A Novel Soluble Squalene-Hopene Cyclase and Its Application in Efficient Synthesis of Hopene

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Hopene is a important precursor for synthesizing bioactive hopanoids with great commercial value. However, the chemical methods for synthesizing hopene are not efficient to date. Hopene is commonly obtained by extracting from plants or bacteria like other terpenoids, but the complicated extraction process is inefficient and unfriendly to the environment. Hopene can be biological synthesized by squalene-hopene cyclase (SHC) from squalene. However, hopene production by SHC remained at a low level until now. In this work, we found a novel SHC named OUC-SaSHC from Streptomyces albolongus ATCC 27414. An easy procedure for expression and purification of OUC-SaSHC was established. The conditions for OUC-SaSHC to convert squalene into hopene are optimized as in 100 mM sodium phosphate buffer (pH 7.0) containing 0.5% Tween 80, 20 mM squalene and 0.14 mg/mL OUC-SaSHC at 30°C. In the scale-up reaction with the final volume of 100 mL, the yield of squalene could be up to 99% at 36 h, and 8.07 mg/mL hopene was produced. Our work showed a great potential of OUC-SaSHC as biocatalyst on scale-up production of hopene, hence improves the SHC-catalyzing enzyme synthesis of hopene from laboratory level to application level.

Keywords: squalene-hopene cyclase, Streptomyces albolongus, squalene, hopene, biocatalysis

INTRODUCTION

Hopanoids are a class of pentacyclic triterpenoids which composed of hopene skeleton and a series of different side chains (Kannenberg and Poralla, 1999; Schmerk et al., 2011). These natural products play an important role in stabilizing the structure of the bacterial membrane just like the effect of sterols in eukaryote (Sáenz et al., 2012; Doughty et al., 2014). Many hopanoids have been reported to display physiological activity. For example, diploterol had been shown to be toxic to mouse leukemia cells, and the bacteriohopane-32, 33, 34, 35-tetrol has been reported to be a valuable and powerful anti-inflammatory and anti-oxidation natural product (Chen et al., 1995; Moreau and Hicks, 2001). To the best of our knowledge, hopene is the common precursor in the pathways of all known hopanoids (Bradley et al., 2010), which means hopene is an important precursor for synthesizing bioactive hopanoids (Schmerk et al., 2015; Sohlenkamp and Geiger, 2016). Compared with the chemical methods to synthesize hopene, the enzyme in multiple bond-forming reaction had a high selectivity for the overall transformation (Yoder and Johnston, 2006). Besides that, due to the low content of hopene in the natural accumulating plants or microbes (Stroeve et al., 1998), the process of extraction and purification would be complicated,
not to mention the waste of resources and environment pollution. Therefore, enzymatic catalysis is considered to be a not only mild but also environmental friendly method for efficiently synthesizing hopene.

In the biosynthetic pathway of hopene, the squalene-hopene cyclase (SHC) could catalyze the formation of hopene from its precursor squalene (Nakano et al., 2019; Figure 1). SHC is a famous enzyme for several reasons: first, the reaction of converting squalene to hopene catalyzed by SHC is the most complex already known natural one-step biochemical reaction, in which five rings and nine stereocenters were formed and thirteen covalent bond were changed (Yoder and Johnston, 2006; Abe, 2007). Second, SHCs contain QW-sequence repeats which are unique to the known triterpene cyclases (Frickey and Kannenberg, 2009). Third, all reported SHCs belong to a rare kind of membrane protein classified as “monotopic,” for SHC submerged from one side into the non-polar part of the phospholipid bilayer without protruding through it (Wendt et al., 1999). Hence, SHC and the reaction it catalyzed are investigated by many biochemists (Kannenberg and Poralla, 1999; Giner et al., 2005; Gustafsson et al., 2017; Kühnel et al., 2017; Fukuda et al., 2018; Takahashi et al., 2018; Nakano et al., 2019).

