 347.1521, and the fragmentation patterns are both consistent with those of 18-hydroxy-CLA previously reported (C19H24O6; fig. S7B) (27).
Different from OB, 5DS was detected in the extract of both medium and cell pellet, which aligns with the lower hydrophilicity of 5DS than OB (fig. S13). Only a small quantity of CLA was detected (fig. S13A), suggesting the high efficiency of GaCYP722C. The putative 18-hydroxy-CLA was also detected in the medium of the 5DS-producing consortium (fig. S13B). This peak exists in the medium of all the canonical SL-producing consortiums we constructed, which supports the hypothesis that 18-hydroxy-CLA is a common intermediate in canonical SLs biosynthesis.

**Functional mapping of various CYP722Cs**

CYP722C genes are widely distributed in flowering plants. GaCYP722C share 65% amino acid identity with VuCYP722C, yet they catalyze different reactions. It is intriguing to investigate whether the enzymatic function of homologous proteins is conserved across different plant species. The successful functional reconstitutions of GaCYP722C and VuCYP722C in the microbial consortium hint the potential of using this biosynthetic platform to establish a sequence-function correlation of CYP722Cs, which will enable predicting the function of unknown CYP722Cs. We used GaCYP722C protein sequence as
The CYP722C characterization results are consistent with the version of CLA or synthesis of 5DS or OB (table S2 and fig. S14). We also included two CYP722Cs with the functions to be confirmed [SICYP722C (27) and LjCYP722C (43)] and CYP722A and CYP722B sequences as the outgroup (Fig. 4).

Phylogenetic analysis indicated that CYP722C subfamily can be divided into two groups (Fig. 4 and fig. S15): group I and group II. The characterized OB-producing VuCYP722C and SICYP722C are members of group I. The speculative 5DS synthase LjCYP722C and characterized 5DS-producing GaCYP722C are members of group II. To examine whether we can simply predict the function on the basis of the phylogenetic analysis, we synthesized eight CYP722C genes (table S3) from different branches and screened their functions by introducing each gene to the CLA-producing microbial consortium on a low copy number plasmid regulated by GPD promoter (table S2).

Among the eight CYP722C genes we examined, all the CYP722Cs from group I [S. lycopersicum, C. annuum, T. pratense, Glycine max (Soybean), Citrus sinensis (Sweet orange), and Vitis vinifera (Grape)] indeed converted CLA to OB (Fig. 4 and fig. S16A), while the two CYP722Cs from group II (F. vesca and L. japonicus) synthesized 5DS from CLA (Fig. 4 and fig. S16B). We also examined the function of OsCYP722B (Oryza sativa) and SbCYP722B (Sorghum bicolor) using the CLA-producing consortium and did not detect any conversion of CLA or synthesis of 5DS or OB (table S2 and fig. S14). The CYP722C characterization results are consistent with the previously reported SL profiles from the corresponding plants (2, 25, 26, 37, 41, 42) and suggest that sweet orange and grape are capable of synthesizing OB, although their SL profiles have not been reported.

The SL profile of most plants is not reported. The phylogenetic analyses and functional characterization of CYP722Cs from various plant species in the microbial system imply a sequence-function correlation, which can be used to predict the SL synthetic capacity of the corresponding plants.

### Metabolic engineering to improve 5DS production

After demonstrating microbial biosynthesis of various SLs, we used metabolic engineering to improve product titer, which should facilitate future gene characterization efforts. We chose 5DS as the model SL. We first improved the CL-producing E. coli (ECL-4). In this strain, the building blocks of CL (isopentenyl diphosphate and dimethylallyl diphosphate) were supplied by an endogenous, non-engineered methylenetetrahydrofolate (MTHF) pathway. Biosynthesis of many isoprenoids was found in prior studies to be limited by this pathway (44–47), possibly due to the fact that E. coli evolved to regulate this pathway to have low flux because of weak demand of isoprenoids by the cell. To up-regulate the MTHF pathway, we overexpressed its first enzyme, 1-deoxy-d-xylulose-5-phosphate synthase (EcDXS; coded by dxs). Since dxs is in the same operon in the E. coli genome with ispA (a gene also involved in the CL biosynthetic pathway), the dxs-ispA operon including its native promoter and terminator was inserted into pAC-BETAipi. IspA encodes farnesyl diphosphate synthase (EcEPPS) and would also be up-regulated in the new strain (ECL-6; table S2). When ECL-4 was replaced by ECL-6 in the

