Applications of protein engineering in the microbial synthesis of plant triterpenoids

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ABSTRACT

Triterpenoids are a class of natural products widely used in fields related to medicine and health due to their biological activities such as hepatoprotection, anti-inflammation, anti-viral, and anti-tumor. With the advancement in biotechnology, microorganisms have been used as cell factories to produce diverse natural products. Despite the significant progress that has been made in the construction of microbial cell factories for the heterogeneous biosynthesis of triterpenoids, the industrial production of triterpenoids employing microorganisms has been stymied due to the shortage of efficient enzymes as well as the low expression and low catalytic activity of heterologous proteins in microbes. Protein engineering has been demonstrated as an effective way for improving the specificity, catalytic activity, and stability of the enzyme, which can be employed to overcome these challenges. This review summarizes the current progress in the studies of Oxidosqualene cyclases (OSCs), cytochrome P450s (P450s), and UDP-glycosyltransferases (UGTs), the key enzymes in the triterpenoids synthetic pathway. The main obstacles restricting the efficient catalysis of these key enzymes are analyzed, the applications of protein engineering for the three key enzymes in the microbial synthesis of triterpenoids are systematically reviewed, and the challenges and prospects of protein engineering are also discussed.

1. Introduction

Triterpenoids are a class of secondary metabolites with diverse structures widely existing in plants. As the main active ingredients in traditional herbal medicines, they have pharmacological activities such as liver protection, anti-inflammation, anti-virus, and anti-tumor [1–3]. The structural diversity of triterpenoids makes them potential lead compounds for new drugs, and thus attracted the attention of the pharmaceutical industry [4]. Triterpenoids are mainly derived from the cyclization of 2,3-oxidosqualene, in which oxidosqualene cyclases catalyze the cyclization of 2,3-oxidosqualene to form triterpenoid skeletons, such as α-amyrin, β-amyrin, and lupeol, laying the foundation for the structural diversity of triterpenoids. Subsequently, the triterpenoid skeletons could be further mono-oxygenated by cytochrome P450s to produce different triterpenoid compounds such as ursolic acid, oleanolic acid and betulinic acid, etc. (Fig. 1). However, the clinical application of many triterpenoids is limited by their low solubility and bioavailability [5]. The glycosylation reaction mediated by UGTs can transfer sugar groups such as glucose and galactose from sugar donors to the hydroxyl or carboxyl groups of triterpenoids, therefore improving the water-solubility and bioavailability of triterpenoids. The products of the glycosylation reaction, triterpenoid saponins, thus have better pharmacological activities [6].

In recent years, with the development of high-throughput sequencing technologies and web-based BLAST servers, as well as the publication of lots of plant transcriptome databases, more and more key enzymes for terpenoid synthesis have been mined. Meanwhile, the synthetic pathways of many terpenoids in plants have been analyzed and obtained [7]. Therefore, the synthesis of plant natural products using microbial cell factories becomes a research hotspot. For example, CYP72A63 has been identified to be a P450 enzyme involved in the synthesis of glycyrrhetinic acid (GA), and can significantly increase its yield [8,9]. When expressed in microbes, however, the plant-derived enzymes suffer from low catalytic activity and poor specificity, limiting the industrial production of natural products. Rational or semi-rational design based on crystal structure analysis or molecular
dynamics simulations for protein engineering has been used to overcome these challenges [10,11]. Notably, the crystal structures of enzymes are of crucial importance in the rational or semi-rational design of protein engineering. While the known crystal structures for enzymes in the synthesis of plant triterpenoids are limited, indirect methods such as sequence alignment, protein homology modeling, and molecular simulation are employed in rational or semi-rational protein design. In addition, AlphaFold 2, a new tool in structural biology, can predict the structure of a typical protein in minutes with unprecedented accuracy. It can enable protein structure elucidation techniques to keep the pace of the genomic revolution in the future [12]. To date, key enzymes in several synthetic pathways have been modified via protein engineering, endowed with specific chemoselectivity, substrate selectivity, and catalytic activity [13,14]. These advancements facilitate the application of protein engineering in the microbial synthesis of triterpenoids and triterpenoid saponins. This review mainly introduces the current studies of key enzymes, OSCs, P450s, and UGTs, in the synthetic pathway of triterpenoids and triterpenoid saponins. An overview of the optimization of OSCs, P450s and UGTs by protein engineering is also provided.

2. Protein engineering

As the synthetic pathways of terpenoids in plants have been elucidated constantly, heterologous synthesis of triterpenoids in microorganisms has been achieved through various metabolic regulation strategies. Zhang et al. [15] overexpressed typical rate-limiting enzymes isopentenyl pyrophosphate isomerase (IDI), farnesyl-diphosphate synthase (FFPS), and squalene synthase (SQS) in *Saccharomyces cerevisiae* to enhance the supply of precursors and increase the yield of oleanane triterpenoids by 1.55-fold. Li et al. successfully synthesized the precursor 2,3-oxidosqualene in *Escherichia coli*, followed by the introduction of the enzymes with truncated N-terminal transmembrane domains, producing dammarenediol II with a final yield of 8.63 mg/L [16]. Akin to *Saccharomyces cerevisiae*, *Yarrowia lipolytica* is another excellent host for the production of triterpenoids. Li et al. [17] constructed a recombinant ginsenoside compound K pathway in *Y. lipolytica* which overexpressed MVA rate-limiting enzyme, employed a fused protein of P450 (PPDS) and NADPH-cytochrome P450 reductase (ATR1), leading to a final product of 30.4 mg/L. In order to increase the production of betulinic acid, cytochrome P450 oxidoreductases (CPR) from *Medicago truncatula* and CYP716A180 from *Betula platyphylla* were screened and paired by Jin et al. [18,19]. By co-expressing with ERG1, ERG9, and HMG1 in yeast, the titer of betulinic acid was increased to 51.87 ± 2.77 mg/L. Although the regulation and optimization of metabolic pathway can improve carbon flux to a certain extent, it is difficult to achieve large-scale production for most natural products. One of the bottlenecks
is the poor catalytic activity and substrate specificity of key enzymes.