To date, several SHCs have been characterized. As early as 1986, the first prokaryotic SHC was purified from the thermophilic bacterium *Alicyclobacillus acidocaldarius* and its activity was tested in vitro (Seckler and Poralla, 1986). Following that, characterization and heterologous expression of it was also carried out (Ochs et al., 1990, 1992). Later, SHC from *A. acidocaldarius* became the enzyme model of this field, the structure and reaction mechanism of it have been revealed (Wendt et al., 1997; Hoshino et al., 2004; Kühnel et al., 2017; Ideno et al., 2018). In the work on SHC from *Zymomonas mobilis*, the substrate specificity of SHC was investigated. It was concluded that SHC can require many terpenoids as substrates (Seitz et al., 2012, 2013). For the SHC from *Streptomyces peucetius*, the optimal conditions of converting squalene to hopene is at pH 7.0 and 35°C (Ghimire et al., 2009). Other SHCs from *Pseudomonas mendocina*, *Bradyrhizobium japonicum*, *Methyllococcus capsulatus*, *Rhodopseudomonas palustris*, and *Tetrahymena thermophila* have also been reported (Saa et al., 1991; Kleemann et al., 1994; Perzl et al., 1997; Tippelt et al., 1998; Nair and Kochupurakal, 2019).

In this work, we found a novel soluble SHC from *Streptomyces albolongus*. After heterologous expression in *Escherichia coli*, high activity of this SHC to convert squalene into hopene was detected. Furthermore, the soluble feature of our SHC enables simple Ni²⁺-NTA purification, so enough enzymes could be obtained for scale-up reaction. Our work makes the large scale production of hopene from squalene feasible.

**MATERIALS AND METHODS**

**Bacterial Strains and Culture Conditions**

*S. albolongus* ATCC 27414 was cultivated at 28°C, pH 7.3 with shaken at 180 rpm in 50 mL medium which consisted of 1% (w/v) soluble starch, 0.2% yeast extract. *E. coli* strains were cultivated at 37°C, 220 rpm in LB medium [1% (w/v) tryptone, 0.5% (w/v) yeast extract and 1% (w/v) NaCl] or using ZYP-5052 medium [1% (w/v) tryptone, 0.5% (w/v) yeast extract] containing 100 µg/mL ampicillin and 34 µg/mL chloramphenicol (Solarbio, China) in 20°C (Liang et al., 2017).

**Sequence Analysis**

Clustal × program 25 was used for multiple sequence alignment analysis and using ESPript 3.0 to compare results (Larkin et al., 2007). Neighbor-joining phylogenetic tree was constructed using the molecular evolutionary genetics analysis (MEGA) 6.0 software (Frickey and Kannenberg, 2009).

**Cloning, Heterologous Expression, and Purification of Recombinant OUC-SaSHC**

A TIANamp Bacteria DNA Kit (Tiangen Biotech, Beijing, China) was used to extract the genomic DNA, which was used as a template to amplify the nucleotide sequence of *sashc* with primers (F-GGAAGATCTGTGACAGCAACCGGAC, R-CTAGCTAGCTGCTCCCGCTCCTCGTCAG) and restriction sites is Nhel and BglII. After designing and synthesizing of the primers, the gene was amplified and connected with vectors (pET-His) using Vazyme One Step Clone Kit (Nanjing, China). After transformation of vector into *E. coli* BL21(DE3) pLySs, expression of OUC-SaSHC was carried out by cultivating the strain in the ZYP-5052 medium (ZYP-5052) at 20°C, 220 rpm for 48 h. 1 mL inoculum (2%) was inoculated into 50 mL medium contained in the 250 mL Erlenmeyer flask. After culturing, cells were collected by centrifugation with 50 mL centrifuge tube at 8000 × g for 15 min at 4°C, and then resuspended in saline to wash. After the second centrifugation, the cells from 50 mL fermentation broth were resuspended in 10 mL buffer (20 mM Tris-HCl, pH 8.0) for disruption (amplitude 40%, 300 W, on 3 s, off 3 s for 30 min). After disrupted by sonication, the supernatant was collected after centrifugation with 50 mL centrifuge tube at 10000 × g for 20 min at 4°C. 50 mL supernatant was loaded onto the Ni²⁺-NTA (5 mL, Qiagen, R-CTAGCTAGCTGCTCCCGCTCCTCGTCAG) and restriction sites is Nhel and BglII. After designating and synthesizing of the primers, the gene was amplified and connected with vectors (pET-His) using Vazyme One Step Clone Kit (Nanjing, China). After transformation of vector into *E. coli* BL21(DE3) pLySs, expression of OUC-SaSHC was carried out by cultivating the strain in the ZYP-5052 medium (ZYP-5052) at 20°C, 220 rpm for 48 h. 1 mL inoculum (2%) was inoculated into 50 mL medium contained in the 250 mL Erlenmeyer flask. After culturing, cells were collected by centrifugation with 50 mL centrifuge tube at 8000 × g for 15 min at 4°C, and then resuspended in saline to wash. After the second centrifugation, the cells from 50 mL fermentation broth were resuspended in 10 mL buffer (20 mM Tris-HCl, pH 8.0) for disruption (amplitude 40%, 300 W, on 3 s, off 3 s for 30 min). After disrupted by sonication, the supernatant was collected after centrifugation with 50 mL centrifuge tube at 10000 × g for 20 min at 4°C. 50 mL supernatant was loaded onto the Ni²⁺-NTA (5 mL, Qiagen, Hilden, Germany) to purify OUC-SaSHC. Tris-HCl buffer (pH 8.0, 100 mM) containing different concentrations of imidazole (20–500 mM) have been used for protein elution. Different aliquots were determined by SDS-PAGE. Protein concentration was detected by Bradford method using coomassie brilliant blue. After purification, OUC-SaSHC was stored at −80°C until further use.

