Direct bioproduction of DHAA (dihydroartemisinic acid) rather than AA (artemisinic acid), as suggested by previous work would decrease the cost of semi-biosynthesis artemisinin by eliminating the step of initial hydrogenation of AA. The major challenge in microbial production of DHAA is how to efficiently manipulate consecutive key enzymes ADH1 (artemisinic alcohol dehydrogenase), DBR2 [artemisinic aldehyde Α11(13) reductase] and ALDH1 (aldehyde dehydrogenase) to redirect metabolic flux and elevate the ratio of DHAA to AA (artemisinic acid). Herein, DHAA biosynthesis was achieved in Saccharomyces cerevisiae by introducing a series of heterologous enzymes: ADS (amorpha-4,11-diene synthase), CYP71AV1 (amorphadiene oxidase), ADH1, DBR2 and ALDH1, obtaining initial DHAA/AA ratio at 2.53. The flux toward DHAA was enhanced by pairing fusion proteins DBR2-ADH1 and DBR2-ALDH1, leading to 1.75-fold increase in DHAA/AA ratio (to 6.97). Moreover, to promote the substrate preference of ALDH1 to dihydroartemisinic aldehyde (the intermediate for DHAA synthesis) over artemisinic aldehyde (the intermediate for AA synthesis), two rational engineering strategies, including downsizing the active pocket and enhancing the stability of enzyme/cofactor complex, were proposed to engineer ALDH1. It was found that the mutant H194R, which showed better stability of the enzyme/NAD+ complex, obtained the highest DHAA to AA ratio at 3.73 among all the mutations. Then the mutant H194R was incorporated into above rebuilt fusion proteins, resulting in the highest ratio of DHAA to AA (10.05). Subsequently, the highest DHAA reported titer of 1.70 g/L (DHAA/AA ratio of 9.84) was achieved through 5 L bioreactor fermentation. The study highlights the synergy of metabolic engineering and protein engineering in metabolic flux redirection to get the most efficient product to the chemical process, and simplified downstream conversion process.

**Keywords:** dihydroartemisinic acid, artemisinic acid, protein engineering, synthetic biology, Saccharomyces cerevisiae
INTRODUCTION

Artemisinin, a well-known sesquiterpene lactone isolated from extracts of *Artemisia annua* with excellent anti-malaria properties, had been designated as first-line antimalarial drugs by WHO in 2002 (Paddon and Keasling, 2014). Artemisinin production and prices vary greatly as they depend on plant extraction, which relies on the supply of plant materials (Chunyin and Cook, 2012; Peplow, 2013). Therefore, a stable and sustainable supply of artemisinin is highly desirable. This breakthrough was achieved by the Amyris, Inc., in which a semi-synthetic process of artemisinin production was developed (Westfall et al., 2012; Paddon et al., 2013). The semi-synthesis of artemisinin consists of two parts: (1) *de novo* biosynthesis of AA (artemisinic acid) with a very high titer (25 g/L) by an engineered *Saccharomyces cerevisiae* and (2) extract AA from yeast culture and transform it to artemisinin by chemical process. To be noted, the first step of the chemical process was reduction of AA to dihydroartemisinic acid (DHAA). Although the biosynthesis of DHAA has been demonstrated (Zhang et al., 2008), the productivity (about 15.7 mg/L) has not yet to be optimized in a heterologous host. Although hydrogenation conversion of AA to DHAA was efficient (conversion efficiency 99%), the production of unexpected isoform (R, S)-DHAA (about 6%) was not avoided (Paddon et al., 2013; Turconi et al., 2014; Pieber et al., 2015). Instead, biosynthesis of DHAA produced very little (R, S) isomer in yeast (Zhang et al., 2008). In order to execute a more efficient way of synthesizing artemisinin, engineering DHAA biosynthesis in microbes would open up a promising alternative route.