![Fig. 4. Phylogenetic analysis of CYP722C and the functional mapping.](image)
5DS-producing coculture (ECL4/YSL-5), the pool size of CL was increased by 150% (Fig. 5A), but the 5DS titer was not increased (Fig. 5B). Since the increase in the CL pool size could also be due to reduced consumption of CL, we analyzed a by-product (βI) of the CL biosynthetic pathway. Because there was no reaction known to consume βI in this culture, its titer can be used as an indicator of the CL production rate. The βI titer of the ECL-6/YSL-5 coculture was 330% (Fig. 5C) higher than that of the ECL-4/YSL-5 coculture, suggesting that the overexpression of dxs and ispA indeed improved the CL production in the coculture and that it is needed to improve the conversion of CL into 5DS in YSL-5.

Since the SL biosynthesis is less understood, it was more difficult to pinpoint the rate-limiting step. We then decided to increase the concentration of all the enzymes involved in functionalizing CL by enlarging the population of YSL-5. The yeast growth was limited by acetate supply (acetate concentration constantly below detection limit, 0.1 g/liter; during the coculture, acetate was the main carbon source of the yeast and was produced by E. coli), so ethanol was added as a supplement carbon source for S. cerevisiae. Ethanol can be converted into acetate in the yeast and cannot be metabolized by E. coli (48). The yeast growth was indeed improved by supplementing ethanol (4 g/liter; Fig. S17), and almost no CL was accumulated under this condition (Fig. 5A). 5DS titer was increased by 220% (to 47.3 μg/liter; Fig. 5B) compared with the coculture without the ethanol addition.

In addition, OsD27 in ESL-6 was replaced with six D27 variants from different plant species individually (Oryza nivara, Oryza rufipogon, Oryza meridionalis, Triticum aestivum, Zea mays, and Populus trichocarpa). Although none of these replacements further increased the 5DS production (fig. S18), these exercises quantitatively compared the activities of variants of SL biosynthetic enzymes from different plant species, which may shed light on evolution of SL biosynthesis (elaborated below).

**DISCUSSION**

The biosynthetic pathway from 3K to CL is conserved in most plant species, which may shed light on evolution of SL biosynthesis (elaborated below).

**Fig. 5.** Pathway engineering and process optimization improved 5DS titer. Information of all the microbial strains mentioned in the caption can be found in table S2.

The native ispA-dxs operon was introduced to ECL-4 (resulting strain: ECL-6) to overexpress E. coli EcDXS and EcFPPS. At 72 hours after starting the coculture, we measured 5DS titer of (A) CL, (B) 5DS, and (C) βI of ECL-4/YSL-5 coculture, ECL-6/YSL-5 coculture with and without ethanol feeding (ethanol was fed at 24 and 48 hours to increase ethanol concentration in the medium by 2 g/liter). All the cocultures were performed at 22°C. 5DS and βI were extracted and quantified by using authentic standards as described in Materials and Methods. Because of the lack of the authentic standard, the quantities of CL were estimated using raw data (the peak area from chromatogram of absorbance at 269 nm, RT: 31.72 min, and the unit of peak area is mAU*min and was defined as arbitral unit (a.u.)). The error bars represent the SE of three biological replicates.
shoot branching inhibition in plants and hyphal branching activity in arbuscular mycorrhizal fungi (AMF) (49). The limited access to natural SLs is hindering comprehensive investigations on the structure-activity correlation of this group of phytohormones in plant. The development of efficient microbial bioproduction of various natural SLs will advance these investigations and agricultural applications.

Much remains to be investigated into the biochemistry and evolution of CYP722Cs. Consistent with the pioneering in planta and biochemical characterizations of CYP722Cs (27, 28), our study also implies that CYP722Cs are divided into two major groups, one group converts CLA to OB (group I) and the other one synthesizes 5DS (group II). Both types of CYP722Cs are believed to synthesize OB or 5DS through first catalyzing 18-hydroxylation of CLA (27, 28). Group I CYP722C further catalyzes the oxidation of the alcohol at C18 to aldehyde that triggers nucleophilic attack to afford B ring and C ring closures for the synthesis of OB (the oxygen atom is retained at C18). This process would not produce 4DO (fig. S19) (27). On the other hand, group II CYP722C does not catalyze further oxidation of 18-hydroxy; it may protonate the 18-hydroxyl to serve as a leaving group to afford the synthesis of 5DS (fig. S19) (28). In the functional identification of various CYP722Cs, we did not detect the synthesis of 4DO in the CYP722C-expressing SL-producing microbial consortium. Further investigations into CYP722Cs from other plants may help answer whether there is a 4DO-synthesizing CYP722C. It is also interesting to understand how CYP722Cs evolve to control the C18 oxidation and stereospecificity of ring closing.