The emergence of protein engineering has greatly accelerated the industrialization of biosynthesis for triterpenoids and triterpenoid saponins. On the other hand, the update of many open source software and websites provides a theoretical basis and feasibility basis for the development of protein engineering. For example, the online BLAST tool can quickly match homologous templates for target enzymes from massive databases; The MODELER program in Discovery Studio v2.5, SWISS-MODEL, or AlphaFold2 [20–22] can use the method of homology modeling to predict the three-dimensional structure of a known sequence; The quality of the 3D model structure is evaluated by the software such as Seq Identity, GMQE, QMEAN, ProCheck, Molprobity, and Errat; The docking of the target protein and the catalytic product is simulated by Autodock Vina [23,24], which predicts the conformation of the ligand at the target binding site, calculates the binding energy, and predicts specific amino acid residues. The activity centers of key enzymes, catalytic products, and functional residues have been studied by researchers through computer-aided design combined with the crystal structures resolved. The section focuses on the application of protein engineering in the optimization of OSCs, P450s, and UGTs and the research progress in triterpenoid synthesis.

2.1. Protein engineering of OSCs

2.1.1. Crystal structure of OSCs

With the in-depth study of the three-dimensional structure of enzyme molecules and the rapid development of bioinformatics, rational design based on crystal structure followed by experimental analysis and computer simulation has become an important method for enzyme molecule modification. Therefore, protein engineering of OSCs from the perspective of structural biology and biochemistry is necessary to improve their performance and applicability. OSC is a membrane protein with a complex structure, which is difficult to be analyzed. To date, only two crystal structures of the human membrane protein OSC, namely OSC-lanosterol complex structure (PDB ID: 1W6K) and OSC-inhibitor Ro 48-8071 complex structure (PDB ID: 1W6J) have been resolved [25].

The crystal structure of OSC-lanosterol (Fig. 2) shows that one hydrogen bond is formed between the lanosterol 3-hydroxy group and the side chain of D455. This reveals the cyclization mechanism of 2,3-oxidosqualene, where D455 donates a proton to the epoxy group of 258. This is consistent with the catalytic mechanism of the OSC-lanosterol complex structure. Due to the difficulty in obtaining the crystal structure of OSCs enzyme, almost all the studies related to the molecular modification of OSCs in recent years have been carried out using human OSC as the template, such as ERG7 from S. cerevisiae, and AshS1 from Avena strigosa [26,27].

2.1.2. Regulation of product preference through protein engineering of OSCs

From the perspective of natural synthetic pathways, OSCs cyclize 2,3-oxidosqualene to generate a variety of compound skeletons for plant triterpenoid through a Chair-Boat-Chair (CBC) or a Chair-Chair-Chair (CCC) conformation and finally forms triterpenoids (e.g. cucurbitadienol, lupeol, and glutinol) (Fig. 3). At present, a considerable number of OSCs have been isolated, purified, cloned, and characterized through plant or transcriptome database mining. For example, Yu et al. [39] identified MdOSC1 from apple, which increased the production of α-aminin in S. cerevisiae by 5.8-fold with a titer of 11.97 ± 0.61 mg/L. However, in addition to the target compound, there are small amounts of β-aminin and lupin alcohol in the product. The MdOSC4 and MdOSC5 identified from apples by Andre et al. [40] also have product conformation, among which germanicol and lupeol are the main products. SITTS2 from tomato catalyzed the production of triterpenoids with six compounds (e.g. α-, β- and δ-aminin) [41]; KmCS from Kandelia candel was verified to be highly homologous to lupeol synthase, but the content of lupeol only accounted for half of the total product [42]. It can be concluded the plant-derived enzymes often have problems of poor specificity when introduced into microbes. Therefore, protein modification of the key enzymes that have been mined is crucial, and the discovery of new enzymes provides a resource-rich database for protein engineering.

Combining advanced computational techniques and molecular biology methods, we can deeply understand the structural characteristics and functions of the enzyme. The correct docking of the substrate can provide important candidates for mutation, including active sites such as key catalytic or binding residues (Table 1). Liang et al. [27] discovered two novel hopane-type triterpenoid synthases, namely, AshS1 and AshHS2, which produce hopenol B and hop-17(21)-en-3β-ol, respectively. AlphaFold2 modeling and QM/MM calculation showed that aromatic residues F257 and F279 of AshS1 promoted the rearrangement reaction by interacting with hopyl C22 cations. The bulky side chain of Y410 affects the interaction between F257 and carbocation through steric hindrance, which is considered to be the key element to product differentiation. When Y410 was mutated to a small alanine (Y410 corresponds to A410 in AshS2), the mutant mainly catalyzed the production of hop-17(21)-en-3β-ol, which was the same as AshS2. This result reveals the possibility of changing product preference through amino acid substitution mutations. Eukaryotic OSCs specifically recognize (3S)-2,3-oxidosqualene as a substrate, rather than squalene or (3R)-2,3-oxidosqualene. Tsutou et al. [43] compared the sequences of OSCs with some prokaryotic squalene-hopene cyclase (SHCs), and proposed that G600 was important for SHC’s recognition of multiple substrates according to the fact that G600 was highly conserved in SHCs but not in OSCs. They knocked out G600 of squalene cyclase from Allomyces alslexicus and the obtained mutant only took (3S)- and 3-oxidosqualene as the substrate. Conserved motifs, 256LIC257 and 258H259, are present in lupeol synthase, β-aminin synthase, respectively, and are actively involved in product differentiation. Kushiro et al. [33] identified the product from lupeol to β-aminin by mutating L256 of lupeol synthase to L256W. Similarly, W259 of β-aminin synthase is replaced by leucine, resulting in the termination of the reaction at the Lupeyl cation stage, and the product is dominated by lupeol. It can be seen that the unique active sites or functional domains of OSCs play an important role in stabilizing carbocation intermediates and product specificity. Interestingly, corresponding residues do not
necessarily produce the same interactions. For example, the F474 residue of β-amyrin synthase (EtAS) from Euphorbia tirucalli corresponds to F365 of SHC, which stabilized the transient cation via cation–π interaction. But when F474 was mutated to aliphatic amino acids lacking π-electrons such as Val, Leu and Met, the polycyclic reaction still proceeded and produced pentacyclic triterpenoids without terminating at the bicyclic stage, which proved that SHC F365 has a distinct mode of action from EtAS F474 [29,30]. Another example is the catalytic efficiency of EtAS, which was significantly reduced after the mutation from phenylalanine to tryptophan at site 728. F728 promotes ring-expansion processes by stabilizing the cationic intermediates, while the steric bulk near the active site also affects enzyme-substrate binding, actively participating in the polycyclization pathway and the direction of the folded conformation [32].