The biosynthesis pathways of DHAA and AA both start from amorpha-4,11-diene that is oxidized to AO (artemisinic aldehyde) through AOH (artemisinic alcohol) by CYP17AV1 (amorphadiene oxidase) and ADH1 (artemisinic alcohol dehydrogenase) (Figure 1). AO is a joint intermediate followed by two branch biosynthesis pathways: (1) directly oxidized to AA by ALDH1 (aldehyde dehydrogenase) or (2) reduced to dihydroartemisinic aldehyde (DHAO) by DBR2 (artemisinic aldehyde Δ11(13) reductase) and then oxidized to DHAA by ALDH1 (Chen et al., 2017). The enzymatic parameters of DBR2 and ALDH1 to their substrates AO and DHAA were summarized in Supplementary Table S1 (Zhang et al., 2008; 2009). Because of the slightly higher reported affinity of ALDH1 [For substrate AO, Km(DBR2) = 19 µM, Km(ALDH1) = 2.58 µM], there is no advantage for DBR2 to bind with AO. Furthermore, no enzymes (even DBR2) have been reported to directly catalyze AA to DHAA. Thus, it was inevitable to produce a large amount of AA as by-product of DHAA production. So far, the highest ratio of DHAA/AA reaches just 1.67 achieved by enzymatic reaction *in vitro* (Chen et al., 2017). Therefore, the major challenge in production of DHAA by recombinant yeasts is efficient redirection of carbon flux to DHAA rather than AA.

Branch-point regulatory mechanisms are involved in many natural metabolic pathways (such as TCA cycle) (Chen et al., 2017). In these pathways, the enzymes can work together spatiotemporally due to protein–protein interactions and channel the metabolites between sequential enzymes without equilibration in the aqueous phase inside cells (Sweetlove and Fernie, 2018). The assemblies of consecutive enzymes are formed either from large clusters of multiple copies of enzymes, or by pairwise interactions of enzymes from single complexes which are beneficial for enzymes to reach substrate saturation (Zhang, 2011) so that the reaction flux is regulated at a branch point. Pham et al. (2015) once co-localized enzymes responsible for GABA (Gamma-aminobutyric acid) biosynthesis together to switch the metabolic flux toward GABA from TCA cycle and finally increased the production of GABA by 2.7 fold.

**FIGURE 1** | Schematic diagram of tailoring the ratio of DHAA to AA by protein fusion and modifying ALDH1 catalysis preference. The broader blue arrow indicates enhanced flux toward DHAA. AOH, artemisinic alcohol; AO, artemisinic aldehyde; DHAO, dihydroartemisinic aldehyde; ADH1, artemisinic alcohol dehydrogenase of *A. annua*; DBR2, artemisinic aldehyde Δ11(13) reductase of *Artemisia annua*; ALDH1, artemisinic aldehyde dehydrogenase of *A. annua*.

DHAA/AA=2.53:1

Redirect the metabolic flux

DHAA/AA=10.05:1
Refraining to DHAA biosynthetic pathway itself and key enzymes, two obstacles should be overcome to achieve a high ratio of DHAA/AA and a high DHAA titer: (1) redirecting the metabolic flux from AA toward DHAA via the expression of DBR2 requires the assembly of the pathway enzyme in a desired order and promotes the reactions of metabolites along a specified pathway (Miles et al., 1999); (2) rationally engineering ALDH1 to shift the substrate specificity from AO to DHAO.

Herein, the biosynthesis pathway of DHAA was successfully rebuilt in S. cerevisiae with high FPP supply. In order to improve the ratio of DHAA/AA by increasing the substrate accessibility of AO by DBR2 as well as that of DHAO by ALDH1, fusion proteins of paired enzymes (ADH1-DBR2 or DBR2-ALDH1) were adopted to reorganize the biosynthetic pathway of DHAA (Figure 1). Meanwhile, ALDH1, as the joint enzyme for biosynthesis of DHAA and AA, was also rationally engineered to shift the substrate specificity from AO to DHAO. Correspondingly, the ratio of DHAA/AA was enhanced by 3.34 fold (from 2.53 to 10.05) through integrating these above two strategies, without the compromise of DHAA production. Eventually, the highest DHAA titer of 1.70 g/L (DHAA/AA ratio of 9.84) was achieved in a 5 L bioreactor through high density fermentation. The study highlights the importance of redirecting metabolic flux toward a desired target via consecutive enzyme-enabled reorganization.