**Cyclization Assay**

The crude extracts or purified OUC-SaSHC was lyophilized by vacuum freezing, then the enzyme powder was used for cyclization assay. The cyclization reaction was started by adding 0.08 mg enzyme powder into 2 mL 100 mM citrate buffer (pH 4.5) containing squalene whose final concentration is 20 mM, fill nitrogen gas to avoid substrate oxidation, then at 20°C for 16 h (Siedenburg et al., 2012).
Product Extraction and Detection
After the catalyzed reaction was stopped (interrupted), the reaction mixture was extracted with four-fold volume of hexane. The organic phase was collected by centrifugation at 8000 \( \times \) g, then dried by \( \text{N}_2 \) and resolved in 300 \( \mu \text{L} \) of isooctane. After that, the samples were analyzed by gas chromatography (GC).

Squalene and hopene were assayed qualitatively and quantitatively by using an Agilent 6890A GC apparatus equipped with a HP-5 column (30 m \( \times \) 0.25 mm \( \times \) 0.25 \( \mu \)m) and a flame ionization detector. Two microliters of the solvent extract was applied onto the column (splitless, helium flow rate: 0.5 mL/min). The injector temperature was 250\(^\circ\)C. The oven temperature was initially 60\(^\circ\)C, kept constant for 3 min,
temperature for 16 h. The mixture, then carrying out the reaction at the optimal pH and 2
10 mM Na	2
 Tween 20, Tween 60, Tween 80, or Triton X-100, 1 or activity of OUC-SaSHC was determined by adding 0.5% (w/v)
buffer (pH 9.0-10.0).

Characterization of OUC-SaSHC

The optimal temperature was determined by conducting the reaction at 20, 30, 40, 50, and 60°C in 100 mM citrate buffer (pH 4.5) with 10 mM squalene as substrate by adding 0.08 mg enzyme powder for 16 h. The optimal pH of OUC-SaSHC was determined by carrying out the reaction at the optimal temperature for 16 h in different buffers, including citrate buffer (pH 4.0-6.0), sodium phosphate buffer (pH 6.0-8.0), Tris-HCl buffer (pH 8.0-9.0) and Na	2
CO	3-NaHCO	3
buffer (pH 9.0-10.0).

The effect of detergents, EDTA and metal ions on the activity of OUC-SaSHC was determined by adding 0.5% (w/v) Tween 20, Tween 60, Tween 80, or Triton X-100, 1 or 10 mM Na	2
-EDTA, CoCl	2
, KCl, NaCl, FeSO	4
, CuCl	2
, MnCl	2
, CaCl	2
, MgCl	2
, ZnCl	2
, NiCl	2
, or BaCl	2 into the reaction mixture, then carrying out the reaction at the optimal pH and temperature for 16 h.