2.1.3. Improved product specificity through protein engineering of OSCs

Although OSCs have generated sequence diversity during evolution, terpenoid synthases belonging to the same type have conserved motifs near the catalytic center (Fig. 4). For instance, β-amyrin synthase has a highly conserved MWCYCR motif, while lupine alcohol synthase keeps the conserved MLCYCR motif [28]. Commonly seen approaches based on the conserved motifs to improve the product specificity and catalytic activities of enzymes have been developed as follows: infrequent amino acids in conserved motifs are identified by comparing multiple amino acid sequences and are replaced with consensus amino acids or treated with site-directed mutations. Taking α-amyrin synthase as an example, there is seldom OSC that can specifically synthesize α-amyrin in nature. By comparing the amino acid sequences of ItOSC2 and representative plant OSCs, Wu et al. [28] identified Y531/L256/L258 as the three key active sites of ItOSC2. While amino acids I256 and L258 are close to lupanyl cation intermediates, preventing premature deprotonation of intermediates, Y531 is involved in the correct orientation of the lupanyl cation and other late intermediates, facilitating the deprotonation of the intermediates through the formation of hydrogen bond network between the hydroxyl group and water molecules. This results in highly specific α-amyrin products. However, site-directed mutagenesis based on sequence conservation analysis has certain limitations [44]. Since multiple amino acid mutations are prone to unpredictable epistasis, there are usually no more than 4 mutation sites, and the improvement in catalytic activity is limited to within 10-fold.

2.2. Protein engineering of P450s

In general, P450 oxidase and its reductase can specifically oxidize different sites of the triterpenoid skeleton to produce triterpenoid compounds with various structures by introducing hydroxyl, carboxyl, or epoxyl groups. P450s have attracted much attention due to their wide range of biocatalyst activities. However, as membrane-localized proteins, plant P450s have low expression levels, low activity, and poor stability in microorganisms, which have been considered key bottlenecks in pathway engineering. In this section, protein engineering in improving the catalytic properties of P450 and increasing the yield of

Fig. 3. Formation of carbocation intermediates and triterpenoid skeletons under the catalysis of OSCs.
The redox partner of P450. It is considered to be an important factor from NAD(P)H to the heme center of Class II microsomal P450s, and is affecting the catalytic activity of P450. Therefore, P450s mining and chaperones, protein fusion, engineering of transmembrane regions, and Protein engineering of OSCs.

Table 1

| OSC   | GenBank    | Substrates       | Products   | Key Residues and Mutants | Mechanism of action | Results | References |
|-------|------------|------------------|------------|--------------------------|---------------------|---------|------------|
| ItOSC2| MT104569.1 | 2,3-oxidosqualene | β-amyrin   | Y531/L256/258            | formed the unusual Y-LL triplet at the active site | Yield greater than 87% | [28]       |
| β-amyln| AB20470.1  | 2,3-oxidosqualene | β-amyrin   | F474                    | Appropriate volume of space near the B-ring formation site | F474L and F474 M increased yield by 60% and 82%, respectively | [29,30]   |
| β-amyln| AB20670.1  | 2,3-oxidosqualene | lupeol     | F413/Y259/W257           | Stabilization of intermediaries cations via cation–π interactions | Generating lupeol synthase from the β-amyln synthase | [31]       |
| β-amyln| AB20670.1  | 2,3-oxidosqualene | triterpenoids | F728        | Stabilizes cationic intermediates and facilitates the ring expansion process | First demonstration of the criticality of cation–π interaction for OSCs | [32]       |
| β-amyln| AB009030.1 | 2,3-oxidosqualene | lupeol     | W259L                  | W259 controls β-amyrin formation by stabilizing oleanyl cation | Produced lupeol as a major product | [33]       |
| lupleol synthase (Pruv) | AB025343.1 | 2,3-oxidosqualene | β-amyrin   | L256W                  | L256 controls lupeol formation by stabilizing dammaranyl cation | Produced exclusively β-amyrin | [33]       |
| AshS1 | OM401324.1 | 2,3-oxidosqualene | hopenol B  | F257/C366/Y410          | Promote rearrangement; Involved in the deprotonation process | Y410A mutant mainly produces hop-17(21)-en-β-ol | [27]       |
| AshS2 | OM401325.1 | 2,3-oxidosqualene | hop-17(21)-en-β-ol | A410       | The side chain size of site 410 leading to a difference in deprotonation positions | Could synthetize hopenol B alone | [27]       |
| β-amyln| JK467352.1 | 2,3-oxidosqualene | β-amyrin   | Y560                   | Y560 and N610 together form a hydrophilic environment and improve the deprotonation of C13 | GaAS2 with Y560 had accumulated more oleanolic acid | [34]       |
| α-amyln| AB007002.1 | 2,3-oxidosqualene | mono- and pentacyclic triterpene | D377C/V380E/V381A/L258W | The triple amino acids are responsible for the initiation of squalene cyclization | Cyclized 2,3-oxidosqualene to give mono- and pentacyclic triterpene | [35]       |
| α-amyln| LC464980.1 | 2,3-oxidosqualene | α-amyrin   | L258W                  | L258 mediates the lupeol specificity of OSCs | From mainly producing pentacyclic products to mainly producing tetracyclic products | [36]       |
| OSC(OrOS) | MG932735.1 | 2,3-oxidosqualene | oryctacinol | Y257                   | The pi-electrons of Y257 stabilize the intermediate cation through cation–π interaction | OsOS2767L-E resulted in complete loss of function | [37]       |
| MdOSC1 | FJ032006.1 | 2,3-oxidosqualene | α-amyrin   | N111/P250H/P373S      | Introduced hydrogen bonds and expand the substrate binding pocket | The yield of α-amyrin was increased 11-fold | [38]       |

2.2.1. Enhanced triterpenoid production via protein fusion of CYP and CPR

P450 oxidoreductase (CPR) is responsible for the transfer of electrons from NADPH to the heme center of Class II microsomal P450s, and is the redox partner of P450. It is considered to be an important factor affecting the catalytic activity of P450. Therefore, P450s mining and combining with CPRs for optimized electron transfer efficiency are important. Wang et al. [47] introduced CPR (GuCYP5) from *Glycyrrhiza uralensis* into the engineered yeast strain. This successfully reduced the accumulation of β-amyrin and increased the glycyrrhetic acid titer to 545 μg/L. Lodeiro et al. [48] compared the pairing effects of different CPRs with CYP71B1A1 and found that the catalytic efficiency of AtCPR1 from *Arabidopsis thaliana* was 7.4-fold higher than that of RhCPR from *Rhizopus oryzae*. Jin et al. [18] sequentially expressed LiCPR from *Lotus japonicus* and MtCPR from *M. truncatula* in *Y. lipolytica*, and the production of betulinic acid increased from 25.62 mg/L to 32.33 mg/L.