A compound detected in the 4DO-producing microbial consortium (ECL-4/YSL-2) may putatively be 18-hydroxy-CLA (fig. S10B). The synthesis of 4DO by OsCYP711A2 in ECL-4/YSL-2 is likely through a similar mechanism as GaCYP722C except that OsCYP711A2 catalyzes one more oxidation at C19 (to carboxylic acid) in addition to the C18 oxidation (to alcohol; fig. S19) (27). We did not detect any 5DS or OB in the 4DO-producing microbial consortium using OsCYP711A2 (ECL-4/YSL-2). Further biochemical investigations are needed to understand how CYP711As that convert CL to CLA were evolved to gain C18 hydroxylation ability and to catalyze the B ring and C ring closures with strict stereospecificity. The microbial platform we have built should be able to expedite enzyme characterization to answer these questions.

In addition, the establishment of deoxy-SL (4DO and 5DS) producing microbial consortium may provide a handy platform for the discovery of the downstream tailoring enzymes in SL biosynthesis. Hydroxylation on SLs can alter the biological activities. For example, Striga-susceptible maize cultivar only produces 5DS, while mainly sorgomol is detected in the resistant cultivar (60); Sorghum produces both 5DS and sorgomol and can convert 5DS to sorgomol (26). Recently, CYP728B35 encoded by Sorghum has been suggested to be the sorgomol synthase catalyzing the hydroxylation of 5DS to afford the synthesis of sorgomol (61). In addition to Sorghum, strigol-producing cotton converts 5DS to strigol and strigyl acetate (26). The enzymes involved in these oxidations of SLs remain to be characterized.

Moreover, the biosynthesis of noncanonical SLs remains largely unknown. In Arabidopsis, CLA can be methylated to form MeCLA by an unknown methyltransferase (11). Recently, lateral branching oxidoeductase, a 2-oxoglutarate- and Fe-dependent dioxygenase, was suggested to catalyze the hydroxylation of MeCLA to hydroxy-methylcarlactonoate (1’-HO-MeCLA) (62, 63), which may be the precursors of many noncanonical SLs such as avenol (37). In addition to MeCLA and 1’-HO-MeCLA, diverse noncanonical SLs, including hydroxyl CL (3-HO-CL, 4-HO-CL, and 16-HO-CL) and the corresponding hydroxyl CLA derivatives, were also detected in Arabidopsis (63), with little known about the enzymes involved in the formation of these structures. L. japonicus can produce noncanonical SL lotuslac-tone (15), and a 2-oxoglutarate–dependent dioxygenase (2-OGD) (named LLD) gene was proposed to be involved in the biosynthesis of lotuslac-tone (43), yet the catalytic function of LLD was not characterized. The CL/CLA-producing microbial consortium may also provide a convenient way to functionally identify and discover the enzymatic mechanism in the biosynthesis of various SL structures, both canonical and noncanonical.