Catalytic Conversion Reaction

The optimal reaction time was determined according to the productivity of hopene formation from squalene catalyzed by 0.002, 0.004, 0.006, 0.008, or 0.01 mg/mL of OUC-SaSHC in 100 mM sodium phosphate buffer (pH 7.0) containing 0.5% Tween 80, 3 mM squalene at 30°C. The optimal substrate concentration was determined by carrying out the reaction using 0.08 mg/mL OUC-SaSHC at pH 7.0, 30°C for 36 h with the substrate concentration of 0.5, 1, 5, 10, 20, 25, 30, 40, 50 mM. The OUC-SaSHC concentration was optimized according to the conversion ratio of squalene in the reaction containing 20 mM squalene at pH 7.0, 30°C for 36 h by adding OUC-SaSHC with the final concentration of 0.14 mg/mL.

The scale-up experiment was conducted at the optimal conditions: 100 mM sodium phosphate buffer (pH 7.0) containing 0.5% Tween 80, 20 mM substrate, 0.14 mg/mL OUC-SaSHC at 30°C with the reaction mixture volume of 100 mL.

RESULTS AND DISCUSSION

Heterologous Expression and Identification of a Novel SHC

In our work on cultivation of S. albolongus ATCC 27414, it was found that both squalene and hopene were accumulated in the strain cells after fermentation (Data not shown). From the whole genome sequence of S. albolongus, one putative SHC-coding gene designated as sashc was found based on annotation and Basic Local Alignment Search Tool (BLAST). Gene sashc contains 2076 bp nucleotides and encodes putative SHC named OUC-SaSHC (GenBank accession No. MH121056) with 691 amino acids. In addition, the phylogenetic analysis result shows that OUC-SaSHC is a member of ISOPREN-C2-like super family (Figure 2A) which contains squalene-hopene cyclase and 2,3-oxidosqualene cyclase. These enzymes catalyze a cationic cyclization cascade converting linear triterpenes to fused ring compounds (Rohmer et al., 1979). Moreover, multiple sequence alignment revealed that OUC-SaSHC had one DXDDTA motif and seven QW motifs [(K/R) (G/A)X	3
−(F/Y/W) (I/I/V)X	3
QX	5
GXW] (Figure 2B). The function of DXDDTA motif is initiation of polycyclization reaction (Wendt, 2005) and the QW motif is necessary for reinforcement of the protein structure (Frickey and Kannenberg, 2009). These results indicated that OUC-SaSHC was a new member of ISOPREN-C2-like superfamily.

After heterologous expression of sashc in E. coli BL21(DE3) pLY5s, the crude extracts of the expressing sashc could convert squalene to hopene (data not shown), which was verified by both GC and GC-MS with the hopene standard sample. This showed that we found a novel SHC from the actinomycete S. albolongus.

OUC-SaSHC Is a Soluble Protein

Ni	2
+-NTA sepharose column was used to purify OUC-SaSHC with the C-terminal 6 × His tag from the crude extracts, the SDS-PAGE result showed a single belt with the relative

**FIGURE 3 |** SDS-PAGE analysis of OUC-SaSHC. Lanes: M, protein marker; 1, supernatant of OUC-SaSHC cell lysate; 2, pellet of OUC-SaSHC cell lysate; 3, purified OUC-SaSHC.
molecular mass of around 63–75 kDa, which was consistent with the theoretical value of 69 kDa (Figure 3). The purified protein was used to catalyze the cyclization of squalene. The specific activity of OUC-SaSHC was tested as 1138 U/mg, and the product was identified as hopene by GC-MS, since both the retention time (Figure 4A) and the most indicative fragments of the mass spectrum [410.4 (M⁺, 21%), 367.3 (M⁺–C; H, 9%), 341 (below 5%), 191-2 (ring C cleavage, 100%), 189.2 (ring C cleavage, 79%), m/z (%)] (Figure 4C) were consistent with the hopene standard sample, which suggests the product was indeed hopene (Figure 4B).