**MATERIALS AND METHODS**

**Strains and Medium**

All *E. coli* used for plasmid construction were cultured at 37 °C in Luria-Bertani (LB) medium (1% tryptone, 0.5% yeast extract, and 1% NaCl) with 100 µg/ml ampicillin or 34 µg/ml chloramphenicol if necessary.

All engineered yeast strains were derived from *S. cerevisiae* CEN.PK2-1C (Entian and Kötter, 2007) obtained from EUROSCARF (Frankfurt, Germany) and were listed in Table 1. *S. cerevisiae* strains were cultured in YPD medium (2% tryptone, 1% yeast extract, and 2% glucose) or in synthetic formulations for yeast culture are available in our previous work (Su et al., 2015).

**Plasmid Construction**

Plasmids used in this study are listed in Table 2. The genes *ADS* (ACCESSION Q9AR04), *CYP71AV1* (ACCESSION Q1PS23), *DBR2* (ACCESSION KC505370.1), *ALDH1* (ACCESSION JQ609276.1), *ADH1* (ACCESSION JF910157.1), *CYB5* (ACCESSION JQ582841.1), *CPR1* (ACCESSION DQ318192.1) from *Artemisia annua* were codon-optimized for expression in yeast and synthesized by GenScript, Inc. (China). To overexpress three key genes for DHAA biosynthesis (including *ADS*, *CYP71AV1*, and *DBR2*), plasmid pZBX040 were first constructed based on the multi-copy plasmid pRS425. Then the plasmid with fusion protein *DBR2-ADH1* or *ADH1-DBR2* (including pZBX059, pZBX060, pZBX067, pZBX069) were constructed based on pZBX040. To substitute ALDH1 integrated in the genome with fusion protein *DBR2-ALDH1*, the cassette *HIS3*-T*HIS3*-P*GAL7*-DBR2-Linker-ALDH1-T*DH1* were assembled in pSB1C3 (obtained from the Registry of Standard Biological Parts1) to form plasmid pZBX100, pZBX101. The cassettes with mutated ALDH1 were constructed based on pZBX101. All the primers used in this work are listed in Supplementary Table S2.

**Construction of pZBX040**

The DNA fragments *ADS*, *CYP71AV1*, P*GAL10*, T*PGK1*, and T*ADH1* were amplified by PCR and joined together by overlap extension PCR (OE-PCR) to obtain cassette T*ADH1*(RC)-CYP71AV1(RC)-P*GAL10*(RC)-P*GAL1*-ADS-T*PGK1*. The cassette was digested with BamHI and XhoI and inserted into pRS425 to obtain plasmid pZBX020. Fragments *DBR2*, P*GAL7*, T*CYC1* were amplified by PCR and joined together by OE-PCR to obtain