To improve the efficiency of P450, the fusion protein is employed. The fusion of plant P450 and CPR has many advantages: (1) the fusion protein enhances the coupling between CYP and CPR, as well as improves the electron transfer efficiency; (2) the continuous reaction is accelerated to obtain higher products titer by reducing the physical distance between the two enzymes [49]; (3) the loss of electrons caused by the lack of inner membrane structure is avoided.

It has been reported that the uncoupling of P450 and CPR generates reactive oxygen species (ROS), which can reduce the efficiency of electron transfer and even lead to the death of heterologous host cells in severe cases [50,51]. For example, when protopanaxadiol synthase (PPDS) and CPR (ATR1) are overexpressed in the protopanaxadiol synthesis pathway, the coupling between the two is poor, and a large amount of reactive oxygen species is released, seriously threatening the health of the host *S. cerevisiae*. Linker of different lengths was used by Zhao et al. [52] to construct the self-sufficient PPDS-ATR1 fusion.

Compared with the previous modification, the catalytic activity of all fusion enzymes increased by about 4.5-fold, and the conversion efficiency of dammarenediol-II was as high as 96%. Meanwhile, the growth of host cells was effectively improved. In order to enhance the electron transfer efficiency between CYP716A12 and ATR1, Li et al. [53] fused the two proteins employing the linker GSTSSG. This reduced the precursor β-amyrin and increased the oleanolic acid production in *Y. lipolytica*, reaching 540.7 mg/L. The fusion of *optCYP706M1* and *opt46AtCPR1* accelerated electron transfer from NADPH to CYP706M1, enhanced the oxidation reaction, and the yield of the target product was 6-fold higher [54]. In Prokaryotes where the sites for electron transfer such as the endoplasmic reticulum are lacking, the P450/CPR fusion is more needed. However, in the taxadiene production pathway of *E. coli*, this fixed impression was broken. The yield of taxadiene decreased after the fusion of CYP725A4 with CPR, even though the plasmid copy number and promoter were adjusted or replaced. Although its origin is unsettled, it is considered that the overexpression of CPR has a negative impact. Resource competition or inefficient NADPH utilization may be the reason [55,56].

In addition, the formation of triterpenoids by P450 oxidation may involve the continuous catalysis of multiple enzymes, i.e., the catalytic product of the previous enzyme is the reaction substrate of the next enzyme. Reducing the spatial distance between multiple enzymes through fusion proteins will help improve the transfer efficiency of substrates between them and minimize the loss of intermediates during
Fig. 4. Alignment of partial amino acid sequences of β-amyrin synthases.

Table 2

| P450       | Gene ID     | Source           | Substrates          | Products                                     | Key Residues and Mutants | Mechanism of action                                                                 | Results                          | References |
|------------|-------------|------------------|---------------------|----------------------------------------------|--------------------------|--------------------------------------------------------------------------------------|----------------------------------|------------|
| CYP72A63   | 11410378    | Medicago truncata | 11-oxo-β-amyrin     | glycyrrhetol                                  | L509I                    | Increases the hydrophobicity of the active pocket ends                               | Yield is 90.6%                   | [9]        |
| CYP72A63   | 11410378    | Medicago truncata | glycyrrhetol        | glycyrrhetinic acid (GA)                     | T338S                    | Improve the hydrophilicity of the binding pocket                                    | GA increased by 7.1-fold         | [9]        |
| CYP72A63   | 11410378    | Medicago truncata | 11-oxo-β-amyrin     | 29-OH-11-oxo-β-amyrin                         | L398I                    | Rotate the substrate 180° to reverse the relative positions of C-30 and C-29        | 100% regional selectivity        | [9]        |
| CYP72A65   | 11410378    | Medicago truncata | glycyrrhetol        | GA                                            | T338S/W205A               | Eliminate limitations on proton transfer                                            | 36.4 ± 3.0 mg/L                  | [9]        |
| CYP72A62v2 | 11409246    | Medicago truncata | β-amyrin            | 30-OH-β-amyrin                                | L149/L398                | Further enhance the production of high-value triterpenoid compounds                |                                 | [4]        |
| CYP72A62v2 | 11409246    | Medicago truncata | β-amyrin            | 30-OH-β-amyrin                                | V398L                    | Changed from being regioselective at C-29 to producing only C-30 oxidation products |                                 | [4]        |
| CYP87D20   | 101213710   | cucumber          | cucurbitadienol     | 11-H-cucurbitadienol                          | E286A                    | E286A plays a role in shuttling protons into the active site                         | Decreases the production of 11C-Cool and 11C-20H-Cool | [45]      |
| TwCYP712K1 | 2081630515  | Tripterygium wilfordi | friedelin         | polpunonic acid                               | S110F                    | Made the substrate closer to the reaction centre                                     | S110F increased the production of polpunonic acid | [46]      |
transport. Enzyme cascade protein fusion and synthetic scaffold protein have been proven to be effective tools in reducing the distance [57]. For example, Moon et al. [58] constructed a peptide scaffold using the protein-protein interaction domain and co-localized a variety of key enzymes to increase the effective concentration of isonitol. As a result, the titer of d-glucaric acid was increased by about 5-fold. Baeck et al. [59] utilized the synthetic scaffold to realize the spatial co-localizations of the heterogeneous pathway enzymes in E. coli, resulting in a 3-fold increase in butyrate production.

2.2.2. Improved expression of CYP and CPR by engineering the transmembrane domain

Both CPR and CYP are inner membrane proteins anchored to the endoplasmic reticulum through their N-terminal. In prokaryotes lacking organelle membranes (such as E. coli), these two enzymes cannot be expressed normally and mostly exist in the form of inclusion bodies [60]. Truncation or modification of the N-terminal transmembrane region of P450 oxidase is an effective strategy to regulate P450 activity and solubility in prokaryotes. A stretch of polar charged amino acid AKKTSSKG was inserted into the N-terminal of CYP71A16 to generate a CYP71A16 mutant, resulting in up to ~50 mg/L active mono-oxygenase [61]. To express the CPR from Candida apicola in a functional form in E. coli, Girhard et al. [62] truncated the N-terminal of CPR with different amino acid lengths, and the modified recombinant protein showed higher expression and stronger solubility. This is the first report of eukaryotic CPR to transfer electrons to bacterial P450s. Improving the compatibility between enzyme and host is an important condition for efficient heterologous expression of P450s. Thus, small libraries of N-terminal expression tag chimeras were developed to enhance the expression of hydrophobic P450 enzymes in cell factories. Diterpene synthase EpCYP76A1 has not been previously expressed in microbial hosts, but when its N-terminal was inserted with a tag, the expression of the EpCYP76A1 mutant in E. coli increased by 9-fold [63].