Another intriguing yet mysterious question to be answered is the origin and evolutionary history of SL biosynthesis and perception. SLs have been reported to be present in several bryophytes and green algae (29, 30, 64), which do not always encode the full set of SL biosynthetic enzymes as described above (29, 30). These facts spur debate about the origin and evolution of SL biosynthesis and perception (29, 30). The moss Physcomitrella patens encodes one set of D27, CCD7, and CCD8, but no MAX1 analog, which is consistent with previous investigation that P. patens only synthesizes CL (29, 37). The activities of P. patens CCD7 (PpCCD7) and CCD8 (PpCCD8) toward the synthesis of CL have been validated through in vitro reconstitution (65). On the other hand, CCD8 and MAX1 analogs are absent from the genome of liverworts Marchantia polymorpha (29, 66). In this study, the activity of P. patens D27 (PpD27), PpCCD8, and M. polymorpha CCD7 (MpCCD7) were examined in the CL-producing ECL-6 through replacing the corresponding ortholog (e.g., PpD27 replacing OsD27; fig. S20). Replacing OsD27 with PpD27 retained the synthesis of CL, while replacing AtCCD7 or AtCCD8 with MpCCD7 or PpCCD8 completely abolished CL production (fig. S20). The inconsistency in the activity of PpCCD8 between our study and previous investigations might be due to different N terminus truncation or codon optimization method (we truncated four more amino acids to be consistent with trAtCCD8; table S6). MpCCD7 does not encode a chloroplast transfer peptide and is more than 10% longer in length than regular CCD7s, and the activity of MpCCD7 might be different from regular CCD7s and requires further investigations (e.g., assay with more substrates than 9C8C). Furthermore, since MAX1 and CYP722 analogs are universally present in flowering plants yet generally absent from primitive plants (29, 67), we also examined the activity of some MAX1 and CYP722 analogs from Sphagnum fallax (moss, SMAX1, 46.1% identity to AtMAX1; SfCYP722C, 36.2% identity to GaCYP722C) and Klebsormidium nitens (green algae, four MAX1 orthologs, KnMAX1a-1d, 27.9 to 31.2% identity to AtMAX1) in the CL-producing consortium (66). Unfortunately, none of these CYPs examined converted CL to CLA or 5DS or OB in the microbial consortia (fig. S21), suggesting different functions from AtMAX1 or group I or II CYP722Cs. These species may produce SL-like compounds that have not been found yet, with distinct biosynthetic pathway from seed plants, which need to be characterized. Comprehensive investigations into the putative SL biosynthetic enzymes from primitive plants are necessary to fully elucidate the SL biosynthesis from these plants as well as the origin and evolution of SL biosynthesis.

The current titer of 5DS (~50 μg/liter) needs to be further improved to isolate pathway intermediates of SL biosynthesis for structural verification (e.g., 18-hydroxy-CLA) and mechanistic investigation of SL biosynthetic enzymes (e.g., CYP722Cs). We engineered an
E. coli strain that could produce ATβC (~200 mg/liter) by using the recently developed isopentenol utilization pathway (68). But when we introduced the CL-synthesizing pathway into this strain, much less CL was produced compared with using the engineered MEP pathway. A future study should investigate how to efficiently convert ATβC into CL in high ATβC-producing strains. A special attention should be paid to D27, which is the first enzyme in this segment and may control its flux. D27s from different species could be screened, and the best natural D27 could be further improved by directed evolution.

Furthermore, although our study demonstrates that microbial consortium is a promising mechanism for the supply of various SLs, single strain–based SL-producing microorganism is still desired because of the easier manipulation during fermentation. To reconstruct CL-producing E. coli strains, further engineering on the functional reconstitution of multiple CYPs (i.e., CYP722Cs or CYP711As) in E. coli is critical. On the other hand, mechanistic investigation into the failed reconstitution of D27 and CCDs in yeast may enable the rational design of microenvironment to functionally reconstitute these plastid-localized enzymes in yeast.

As a conclusion, in this study, we have successfully reconstituted the biosynthesis of CL, CLA, and three canonical SLs (4DO, 5DS, and OB) in E. coli–yeast microbial consortia. Our study highlights the usefulness of the microbial platform as a convenient and rapid way to characterize the function of SL biosynthetic enzymes.