**Characterization of OUC-SaSHC**

To find out the conditions for OUC-SaSHC exhibiting high activity, effects of temperature, pH, detergents and metal ions on the activity of OUC-SaSHC were investigated. As shown in Figure 5A, OUC-SaSHC showed highest activity at 30°C, and maintained approximately 75% of the highest activity at
FIGURE 5 | Characterization of OUC-SaSHC. (A) Effect of temperature on OUC-SaSHC activity. (B) Effect of pH on OUC-SaSHC activity. (C) Effect of different kinds of detergents (0.5%, w/v) on OUC-SaSHC activity. The reaction without detergents was control which activity was defined as 100%.
TABLE 1 | Effect of metal ions on OUC-SaSHC activity.

| Ion      | Relative activity (%) |
|----------|-----------------------|
| Control  | 100.00 ± 1.2          |
| Co²⁺     | 103.26 ± 5.28         |
| K⁺       | 125.31 ± 0.79         |
| Na⁺      | 92.76 ± 2.26          |
| Fe²⁺     | 102.35 ± 3.78         |
| Cu²⁺     | 92.17 ± 5.85          |
| Mn²⁺     | 153.21 ± 4.53         |
| Ca²⁺     | 103.57 ± 3.27         |
| Mg²⁺     | 101.76 ± 5.23         |
| Zn²⁺     | 57.69 ± 0.69          |
| Ni²⁺     | 85.25 ± 2.72          |
| Na₂-EDTA | 94.17 ± 3.67          |

40°C. When the temperature was over 50°C or below 20°C, the activity of OUC-SaSHC decreased rapidly. The optimal pH for OUC-SaSHC is 7.0. At pH 6.0, OUC-SaSHC exhibited 82% of its highest activity, while in the case of pH > 8.0 or pH < 5.0, the activity of OUC-SaSHC was less than 40% of its optimal value. This indicated that OUC-SaSHC could only exhibit high activity at neutral conditions (Figure 5B). The optimal reaction temperature for SHC from S. peucetius and A. acidocaldarius has been reported as 35 and 60°C, respectively. Similar with OUC-SaSHC, their optimal reaction conditions are in accordance with the optimum growth temperature of their original microorganisms (Ghimire et al., 2009; Siedenburg and Jendrossek, 2011).

Squalene is not only insoluble in water but also easily oxidized (Siedenburg et al., 2012). On the contrary, OUC-SaSHC was hydrophilic since it was a cytoplasmic protein. In order to promote the contact of OUC-SaSHC with squalene, 0.5% detergent was added to the reaction mixture. As shown in Figure 5C, three detergents, Tween 20, Tween 80, and Triton X-100, could increase the activity of OUC-SaSHC. When using Tween 80, the activity was increased by 29.72%.

Effect of metal ions on the activity of OUC-SaSHC is shown in Table 1. When adding EDTA, OUC-SaSHC still remained 75.26% of its normal activity, which means metal ions are not necessary for OUC-SaSHC. Many metal ions, such as Co²⁺, Na⁺, Fe²⁺, Mn²⁺, Ca²⁺, and Mg²⁺, had no obvious effects on the activity of OUC-SaSHC. Only 3 metal ions, Cu²⁺, Zn²⁺, and Ni²⁺ could inhibit OUC-SaSHC obviously. K⁺, Mn²⁺, and Ba²⁺ have a significant role in promoting OUC-SaSHC’s activity. Especially, 1 mM Mn²⁺ could increase the activity of OUC-SaSHC by 53.21%. Therefore, 1 mM Mn²⁺ was added in the reaction mixture in the application of OUC-SaSHC for hopene production in the following work.

SHC Has the Potential for Large-Scale Production of Hopene

Enzymatic catalysis has not been used in the large scale preparation of hopene until now. Discovery of the soluble OUC-SaSHC will change this status, because the steady activity and easy purification steps of OUC-SaSHC will greatly decrease the cost of squalene-to-hopene reaction. Therefore, scale up of the OUC-SaSHC catalyzed hopene synthesis from squalene was investigated.