2.2.3. Crystal structure of P450s

As a low soluble membrane protein, the crystal structure analysis of P450 is also difficult to achieve. X-ray diffraction, cryo-electron microscopy, and nuclear magnetic resonance have been applied to the studies of the three-dimensional structure of proteins. To date, however, only several P450 crystal structures such as CYP116B46 heme domain (PDB ID, 6G1I) [64], CYP4B1 (PDB ID: 5eq1. A) [65], and P450BM3 [66] have been obtained. Without structures of P450s proteins involved in triterpenoid biosynthesis, analysis of structure-function relationships cannot be performed comprehensively. Fortunately, even though the sequence similarity of P450 oxidases is only 10%, their three-dimensional architecture is similar [67]. According to this feature, in the absence of high homology P450 oxidase crystal structure, modeling using the resolved crystal structure as a template can also meet the needs of the semi-rational design. Sun et al. [9] used the crystal structure of CYP4B1 with 26.99% and 23.83% homology to CYP72A63 and CYP72A154, respectively, as a template, and conducted homology modeling to study the transformation strategy and precisely tune its catalytic properties to achieve the selective synthesis of target products.

2.2.4. Improved regioselectivity of P450s through protein engineering

In order to solve the problems of low synthesis efficiency and complex by-products of triterpenoids, researchers attempted to obtain candidates of bio-metabolic engineering through sequence database and comparative analysis. However, when plant P450s are expressed in microorganisms, they often show substrate heterogeneity or poor regioselectivity. As a result, some problems such as scattered metabolic flow in the synthetic pathway, low synthesis efficiency, and the coexistence of multiple by-products, often occur and thus limit the effective synthesis of target products. Meanwhile, it aggravates the difficulty of separation and purification of downstream products.

Protein engineering based on rational design or semi-rational design has been proved to be an effective way to improve the regioselectivity of P450s and achieve the specific synthesis of target products. CYP72A63 from M. truncatula has catalytic activity on the C-29 and C-30 positions of 11-oxo-β-amyrin, resulting in a mixed product of various structural analogs [9]. After the mutation from threonine to hydrophilic serine at site 338 of CYP72A63, the distance between glycurrhetol C-30 OH and heme-Fe is drawn closer by hydrophilic interaction, and C-29 is pushed away from heme-Fe. Therefore, the regional selectivity of the mutant T338S to C-30 was improved, and the yield of glycurrhetinic acid was increased 7.1-fold. When the leucine at position 398 was mutated to isoleucine, the substrate 11-oxo-β-amyrin rotated nearly 180° and the relative positions of C-30 and C-29 were reversed. At this point, the mutant generated a highly regioselective 29-OH-11-oxo-β-amyrin, an isomer of glycurrhetol. CYP72B01 from Pinus taeda has a certain degree of substrate promiscuity, and its substrate binding site is almost entirely hydrophobic residues that may mediate interactions with lipophilic substrates. Ignea et al. [68] mutated these residues into amino acids with different volumes and sizes but still hydrophobicity. The shape of the binding cavity of the mutant and the way of binding with the substrate were changed. For example, the G359A mutant retained only the activity of oxidizing C-18 and produced 18-hydroxy-miltiradiene with a 2-fold increase in production efficiency.

Site-directed mutagenesis based on sequence conservation analysis is also beneficial to find important amino acid residues, particularly for those involved in substrate and product specificity. Fanani et al. [4] identified L149 and L398 as the key amino acids to regional selectivity of CYP72A63 to β-amyrin C-30 through structural analysis of the binding protein model based on amino acid sequence alignment. When V398 of CYP72A62v2 was replaced by leucine, the mutant changed from being regioselective at C-29 to producing only C-30 oxidation products such as 30-hydroxy-β-amyrin.

2.2.5. Enhanced catalytic activity of P450s through protein engineering

The proton delivery pathway is important for the catalytic activity of P450s. Molecular docking results of CYP72A63 with the substrate 11-oxo-β-amyrin suggest that K236 has the function of transferring protons from solvent to alcohol-acid pair, while the bulky side chain of W205 may hinder proton transfer, thereby limiting the catalytic activity of the enzyme. After the mutation of tryptophan into alanine with a small volume, the restriction of steric hindrance on proton transfer was removed, and the double mutant CYP72A63 (T338S/W205A) resulted in the yield of glycurrhetinic acid reaching 36.4 ± 3.0 mg/L [9]. Huang et al. remodeled the CYPs microenvironment by increasing NADPH supply based on overexpressing CYP72A63 (T338S/W205A) to achieve de novo synthesis of glycurrhetic acid 3-O-mono-β-D-glucuronide (GAMG) in S. cerevisiae, and the yield was increased from 0.02 µg/L to 92.00 µg/L [69]. Thus, unblocking the restriction that impedes proton transport is an effective way to increase specific P450 activity.

Seifert et al. conducted systematic computational analysis for the crystal structures of 29 P450s and the sequences of 6379 CYPs. Their study showed that two amino acids adjacent to the heme center were both hydrophobic, a phenomenon that existed in most of the analyzed P450s. F87 and A328 of P450BM3 were indicated to be the corresponding two hydrophobic residues. By replacing them with nonpolar amino acids (e.g. Ala, Val, Phe, Leu, and Ile), the mutants showed similar or higher activity. In addition, the large volume amino acid Phe87 located near the binding pocket is also considered to be the key active site [70]. Mantovani et al. [71] mutated phenylalanine into small volume alanine (F87A), which enhanced the coupling efficiency of the epoxidation of uscatadiene, confirming that steric hindrance does affect the binding of the enzyme to the substrate.