**TABLE 1 | Yeast strains used in this study.**

| Yeast strains | Description | Source |
|---------------|-------------|--------|
| CEN.PK2.1C    | MAT a; ura3-52, trp1-289, leu2-3,112, his3Δ1, MAL2-8C, SUC2 | Invitrogen |
| Sc027         | CEN.PK2-1C derivatives; leu2-3,112:G418°_P*GAL7*-CYB5_T*ERG19*(RC)-ERG19*(RC)-P*GAL1*(RC)-P*GAL8*-ERG18; his3Δ1; HIS3_P*GAL7*-ADH1_T*DH1*-P*GAL7*-ERG12*(RC)-ERG12*(RC)-P*GAL1*(RC)-P*GAL7*-ERG10-ERG10-ERG10; ade1Δ_T*HMG1*(RC)-HMG1*(RC)-P*GAL1*(RC)-P*GAL7*-T*DH1*-ADE1; ura3-52:T*ERG13*(RC)-HMG1*(RC)-P*GAL1 (RC)_P*GAL10*-ERG13_T*ERG13; trp1-289; T*ERG13*(RC)-HMG1*(RC)-P*GAL1 (RC)_P*GAL10*-ERG20_T*ERG20_TRP1; gal1/10/7Δmatα_P*GAL7*-CPR1-T*CYC1 | This study |
| Sc057         | Sc027 derivative; GAL80ΔURA3_P*GAL7*-AdaDH1_T*CYC1 | This study |
| Sc077         | Sc027 derivative; GAL80ΔURA3 | This study |
| Sc085         | Sc057 derivative; pZBX040 | This study |
| Sc113         | Sc077 derivative; pZBX059 | This study |
| Sc115         | Sc077 derivative; pZBX067 | This study |
| Sc146         | Sc077 derivative; pZBX060 | This study |
| Sc147         | Sc077 derivative; pZBX069 | This study |
| Sc352         | Sc057 derivative; his3Δalcohol1Δ_hphA | This study |
| Sc361         | Sc077 derivative; pZBX067; ΔALDH1_P*GAL7*-DBR2-Linker1-ALDH1_T*CYC1 | This study |
| Sc467         | Sc057 derivative; pZBX040; ΔALDH1_P*GAL7*-DBR2-Linker1-ALDH1_T*CYC1 | This study |
| Sc468         | Sc057 derivative; pZBX040; ΔALDH1_P*GAL7*-DBR2-Linker1-ALDH1_T*CYC1 | This study |
| Sc470         | Sc085 derivative; ura3Δdown:hphMX6_P*GAL7*-DBR2_T*CYC1 | This study |
| Sc479         | Sc057 derivative; pZBX040; ΔALDH1_P*GAL7*-ALDH1_H134R_T*CYC1 | This study |
| Sc457         | Sc057 derivative; pZBX067; ΔALDH1_P*GAL7*-DBR2-Linker1-ALDH1_H134R_T*CYC1 | This study |

The DNA fragment followed by "*(RC)*" represents that the orientation of the DNA fragment is reversed. Sequences of linker1 and linker2 showed in Figure S3. In strain Sc470, the cassette pphMX6-P*GAL7*-DBR2-T*CYC1 was inserted into the downstream of cassette T*ADH1*(RC)-HMC1*(RC)-P*GAL1*(RC)-P*GAL7*-ERG13_T*ERG13 which had been integrated in the locus in ura3.

1http://parts.igem.org/Part:pSB1C3
TABLE 2 | Plasmids used in this study.

| Plasmid | Description | Source |
|---------|-------------|--------|
| pZBX020 | pRS425_TADH1(ADH1)-CYP71AV1(ADH1)-PGAL10(ADH1)-P_GAL1-ADS_TDXH | This study |
| pZBX040 | pRS425_PGAL1-DBR2_TADH1(CYC1)-PGAL1(ADH1)-CYP71AV1(ADH1)-PGAL10(ADH1)-ADS_TDXH1 | This study |
| pZBX059 | pRS425_PGAL1-ADH1 linker1-DBR2_TADH1(ADH1)-CYP71AV1(ADH1)-PGAL10(ADH1)-ADS_TDXH1 | This study |
| pZBX060 | pRS425_PGAL1-ADH1 linker2-DBR2_TADH1(ADH1)-CYP71AV1(ADH1)-PGAL10(ADH1)-ADS_TDXH1 | This study |
| pZBX067 | pRS425_PGAL1-DBR2 linker1-ADH1_TADH1(ADH1)-CYP71AV1(ADH1)-PGAL10(ADH1)-ADS_TDXH1 | This study |
| pZBX069 | pRS425_PGAL1-DBR2 linker2-ADH1_TADH1(ADH1)-CYP71AV1(ADH1)-PGAL10(ADH1)-ADS_TDXH1 | This study |
| pZBX101 | pSB1C3_HIS3-TDRH1-DBR2 linker1-ADH1_TDXH | This study |
| pZBX100 | pSB1C3_HIS3-TDRH1-DBR2 linker2-ADH1_TDXH | This study |
| pZBX199 | pSB1C3_HIS3-TDRH1-ALDH1-H194R_TDXH | This study |
| pZBX196 | pSB1C3_HIS3-TDRH1-ALDH1-227V_TDXH | This study |
| pZBX197 | pSB1C3_HIS3-TDRH1-ALDH1-227F_TDXH | This study |
| pZBX194 | pSB1C3_HIS3-TDRH1-ALDH1-227V_TDXH | This study |
| pZBX195 | pSB1C3_HIS3-TDRH1-ALDH1-227F_TDXH | This study |
| pZBX218 | pSB1C3_HIS3-TDRH1-DBR2 linker1-ADH1-H194R_TDXH | This study |