Reaction time is an important factor needed to be optimized. Conversion of squalene into hopene by OUC-SaSHC was carried out in 100 mM sodium phosphate buffer (pH 7.0) containing 3 mM squalene, 0.5% Tween 80 and 1 mM MnCl₂ at 30°C. As shown in Figure 6, when the OUC-SaSHC concentration is 0.006 mg/mL or more, hopene concentration is higher than 1.12 mg/mL at 48 h. This indicated that as long as the reaction time is enough, over 90% of squalene could be converted into hopene. Moreover, when using 0.008 mg/mL or 0.01 mg/mL...
OUC-SaSHC, hopene was produced with the concentration of 1.08 mg/mL or 1.11 mg/mL at 36 h, which was similar with that (1.12 mg/mL) using 0.006 mg/mL OUC-SaSHC at 48 h (Figure 6A). This suggested that reaction time could be shortened by increasing biocatalyst concentration. Furthermore, in the case of using 0.008 mg/mL or 0.01 mg/mL OUC-SaSHC, the productivity of hopene was 0.030 g/L/h or 0.031 g/L/h at 36 h, both of which decreased after 36 h (Figure 6B). Therefore, the reaction time was optimized as 36 h, with the corresponding OUC-SaSHC concentration of 0.008 mg/mL.

Substrate concentration was further optimized at optimal reaction time of 36 h. As shown in Figure 7, when the substrate concentration was 5 mM or 10 mM, 100% of the squalene was converted into hopene by 0.08 mg/mL OUC-SaSHC. With the increase of substrate concentration, the conversion ratio decreased gradually. When the substrate concentration increased from 20 mM to 25 mM, the conversion ratio decreased from 90.19 to 82.52%, and the yield of hopene increased from 7.14 to 7.54 mg/mL (Figure 7). This meant 25% of substrate increase only resulted in 5.6% of product increase, which caused waste of squalene. Therefore, 20 mM of squalene was selected as the optimal substrate concentration.

With 20 mM of squalene as substrate, there were still 9.81% squalene remained in the reaction mixture. This suggested that further separation process was needed to obtain purified...
hopene. One effective way to avoid separation is to improve the conversion ratio by adding more enzymes. In Figure 8, it showed that when the OUC-SaSHC concentration was 0.14 mg/mL, the conversion ratio of squalene reached 98.72%, with 7.82 mg/mL hopene produced. It seems that the added OUC-SaSHC increased 75%, while the produced hopene only increased 9.52%. Actually, the preparation program of the soluble OUC-SaSHC is cost-effective, which means the cost of the increased 0.06 mg/mL OUC-SaSHC was much less than the value of increased 0.68 mg/mL hopene after improvement of conversion ratio from 90.19 to 98.72%.

After optimization of the reaction conditions, the efficiency of OUC-SaSHC converting squalene to generate hopene has been greatly improved. The soluble properties and the high conversion ratio indicate that OUC-SaSHC has potential for large scale production of hopene. In order to further verify the feasibility of applying OUC-SaSHC for large scale production of hopene, a 50-fold scale-up reaction was conducted with the reaction volume of 100 mL. As shown in Figure 9, the reaction was carried out in 100 mM sodium phosphate buffer (pH 7.0) containing 0.5% Tween 80, 1 mM MnCl₂, 20 mM squalene and 0.14 mg/mL OUC-SaSHC at 30°C. After reaction of 36 h, 8.07 mg/mL hopene was produced with the conversion ratio of 99%. Better data might benefit from the better mass transfer efficiency in reaction of 100 mL than that of 2 mL. This result indicated that to further improve the scale of the catalytic conversion system was feasible and expecting for producing more hopene with high purity. SHC can also be used to produce other terpenoids, such as epoxydammarane (Ideno et al., 2018) and ambrox (Eichhorn et al., 2018; List and Sun, 2018). These valuable terpenoids can also be produced at a scale-up level by using the soluble OUC-SaSHC found in this study.

CONCLUSION

In summary, we discovered a novel SHC named OUC-SaSHC from S. albolongus. The relative molecular mass of OUC-SaSHC is 69 kDa. When using squalene as substrate, the specific activity of OUC-SaSHC is 1138 U/mg. The soluble OUC-SaSHC could be expressed and purified easily. Cost-effective preparation of OUC-SaSHC makes it possible to produce value-added hopene from squalene at a large scale. Under the optimal reaction condition, 0.81 g hopene with the purity of 99% was produced in a 100 mL catalytic conversion system. Our work shows the feasibility of large scale hopene production from squalene by the soluble OUC-SaSHC. OUC-SaSHC can also be used to large-scale produce other valuable terpenoids which can be synthesized by SHC.