2.3. Protein engineering of UGTs

UDP-glycosyltransferase mediates the glycosylation reaction of triterpenoids. The resulting compounds are called triterpenoid saponins,
which usually have better physicochemical properties and are easier to be stored and transported in plants. From the perspective of structural characteristics, UDP-glycosyltransferase has two binding domains. One domain is located in a highly conserved region consisting of 44 amino acids at the C-terminus, known as the plant secondary product glycosyltransferase (PSPG) motif, involved in the recognition and binding of sugar donors. The other is the sugar receptor binding domain, which is mainly involved in the recognition and binding of sugar receptors. The amino acid with high N-terminal variability is an important reason for the poor substrate selectivity and specificity of glycosyltransferases. The highly conserved histidine, located between the binding domain of sugar donor and sugar receptor, is the catalytic center of glycosyltransferase, and the neighboring aspartic acid plays a conformational and charge balancing role. The mutation of histidine and/or aspartic acid leads to the loss of enzyme activity [72], suggesting that the catalytic pair of His-Asp is an indispensable key amino acid residue (Fig. 5). Glycosylation promotes the structural and functional diversity of natural products and provides many possibilities and opportunities for the development of the food and pharmaceutical industries. For example, while most of the basic glycosides in ginseng, such as protopanaxadiol and protopanaxatriol, have diverse pharmacological activities, ginsenosides (such as ginsenoside Rh2 and ginsenoside Rg3) are the main bioactive ingredients of ginseng and have better applications in the pharmaceutical industry [73]. The glycosyltransferase UGT73C11 was used to catalyze the C-3 hydroxyl of GA to GAMG, which has a sweetness of 941-fold higher than that of sucrose and has better solubility and wider pharmacological activity [74,75]. Precise glycosylation of C-3 OH of triterpenoids can efficiently produce higher value-added derivatives such as GA-3-O-glucose, calenduloside E, and oleanolic acid 3-O-glucose.

Even though the microbial synthesis of some terpenoid saponins has been achieved, it is still a huge challenge to realize their industrialization due to the following difficulties in the application of reported UGTs. Firstly, the low regioselectivity of UGT inhibits the efficient production of target products. For example, UGT73C33 from licorice shows high activity towards C-3 OH and C-30 COOH of pentacyclic triterpenoids but produced three glycosides due to poor substrate selectivity [76]. The glycosyltransferase UGT73C11 from Barbarea vulgaris displays catalytic activity for oleanolic acid C-3 OH and C-28 COOH, resulting in mixed products, which increases the cost of downstream separation and purification [77]. Secondly, the number of introduced sugar groups affecting the pharmacological activity is difficult to control [78,79]. The C-3 OH of GA is more likely to be introduced into two glucuronic acid groups to generate glycyrrhizic acid (GL) than to introduce one to generate GAMG. Meanwhile, the bioavailability of GL is much lower than that of GAMG [77]. Thirdly, the low catalytic activity of UGT leads to the low yield of terpenoid saponins in microbes and thus limits the industrial applications of microbial synthesis. The glucurononyltransferase OAGT identified by Tang et al. [80] from the transcriptome data of Panax zingiberensis showed the catalytic activity of oleanolic acid C-3 OH after recombinant expression in E. coli. However, the target product, calenduloside E, was only obtained at a very low yield. The glycosyltransferase GgGT14 identified in licorice by bioinformatics can specifically synthesize GAMG, but the conversion rate is only 40% [81]. UGT74AG5 expressed in S. cerevisae showed high regioselectivity for ursoyl acid C-28 COOH, and synthesized a rare ursoyl acid 28-O-β-D-glucopyranoside, but the yield was very low [82]. Fourthly, the high specificity of UGT for sugar donors restricts its applications. For example, soyaosapogenol glucuronosyl-transferase (UGASGT), isolated from the seeds of Glycine max, synthesizes oleanolic acid 3-O-β-D-glucuronide using UDP-glucuronic acid rather than UDP-glucose as the sugar donor [83]. The last one is the substrate heterogeneity of UGT. With UDP-Glc, UDP-Rha, UDP-Gal, UDP-Ara, UDP-Xyl as sugar donors, the corresponding sugar groups can be introduced into glycyrrhetinic acid through UGT73C33 and UGT73F24, but the catalytic activity of UGT towards different sugar donors is distinct [76]. GmSGT2 from G. max catalyzes the production of triterpenoid saponins from UDP-Gal, UDP-Glc, UDP-Ara, and UDP-Xyl, of which UGT has the highest catalytic activity towards UDP-Gal [84]. In general, the above five problems still limit the synthesis of terpenoid saponins in microbial cell factories. As shown in Table 3, protein engineering is a helpful strategy to address these problems and achieve high regioselectivity and efficient glycosylation of UGT.

2.3.1. Crystal structure of UGTs

Significantly improving the catalytic efficiency of plant UGT remains a challenge due to the lack of sufficient crystal templates for structural modeling of enzymes and in-depth understanding of their reaction mechanisms. At present, lots of glycosyltransferases (GTs) have been excavated and identified [90]. However, only 13 plant UGT crystal structures have been analyzed and included in Protein Data Bank Database (PDB: https://www.rcsb.org/), such as UGT774F2 from Arabidopsis [91], UGT76G1 from Stevia Rebaudiana [92] and PtUGT1 from Polygonum tinctorum [93]. A smaller number of UGT crystals were identified in the database as responsible for triterpenoid glycosylation, including MtUGT71G1 from M. truncatula [94], ScUGT51 from S. cerevisiae [95], and UGT74AC1 from Siraitia grosvenorii [72]. The crystal structure is important for protein engineering of UGTs. UGT74AC1 from Siraitia grosvenorii (SgUGT74AC1) has a high region-specific but low catalytic activity for glycosylation of mogrol C-3 OH. In order to understand the molecular basis of enzyme and substrate binding, Li et al. [72] used X-ray crystallography to elucidate the crystal structure of SgUGT74AC1 (PDB ID: 6L90) and its complex with substrate UDP-glucose (PDB ID: 6LBZ). In addition, error-prone polymerase chain reaction (EP-PCR) and sequence alignment combined with semi-rational design were used to generate mutants to improve the catalytic efficiency of UGT74AC1, which significantly increased the catalytic capacity of triterpenoid glycosylation by 102-103 times. Finally, they revealed that the enhanced catalytic activity of UGT mutants was attributed to the improved hydrophobic interaction between enzyme and substrate, which promoted the anchoring of triterpenoids at the active center. Currently, the crystal structures of most enzymes have not been resolved, but three-dimensional structures obtained by homology modeling can be used for molecular dynamics simulations to rationally modify proteins.