**MATERIALS AND METHODS**

**Chemicals and general culture conditions**

(±)-deoxy-strigol (purity >98%) and (±)-OB were purchased from Strigolab (Italy). (±)-4DO [also named as (±)-2′-epi-5DS] were acquired from ChemPep Inc. (USA). β is purchased from Thermo Fisher Scientific (USA). ATβC and 9CβC were purchased from Sigma-Aldrich Co. (USA). The chemically competent E. coli strain TOP10 (Life Technologies) was used for DNA manipulation and amplification and was grown at 37°C in lysogeny broth (LB) medium (Thermo Fisher Scientific) supplemented with appropriate amount of antibiotics [ampicillin (100 μg/ml; Thermo Fisher Scientific), kanamycin (50 μg/ml; Thermo Fisher Scientific), chloramphenicol (25 μg/ml; Thermo Fisher Scientific), and/or spectinomycin (50 μg/ml; Sigma–Aldrich)] for plasmid maintenance. For protein expression and CL production, we used chemically competent E. coli strain BL21(DE3) (Novagen) and LB (or XY medium). XY medium contains KH2PO4 (13.3 g/liter), (NH4)2HPO4 (4 g/liter), citric acid (1.7 g/liter), CoCl2 (0.0025 g/liter), MnCl2 (0.015 g/liter), CuCl2 (0.0015 g/liter), H3BO3 (0.003 g/liter), Na2MoO4 (0.0025 g/liter), Zn(CH3COO)2 (0.008 g/liter), Fe(III) citrate (0.06 g/liter), thiamine (0.0045 g/liter), MgSO4 (1.3 g/liter), yeast extract (5 g/liter), and xylene (40 g/liter), pH 7.0. In the first stage of the coulture fermentation, yeast strains were cultured at 28°C in complex yeast extract, peptone, and dextrose (YPD; all components from BD Biosciences) medium or synthetic dropout (SD) medium containing yeast nitrogen base (1.7 g/liter) without amino acids (BD Biosciences), ammonium sulfate (5 g/liter; Thermo Fisher Scientific), dextrose (20 g/liter), and suitable SD mixture at the concentrations specified by the manufacturer (Clontech). XY medium was used in the second stage of the coculture fermentation. Unless specified, all the chemicals used in this study were purchased from Thermo Fisher Scientific or Sigma–Aldrich Co.

**General techniques for DNA manipulation**

Plasmid DNA was prepared using the EconoSpin columns (Epoch Life Science) according to the manufacturer’s protocols. Polymerase chain reactions (PCRs) were performed using Q5 DNA polymerase [New England Biolabs (NEB)] and Expand High Fidelity PCR System (Roche Life Science) according to the manufacturer’s protocols. PCR products were purified by the Zymoclean Gel DNA Recovery Kit (Zymo Research). All DNA constructs were confirmed through DNA sequencing by Source BioScience (LA, USA) or BioBasic (Singapore). Restriction enzymes (NEB) and T4 ligase (NEB) were used to produce and ligate the DNA fragments, respectively. BP Clonase II Enzyme Mix, Gateway pDONR221 Vector and LR Clonase II Enzyme Mix (Invitrogen), and the S. cerevisiae Advanced Gateway Destination Vector Kit (Addgene) were used to perform Gateway cloning (69). Using this method, the yeast expression cassette vectors were constructed. Plasmids were introduced into yeast cells using the Frozen-EZ Yeast Transformation II Kit (Zymo Research). Gibson one-pot, isothermal DNA assembly was conducted at 10-μl scale by incubating T5 exonuclease (NEB), Phusion polymerase (NEB), Taq ligase (NEB), and 50 ng of each DNA fragment at 50°C for 1 hour to assemble multiple DNA fragments into one circular plasmid (70). Integrated yeast strains were constructed through homologous recombination and DNA assembly (71). Plasmids and microbial strains used in this study are listed in tables S1 and S2, respectively. DNA oligonucleotides were synthesized by Integrated DNA Technologies (IDT) and Life Technologies. The plant gene sequences were codon-optimized for expression in S. cerevisiae and synthesized by IDT (Corvalle, IA) and Twist Bioscience (San Francisco, CA). DNA sequences of genes involved in this work are listed in table S6.

For the construction of E. coli expression vectors, the truncated AtCCD7 gene was amplified by PCR and cloned into the pCDFDuet-1 plasmid (Novagen) using Nco I and Not I to yield the plasmid pCDFDuet-trAtCCD7. The OsD27 gene was amplified by PCR, digested by Nde I and Avr II, and ligated into accordingly digested pCDFDuet-trAtCCD7 yielding the plasmid pCDFDuet-trAtCCD7-OsD27. Using the same strategy, the other six D27 homologous genes were cloned into the pCDFDuet-trAtCCD7 vector, respectively, yielding the corresponding coexpression plasmid. The OsD27, truncated OsD27, and AtD27 genes were PCR-amplified, digested by Nde I and Avr II, and ligated into accordingly digested pCDFDuet-1, yielding the plasmid pCDFDuet-OsD27, pCDFDuet-trOsD27, and pCDFDuet-AtD27. The truncated AtCCD8 gene was amplified by PCR and cloned into pET21a using Gibson assembly, yielding pET21a-trAtCCD8. The truncated PpCCD8 was cloned into pET21a, yielding pET21a-trPpCCD8. For the construction of yeast expression cassettes, NADPH-P450 reductase and each individual P450 gene were constructed using Gateway cloning as described previously. The plasmid of pAC-BETAipi-ispA/dxs was constructed by using a Gibson assembly kit (Gibson Assembly Master Mix, NEB). The insert was amplified from genomic DNA of E. coli BL21(DE3) by using colony PCR and assembled with the backbone of pAC-BETAipi generated by using PCR.