DATA AVAILABILITY STATEMENT

The datasets generated for this study can be found in the GenBank Accession No. MH121056.

AUTHOR CONTRIBUTIONS

ZL, XM, CX conceived and designed the experiments. YZ, JS, W-CH performed the experiments. YZ, ZL analyzed the data. XM contributed reagents, materials, and analysis tools. YZ, ZL wrote the manuscript.

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REFERENCES

Abe, I. (2007). Enzymatic synthesis of cyclic triterpenes. Nat. Prod. Rep. 24, 1311–1331. doi: 10.1039/b616857b
Bradley, A. S., Pearson, A. J., Sáenz, P., and Marx, C. J. (2010). Adenosylhopane: the first intermediate in hopanoid side chain biosynthesis. Org. Geochem. 41, 1075–1081. doi: 10.1016/j.orggeochem.2010.07.003
Chen, Z., Washio, T., Sato, M., and Suzuki, Y. (1995). Cytotoxic effects of several hopanoids on mouse leukemia L1210 and P388 cells. Biol. Pharm. Bull. 18, 421–423. doi: 10.1248/bpb.18.421
Doughty, M. D., Dieterle, M., Sessions, A., Fischer, W. W., and Newman, D. K. (2014). Probing the subcellular localization of hopanoid lipids in bacteria using NanoSIMS. PLoS One 9:e84455. doi: 10.1371/journal.pone.0084455
Eichhorn, E., Locner, E., Guillerme, S., Wahler, D., Fourage, L., and Schilling, B. (2018). Biocatalytic process for (-)-ambrox production using squalene hopene cyclase. Adv. Synth. Catal. 360, 2339–2351. doi: 10.1002/adsc.201800132
Frickey, T., and Kannenberg, E. (2009). Phylogenetic analysis of the triterpene cyclase protein family in prokaryotes and eukaryotes suggests bidirectional lateral gene transfer. Environ. Microbiol. 11, 1224–1241. doi: 10.1111/j.1462-2920.2008.01851.x
Fukuda, Y., Watanabe, T., and Hoshino, T. (2018). Mutated variants of squalene-hopene cyclase: enzymatic syntheses of triterpenes bearing oxygen-bridged monocycles and a new 6,6,6,6,6-fused pentacyclic scaffold, named neogammacerane, from 2,3-oxidosqualene. Org. Biomol. Chem. 16, 8365–8378. doi: 10.1039/c8ob02009d
Ghimire, G. P., Oh, T. J., Lee, H. C., and Sohng, J. K. (2009). Squalene-hopene cyclase (Spterp25) from Streptomyces peucetius: sequence analysis, expression and functional characterization. Biotechnol. Lett. 31, 565–569. doi: 10.1007/s10529-008-9903-2
Giner, J., Rocchetti, S., Neunlist, S., Rohmer, M., and Arigoni, D. (2005). Detection of 1,2-hydride shifts in the formation of euph-7-ene by the squalene-tetrahymanol cyclase of Tetrahymanella pyriformis. Chem. Commun. 24, 3089–3091. doi: 10.1039/b504300g
Gustafsson, C., Vassiliev, S., Kurten, C., Syren, P. O., and Brinck, T. (2017). MD simulations reveal complex water paths in squalene-hopene cyclase: tunnel-obstructing mutations increase the flow of water in the active site. ACS Omega 2, 8495–8506. doi: 10.1021/acsomega.7b01804
Hoshino, T., Shimizu, K., and Sato, T. (2004). Deletion of the Gly600 residue of Streptomyces violaceusnigrus squalene-hopene cyclase alters the substrate specificity into that of the eukaryotic-type cyclase specific to (3S)-2,3-oxidosqualene. Angew. Chem. Int. Ed. Engl. 43, 6700–6703. doi: 10.1002/anie.200461523
Sáenz, J. P., Sezgin, E., Schwille, P., and Simons, K. (2012). Functional convergence of hopanoids and sterols in membrane ordering. *Proc. Natl. Acad. Sci. U.S.A.* 109, 14236–14240. doi: 10.1073/pnas.1212141109