2.3.2. Enhanced catalytic activity of UGTs through protein engineering

Biochemical or amino acid sequence information is important for selecting mutated amino acids. The key amino acids in the active pocket
usually determine or directly affect the catalytic activity and substrate specificity of the enzyme. Zhang et al. [76] improved the activity of UG773F24 towards glycyrrhetic acid C3-OH by molecular docking and mutation of key residues, leading to the de novo synthesis of GAMG in engineered yeast. They found that L84 is located on a flexible loop at the substrate pocket entrance, which has a certain steric hindrance effect. After L84 is mutated to asparagine, the steric hindrance decreased and the active pocket increased. The combined mutation of L84 and I23 enhanced the activity of UG773F24 (I23G/L84N) by 4.1-fold [76]. Through nine rounds of iterative saturation mutation of amino acid residues in the glycosyltransferase-receptor complex model, UGT51 was engineered into a glycosyltransferase for the efficient synthesis of ginsenoside Rh2. The catalytic efficiency ($k_{cat}/K_m$) was improved by about 1800 times [87]. In the 3D structure of UGTSL2 from S. rebaudiana, N358 is close to the substrate channel and interacts with UDPG through two hydrogen bonds. Mutation of N358 to phenylalanine in the UGTSL2 (N358F) multi-enzyme reaction system increased the production of rebaudioside D by 60% [96]. Despite the key residues interact directly with the substrate, the catalytic results of the triterpenoid glycosyltransferase mutants UGTPg71A29 and UGTPn50 have the function of catalyzing protopanoxadiol (PPD) to ginsenoside Rh2. By comparing the amino acid sequences of them, it was found that two amino acids were missing in UGTPn50, corresponding to A322 and E323 of UGTPg45. The insertion of corresponding amino acids into UGTPn50 was carried out followed by the modification according to beneficial mutations (Q222H and A322V) of UGTPg45. As a result, the catalytic activity of mutant UGTPn50 was 75.4% higher than that of wild type [88]. Yin et al. [97] identified conservative amino acid residues and mutated them through sequence comparison of methionine adenosyltransferase (MAT) from different sources. The mutant MAT (I303V) showed higher catalytic activity than the wild type. These results suggest that analogizing beneficial mutations by sequence comparison is effective to improve the catalytic efficiency of wild-type glycosyltransferase.

| Table 3 | Protein engineering of UGTs. |
|---------|-----------------------------|
| UGT | Substrates | Products | Key Residues and Mutants | Mechanism of action | Results | References |
| UG773F24 | GA | GA-3-O-glucose | I23G/L84 N | Reduce steric hindrance; Expanded binding pocket | Activity increased by 4.1-fold | [76] |
| UGTSr | stevioside | rebaudioside A | H25/D124 | His25 abstracts a proton from the acceptor; Asp124 stabilizes His25 by forming hydrogen bonds | Stability and specificity increased | [85] |
| UGTPg71A29 | Rh1 | Rh1 | Q283 | Affects conformation through hydrophobic interactions | 80% reduction in activity after mutation | [86] |
| UG774A1C | mogrol | mogroside IE | The hydrophobic interactions between enzyme and substrate that play a critical role in anchoring triterpenoids at the active center | Catalytic ability increased by 10–104 folds | [72] |
| UGT51 | propanoaxadiol | ginsenoside Rh2 | S81A/L82A/V84A/A99A/K92A/179K/S129A/N172D | Residues were 9–20 Å away from the docked substrate and involved in a significant conformational change upon substrate binding | Catalytic efficiency (kcat/Km) increased by –1800-fold | [87] |
| UGTPg45 | propanoaxadiol | ginsenoside Rh2 | Q222H/A322V | Enhanced binding to PPD; Affects the configuration of the binding pocket | Rh2 increased by 70% | [88] |
| UG773P12 | GAGM | gycyrrhizin | R32 | Generates salt bridges and hydrogen bonds; Improves affinity with substrates | Identified Arg32 as the essential residue | [89] |

2.3.3. Expanded sugar-donor selectivity through protein engineering

Most UGTs are highly specific for the selection of sugar donors, which not only increase the production cost of compounds, but also limit the structural diversity of saponins. UDP-glucose is the sole sugar donor for UGT92G1 and UGT92G2 from G. max [98]. UG773P12 from licorice prefers to use UDP-glucuronic acid rather than UDP-glucose or UDP-galactose as the sugar donor [89]. Although the chassis host has been modified and optimized, it is still difficult to provide sufficient and diverse glycosylated donors. Modification of glycosyltransferase through the underlying mechanism of substrate selectivity and specificity of glycosylation can effectively solve this problem. Chen et al. [99] analyzed the structural features of UGT78H2 and found that the 23rd residue of the plant secondary product glycosyltransferase (PSPG) motif, which corresponds to the 360th residue in UGT78H2, influences the interactions of enzyme and ligand by forming hydrogen bonds with the donor. When lysine was mutated to asparagine at site 360 and combined with the beneficial mutation N340P, the catalytic activity of the double mutant for UDP-galactose was increased by 23%, and the selectivity of sugar donors was also broadened. In addition to using UDP-glucuronic acid and UDP-galactose, it could also use UDP-glucosyl as a substrate. Similarly, R350 in the PSPG conserved motif of UGT88D7 is a key amino acid for recognizing UDP-glucuronic acid. When arginine was replaced by tryptophan, the utilization efficiency of UDP-glucuronic acid was significantly reduced. Interestingly, the mutant obtained novel UDP-glucosyl transfer activity [100]. The residue R25 side chain of the N-terminal in UGT94B1 is positively charged and directed towards the carboxylic acid group of the UDP-GlcUA sugar donor. Osmani et al. [101] mutated R25 as the active site to produce mutant proteins R25S, R25G, R25K and R25P, which significantly reduced the catalytic activity of the sugar donor UDP-GlcUA. All the other mutants except R25P showed a preference for UDP-Glc, and the catalytic activity was increased 3–4 fold. When D374 of UGT78H2 was mutated to glutamic acid, the amino acid side chain at position 374 was extended and closer to the –OH groups of the sugar, which promoted the formation of
hydrogen bonds between the mutant and UDP-Gal. And UGT76E2 (D374E) showed UDP-Gal activity that was not found in the wild type [102]. The glycosyltransferase UGT73P12 is responsible for the last step in the de novo synthesis of glycyrrhetic acid, which function is to introduce a glucuronic acid group on the basis of GAMG. The positively charged side chain of R32 is considered to be the key amino acid to the specific recognition of UDP-glucuronic acid by UGT73P12. The reason is that under the combined influence of the resultant salt bridge and electrostatic interaction, the positively charged side chain of R32 is oriented toward the negatively charged carboxyl group in UDP-glucuronic acid that provides high affinity for it. Following mutation of arginine to serine, the sugar-donor substrate of UGT73P12 (R32S) was changed from UDP-glucuronic acid to UDP-glucose and UDP-galactose [89].