**Culture conditions for E. coli–based CL precursors production**

For the in vivo production of 9CβC, E. coli BL21(DE3) was transformed with pAC-BETAipi (Addgene) and pCDFDuet-OsD27, generating E. coli ECL-2. For 9CβA10COL production, E. coli
BL21(DE3) was transformed with pAC-BETAipi (Addgene) and pCDFDuetoatsD27-trAtCCD7 to generate E. coli ECL-3. Then, the yellow colonies were picked and grown in LB with chloramphenicol (25 μg/ml) and spectinomycin (50 μg/ml) at 37°C, overnight. Fifty microliters of the overnight culture was then used to inoculate 5 ml of fresh LB with the corresponding antibiotics with a starting optical density at 600 nm (OD600) ~0.05 and cultured at 37°C and 220 rpm in the 100-ml Erlenmeyer flask. When OD600 reached ~0.6, IPTG was added to make the final concentration at 0.2 mM, with ferrous sulfate supplemented at the same time (final concentration at 10 μM/liter). Then, the cultures were cultivated at 22°C and 220 rpm for 72 hours.

**Culture conditions for E. coli–yeast consortium–based SL production**

For the in vivo production of SLs, E. coli BL21(DE3) was cotransformed with the plasmids pAC-BETAipi (Addgene), pCDFDuetoatsD27-trAtCCD7, and pET21α-trAtCCD8, generating E. coli ECL-4. Single yellow colony was then picked and grown overnight at 37°C in 1 ml of LB supplemented with ampicillin (100 μg/ml), chloramphenicol (25 μg/ml), and spectinomycin (50 μg/ml). Fifty microliters of the overnight culture was then used to inoculate 5 ml of fresh LB with the corresponding antibiotics with a starting OD600 at ~0.05 and cultured at 37°C and 220 rpm in the 100-ml Erlenmeyer flask. When OD600 reached ~0.6, IPTG was added with the final concentration at 0.2 mM, with ferrous sulfate supplemented at the same time (final concentration at 10 μM/liter). Then, the cultures were incubated at 22°C and 220 rpm for 15 hours.

In parallel to preparing the E. coli culture, single colony of each yeast strain harboring the corresponding cytochrome P450-expression constructs was used to inoculate an appropriate SD medium considering the auxotrophic markers for maintaining the plasmid(s). The seed culture was incubated at 28°C and 220 rpm overnight. One hundred microliters of the overnight grown seed culture was used to inoculate 5 ml of the corresponding SD medium in a 100-ml Erlenmeyer flask and grown at 28°C for 15 hours.

The E. coli and yeast cells prepared as described above were harvested by centrifugation at 3500 rpm for 5 min. Then, the E. coli and S. cerevisiae cells were mixed and resuspended in 5 ml of XY medium (OD600 ~ 8.0) and cultured in 100-ml shake flask at 22°C and 220 rpm for 60 or 72 hours (final OD600 ~ 40). In the case of CL production, the parent S. cerevisiae strain (CEN.PK2-1D) was pre-cultivated in YPD and mixed with the CL-producing E. coli cells.

**Isolation and characterization of SLs and their precursors**

Unless specified, 5 ml of culture was used for compound extraction. For the extraction of intracellular and extracellular metabolites, 5 ml of cell culture was transferred into 50-ml centrifuge tube, and the cells and medium were separated by centrifugation at 5000 rpm for 10 min.