Schmerk, C., Bernardi, M., and Valvano, M. (2011). Hopsanoid production is required for low-pH tolerance, antimicrobial resistance, and motility in *Burkholderia cenocepacia*. *J. Bacteriol.* 193, 6712–6723. doi: 10.1128/JB.05597-11

Seckler, B., and Poralla, K. (1986). Characterization and partial purification of squalene-hopene cyclase from *Bacillus acidocaldarius*. *BBA Gen. Subj.* 881, 356–363. doi: 10.1016/0304-4165(86)90027-9

Seitz, M., Klembersberger, J., Siebenhaller, S., Breuer, M., Siedenburg, G., and Jendrossek, D., et al. (2012). Substrate specificity of a novel squalene-hopene cyclase from *Zymomonas mobilis*. *J. Mol. Catal. B ENZYM* 84, 72–77. doi: 10.1016/j.molcatb.2012.02.007

Seitz, M., Syren, P. O., Steiner, L., and Klembersberger, J. (2013). Synthesis of heterocyclic terpenoids by promiscuous squalene-hopene cyclases. *Chembiochem* 14, 436–439. doi: 10.1002/chem.201300018

Siedenburg, G., and Jendrossek, D. (2011). Squalene-hopene cyclase. *Appl. Environ. Microbiol.* 77, 3905–3915. doi: 10.1128/AEM.00300-11

Siedenburg, G., Jendrossek, D., Breuer, M., Juhl, B., Pleiss, J., Seitz, M., et al. (2012). Activation-independent cyclization of monoterpenoids. *Appl. Environ. Microbiol.* 78, 1055–1062. doi: 10.1128/AEM.07059-11

Sodenkamp, C., and Geiger, O. (2016). Bacterial membrane lipids diversity in structures and pathways. *FEMS Microbiol. Rev.* 40, 133–159. doi: 10.1093/femmicro/fvw008

Stroeve, P., Pettit, C. D., Vasquez, V., Kim, V. L., and Berry, A. M. (1998). Surface active behavior of hopanoid lipids: bacteriohopanepentol and phyllocacteate monomer bacteriohopanoptol. *Langmuir* 14, 4261–4265. doi: 10.1021/la980013e

Takahashi, K., Sasaki, Y., and Hoshino, T. (2018). Squalene-hopene cyclase: on the polycyclization reactions of squalene analogues bearing ethyl groups at positions C-6, C-10, C-15, and C-19. *Eur. J. Org. Chem.* 2018, 1477–1490. doi: 10.1002/ejoc.201800010

Tippelt, A., Jahne, L., and Poralla, K. (1998). Squalene-hopene cyclase from *Methylcoccus capsulatus* (bath): a bacterial production hopanoids and steroids. *Biochim. Biophys. Acta* 1391, 223–232. doi: 10.1016/S0005-2760(97)00212-9

Wendt, K. U. (2005). Enzyme mechanisms for triterpene cyclization: new pieces of the puzzle. *Angew. Chem. Int. Ed.* 44, 3966–3971. doi: 10.1002/anie.200538293

Wendt, K. U., Lenhart, A., and Schulz, G. E. (1999). The structure of the membrane protein squalene-hopene cyclase at 2.0 A resolution. *J. Mol. Biol.* 286, 175–187. doi: 10.1006/jmbi.1998.2470

Wendt, K. U., Poralla, K., and Schulz, G. E. (1997). Structure and function of a squalene-hopene cyclase. *Science* 277, 1811–1815. doi: 10.1126/science.277.5333.1811

Wu, T., Ye, L., Zhao, D., Li, S., Zhang, B., et al. (2017). Membrane engineering—a novel strategy to enhance the production and accumulation of β-carotene in *Escherichia coli*. *Metab. Eng.* 43, 85–91. doi: 10.1016/j.ymben.2017.07.001

Yoder, R. A., and Johnston, J. N. (2006). A case study in biomimetic total synthesis: polyolefin carbocyclizations to terpenes and steroids. *ChemInform* 37, 4730–4756. doi: 10.1002/chin.200617620.10.1002/chin.200617620.10.1002/chin.200617620

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