3. Perspective

Metabolic engineering in recombinant microbial cell factories shows great potential for the production of various triterpenoids. At the same time, candidate genes are mined according to the sequence libraries and databases provided by the PhytoMe project. Genetic tools and sequencing technologies and the databases provided by the PhytoMe project [103], the 1 KP project [104] and the Medicinal Plant Genomics Resource project. Although some progress has been made in the heterologous synthesis of plant triterpenoids by microorganisms, great challenges still exist for their commercial production. The lack of efficient key enzymes is one of the most critical limiting factors, especially the key enzymes in the triterpenoid synthetic pathway such as OSCs, P450s, and UGTs. Recently, the discovery of cellulose synthase-like (CSLs) enzymes as triterpenoid glucuronyltransferases break the traditional view that glycosylation is only catalyzed by UDP-dependent glycosyltransferases and fill a knowledge gap in the biosynthetic pathways of triterpenoid saponins. Jawziak et al. [105] successfully catalyzed medicaginid acid to medicaginid acid 3-O-glucuronide using cellulose synthase-like G (CSLG) belonging to GT family 2. At the same time, they performed site-directed mutagenesis of CSLG, revealing that the active site amino acids S442 and K483 are important for the enzyme to specifically recognize UDP-GlcA. Two other cellulose synthase-like M-subfamily enzymes (AeCSLMs) derived from Aralia elata have been reported to catalyze oleanolic acid and echinocystic acid to produce calenduloside E and echinocystic acid 3-O-glucuronopyranoside, respectively, and have been applied to the de novo synthesis of oleanane-type pentacyclic triterpenes in yeast [106]. The cellulose synthase superfamily-derived glycosyltransferase (CSyGT) can introduce glucuronic acid at C-3 of Glycyrrhetinic acid. Therefore, it has been used in the de novo synthesis of glycyrrhizin [107]. Although the catalytic mechanism of the CSyGTs is still unclear, Sool et al. hypothesise that the transmembrane topology of CSyGTs is critical for triterpene glycosylation. More novel enzymes have been discovered in triterpenoid synthesis. Tao et al. [108] discovered two fungal chimeric class I triterpene synthases that use dimethylallyl diphosphate and isopentenyl diphosphate or hexaprenyl diphosphate as substrates to synthesize triterpenoids, which overturns the long-held belief that all triterpenoids are synthesized exclusively from squalene. The discovery and mining of enzymes in the synthesis of triterpenoids provides more candidates that can be employed in the microbial production.

Despite the obtain of novel efficient key enzymes, the modifications of known enzymes are alternative ways for efficient production. Protein engineering of the key enzymes has attracted much attention of the researchers. In recent years, rational design of enzymes involved in triterpenoid biosynthesis has become a trend by means of sequence alignment, molecular evolutionary analysis and protein homology modeling. Successful examples include RoSC2 (Y531/L256/L258) [28], UGT71G1 (Y202A) [109], CYP72A63 (T338S/W205A) [9], etc. Significant advances in protein engineering have accelerated the improvements in enzyme specificity, activity, and stability. Moreover, the protein engineering of key enzymes could lead to novel reactions that do not occur in nature and thus generate different types of specialized metabolites. The crystal structure of enzymes is the basis for the elucidation of enzyme-substrate binding and interactions, which helps to optimize the catalytic properties of enzymes through protein design. Even though the docking of enzyme-substrate active sites can be studied by indirect methods such as homology modelling and molecular simulation, the semi-rational design lacking crystallographic data is still challenging. Therefore, accelerating the acquisition of the three-dimensional structure of enzyme molecules is one of the main problems need to be solved in protein engineering [110].

In recent years, the technologies for obtaining the three-dimensional structure of enzymes mainly include nuclear magnetic resonance and X-ray crystallography. Padyana et al. [111] obtained the crystal structure of human squalene epoxide (SQLE), a key rate-limiting enzyme in cholesterol biosynthesis, through the most classical X-ray crystallography. The key conformational rearrangement necessary for the binding of inhibitors including NB-598 and compound-4 was determined, and the structural basis of catalytic epoxidation of SQLE was clarified. However, there are many difficulties in the crystallization process for some proteins, such as membrane proteins. Fortunately, the improvement of single-particle electron cryomicroscopy (Cryo-EM) resolution makes up for this deficiency. Yao et al. [112] performed high-resolution cryoEM structural analysis of AcrB reconstituted into liposomes by cryo-EM to obtain the membrane protein structure in physiological state. Gong et al. [113] measured the three-dimensional structure of human cholesterol intracellular transporter Niemann-pick C1 (NPC1) and its complex with EboV-GPCL by CRYO-EM. Qian et al. [114] analyzed the near-atomic resolution cryo-EM structure of the full-length human ABCA1 protein, which laid an important foundation for understanding the mechanism of ABCA1-mediated lipid export and the pathogenesis of related diseases. AlphaFold 2, a representative combination of artificial intelligence and biophysical ideas, will become an important tool for structural biology. It can provide an initial structural model for X-ray crystal diffraction, avoiding the embarrassment that the structural model cannot be established due to lack of phase information. More targeted mutation sites to stabilize protein structures can also be provided. AlphaFold 2 can also guide the classification of 2D images in single-particle cryo-EM structure analysis. Obviously, the development of structural biology will be driven by the combination of AlphaFold 2 and experimental structural biology.

Different from structure-guided protein engineering, the enzyme gene functional modification method based on Rosetta and the amino acid co-evolution information calculated by GREMLIN is suitable for enzyme molecules with rare crystal structures, which have been used to optimize high-resolution protein structures [115]. Ma et al. changed the function of cucurbitadienol synthase from cucumber by Rosetta, resulting in a conversion of catalytic activity from tetracyclic products to pentacyclic products [116]. Li et al. [117] transformed the multifunctional P450 enzyme CYP87D20 involved in the synthesis of cucurbitacin C into a mono-oxygenase, which has the function of directionally catalyzing cucurbitadienol to generate 11-hydroxyl cucurbitadienol, the precursor of mogrol. Zhang et al. performed protein surface engineering based on the CYP87D20 mutant, and combined with amino acid site mutations in the active region to further improve the catalytic activity of 11-hydroxycucurbitadienol synthase [45].

Protein engineering is expected to be developed as a useful approach for the commercial production of useful triterpenoids. This strategy also opens a new perspective for synthetic biology by constructing mutant libraries of key enzymes to generate triterpenoid syntheses with novel structures and prominent functions, as well as more and more functional triterpenoid compounds. In future, enzyme catalysts for any chemical reaction could be designed as machine learning, multi-site mutagenesis, high-throughput biosensors, and other technologies are more and more matured. The employment of protein engineering in microbial synthesis of plant triterpenoids will greatly accelerate their industrial application.
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Declaration of competing interest

The authors declare no competing interest.

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