The cell pellets were transferred to a new 2-ml microcentrifuge tube and resuspended in 150 μl of dimethylformamide and shaken vigorously, followed by the addition of 850 μl of acetone and vigorous shaking for 15 min by using a vortex mixer (Thermo Fisher Scientific) and centrifugation at 12,000 rpm for 10 min. Then, the supernatant was transferred to a new 1.7-ml microcentrifuge tube and dried in a vacuum concentrator (Eppendorf Vacufuge plus) at 30°C for 2 to 3 hours. The dried sample was dissolved in 100 μl of acetone. The medium was transferred into a 50-ml centrifuge tube containing 4 ml of ethyl acetate. The mixture was vortexed vigorously for 5 min by using a vortex mixer (Genie) and then centrifuged at 4000 rpm for 20 min. The upper ethyl acetate layer of extracted medium was transferred into a 1.7-ml microcentrifuge tube and evaporated to dryness by using a vacuum concentrator (Eppendorf Vacufuge plus) at 30°C for 30 min. The dried extract was dissolved in 100 μl of acetone.

The samples were centrifuged at 12,000 rpm for 10 min before applying to high-performance LC (HPLC) analysis. Both UV-VIS and MS detectors were used. The used instrument was Shimadzu LC-MS 2020 (Kyoto, Japan) or a Waters UPLC (ACQUITY) coupled with a Bruker quadrupole time-of-flight MS (Q-TOF MS; micrOTOF II).

ATβC and 9CβC were analyzed based on Separation Method I and a C18 column (Kinetic C18, 100 mm by 2.1 mm, 100 Å, particle size of 2.6 μm; Phenomenex, Torrance, CA, USA). In Separation Method I, the column temperature was 25°C, single mobile phase was 0.1% (v/v) formic acid in methanol, the isotropic elution was 0.4 ml/min, and the analysis time was 20 min. The injection volume was 10 μl, and the UV-VIS absorption was monitored in the range of 190 to 800 nm. With Separation Method I, the retention time of 9CβC was 7.78 min and that of ATβC was 8.17 min (characteristic absorption wavelengths: 447 and 471 nm for both carotenes).

9CBA10′COL was analyzed using Separation Method II and a C14 column (Poroshell 120 EC-C18, 100 mm by 3.0 mm, 100 Å, particle size 2.7 μm; Agilent, Santa Clara, CA, USA). In Separation Method II, the column temperature was 40°C, the mobile phase A was 0.1% (v/v) formic acid in water, the mobile phase B was 0.1% (v/v) formic acid in methanol, and the gradient elution was 0.5 ml/min. The gradient was as follows: 0 to 18 min, 5 to 100% B; 18 to 43 min, 100% B; and 43 to 45 min, 100 to 5% B. The injection volume was 10 μl, and the UV-VIS absorption was monitored in the range of 190 to 800 nm. With Separation Method II, the retention time of 9CBA10′COL was 12.64 min (characteristic absorption wavelengths: 373 and 390 nm).

All the SLs (CL, CLA, putative 18-hydroxy-CLA, 4DO, 5DS, and OB) were analyzed using Separation Method III and a C18 column (Kinetic C18, 100 mm by 2.1 mm, 100 Å, particle size 2.6 μm; Phenomenex, Torrance, CA, USA). In Separation Method III, the column temperature was 40°C, the mobile phase A was 0.1% (v/v) formic acid in water, the mobile phase B was 0.1% (v/v) formic acid in methanol, and the gradient elution was 0.4 ml/min. The gradient was as follows: 0 to 28 min, 5 to 100% B; 28 to 35 min, 100% B, and 35 to 40 min, 5% B. The injection volume was 10 μl, and the UV-VIS absorption was monitored in the range of 190 to 800 nm. With Separation Method III, the retention time and the characteristic absorption wavelength of various analytes are listed as follows: βI, 15.91 min (298 nm); CL, 20.58 min (269 nm); CLA, 16.32 min (271 nm); 18-hydroxy-CLA, 12.16 min; 4DO, 15.81 min; OB, 11.03 min; and 5DS, 15.97 min. The compounds without the wavelength information were detected using an MS detector, which operates in the m/z range of 50 to 800 in the positive or negative ion modes. Electrospray ionization (ESI) was used. The desolvation line temperature was 250°C. The nebulizing gas and drying gas flow rates were 1.5 and 15 liter/min, respectively.

The HR-MS analysis of CLA was performed by using an Ultra Performance LC (UPLC, Waters ACQUITY) linked with a TOF MS (Bruker micrOTOF II). The separation was based on Separation...
Method IV and a C18 column (Poroshell 120 EC-C18 column, 2.1 mm by 50 mm, particle size of 2.7 μm; Agilent Technologies). In Separation Method IV, the column temperature was 40°C, the mobile phase A was 0.1% (v/v) formic acid in water, the mobile phase B was 0.1% (v/v) formic acid in acetonitrile, and gradient elution was 0.3 ml/min. The gradient was as follows: 0 to 15 min, 20 to 100% B; 15 to 20 min, 100 to 20% B; and 20 to 22 min, 20% B. The MS/MS analysis was performed in product ion scan mode based on ESI (negative mode). The optimized MS/MS conditions were as follows: capillary voltage, 3500 V; end plate offset, 500 V; desolvation gas flow rate (N₂), 4.0 liter/min; drying gas temperature, 200°C; hexapole RF, 50 Vpp; ion energy, 4 eV; collision energy, 18 eV; and precursor ion m/z, 331.2. The scan range of m/z was from 50 to 1300. Sodium formate (10 mM) solution was used to calibrate the MS before every use. HR-MS analysis of 18-hydroxy-CLA was performed on a Synapt G2-Si Q-TOF MS (Waters) coupled to an i-UPPLC system (Waters). The separation was conducted using Separation Method III and the C18 column (Kinetex C18, 100 mm by 2.1 mm, 100 Å, particle size 2.6 μm; Phenomenex, Torrance, CA, USA) as mentioned above. The injection volume was 5 μl. The MS were obtained using the negative ion mode, the scan range of m/z was from 50 to 1200 with a 0.2 s scan time. MS/MS was acquired in a continuum data format and data-dependent fashion with collision energy 25 eV. Source and desolvation temperatures were 150°C and 600°C, respectively. Desolvation gas was set to 1100 liter/hour, and cone gas was set to 150 liter/hour. All gases were nitrogen except the collision gas, which was argon. Capillary voltage was 1.5 kV in negative ion mode.

In the experiment aiming to improve 5DS biosynthesis, ββ was analyzed without drying to avoid evaporation loss. A total of 0.5 ml of cell culture was transferred into a 2-ml Eppendorf Safe-Lock Tubes containing 0.1 g of glass beads (Sigma-Aldrich, G8772) and 0.5 ml of ethyl acetate. The mixture was incubated at 25°C and 1500 rpm for 1 hour by using a Thermomixer (Eppendorf) and then centrifuged at 14,000 rpm for 10 min. One hundred microliters of the upper organic phase was used for gas chromatography–MS (5977B GC/MSD, Agilent Technologies) analysis. Five microliters of the filtered sample was injected in a spitless mode. HP-5MS capillary column (30 m by 0.25 mm, 0.25-μm film thickness; Agilent Technologies) was used, with helium as the carrier gas at the flow rate of 1 ml/min. The following oven temperature program was carried out: 50°C for 1 min, 50°C to 100°C at a rate of 5°C/min, 100°C to 300°C at a rate of 50°C/min, and 300°C for 1 min. Commercially available ββ (Sigma-Aldrich, N12603) was used to prepare standard solutions. The retention time was 11.02 min.

In the experiment aiming to improve 5DS biosynthesis, 5DS was quantified by using the same procedure as the HR-MS experiment, except the used mode was scan instead of product ion scan. No collision energy was applied. The quantification was based on extracted ion chromatogram (m/z: 331.20 ± 0.01). The retention time was 7.95 min.

In the experiment aiming to improve 5DS biosynthesis, the quantification of ATβC and 9CβC was done by using Separation Method V and a C30 column (YMC Carotenoid, 250 mm by 4.6 mm, 5 μm). In Separation Method V, the column temperature was 30°C; the single mobile phase was 75% (v/v) ethanol, 20% (v/v) methanol, and 5% (v/v) tetrahydrofuran; and the isocratic elution was 1 ml/min for 20 min. The injection volume was 10 μl, and the used detector was a UV–VIS detector (475 nm). The model of the HPLC was Agilent 1260 Infinity. The retention time of ATβC and 9CβC was 12.26 and 13.98 min, respectively.

In the experiment aiming to improve 5DS biosynthesis, the quantification of CL was done using Separation Method III on Agilent 1260 Infinity HPLC with a UV detector. The column was Poroshell 120 EC-C18 (150 mm by 4.6 mm, 100 Å, particle size of 4 μm; Agilent Technologies). The injection volume was 10 μl. The UV detector with a wavelength of 269 nm was used. The retention time of CL was 31.72 min.

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at https://science.org/doi/10.1126/sciadv.abh4048

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