Construction of Plasmid for Fusion Protein DBR2-ALDH1

Fragment HIS3-P_GAL17 was amplified by PCR from the genome of Sc085 followed by digestion of EcoRI and BsaI. The fragments of DBR2, and ALDH1-TDXH1 were amplified by PCR from Sc085. Fragment DBR2 was digested with BsaI, and fragment ALDH1-TDXH1 was digested with BsaI and PstI. The vector pSB1C3 was digested with EcoRI, and BsaI. All above four fragments were ligated together by T4 ligase to construct all the plasmid with the fusion protein DBR2-ALDH1 (including pZBX099, pZBX100, pZBX101). The cassettes of the plasmids were released from plasmid by digestion of PmeI before transformation to the strain Sc352 for integration.

Construction of Plasmid for ALDH1 Mutants

Each DNA fragment of an ALDH1 mutant was divided into two parts P_GAL17-ALDH1a and ALDH1b with 40 bp overlapping according to the mutated site, and was amplified by PCR. Each pair of P_GAL17-ALDH1a and ALDH1b was ligated to form intact ALDH1 mutant by OE-PCR followed by digestion of SpeI and KpnI. The vector pZBX100 was digested with SpeI, and KpnI for insertion of the fragment of P_GAL17-ALDH1a mutant, obtaining pZBX194, pZBX195, pZBX196, pZBX197, and pZBX199.

The process of construction of pZBX218 was the same as that of pZBX101 except that the fragment ALDH1-TDXH1 was amplified from plasmid pZBX199. All the cassettes of the plasmid described here were released from plasmid by digestion of PmeI and transformed to the strain Sc352.

Fermentation Condition

Medium for Fermentation

All the fermentation medium (FM) recipe used in this work was prepared similarly to that used in Westfall's work with some modifications (Westfall et al., 2012). The medium was composed of 8 g/L KH2PO4, 15 g/L (NH4)2SO4, 6.2 g/L MgSO4.7H2O, 40 g/L glucose, 12 ml/L vitamin solution, and 10 ml/L trace metal solution. The vitamin solution included 550 mg/L biotin, 1 g/L calcium pantothenate, 1 g/L nicotinic acid, 25 g/L myo-inositol, 1 g/L thiamine HCl, 1 g/L pyridoxal HCl, 0.2 g/L p-aminobenzoic acid, and 2 g/L adenine sulfate. The trace metal solution was composed of 5.75 g/L ZnSO4.7H2O, 0.32 g/L MnCl2.4H2O, 0.32 g/L Anhydrous CuSO4, 0.47 g/L CoCl2·6H2O, 0.08 g/L Na2MoO4·2H2O, 2.9 g/L CaCl2·2H2O, 2.8 g/L FeSO4·7H2O, and 80 ml/L EDTA solution (containing 0.5 mol/L Na2EDTA pH 8.0). The amino acid solution (10 ml/L), including 2 g/L methionine, 6 g/L tryptophan, 8 g/L isoleucine, 5 g/L phenylalanine, 10 g/L sodium glutamate, 20 g/L threonine, 10 g/L aspartate, 15 g/L valine, 40 g/L serine and 2 g/L arginine, was supplemented to the medium in our study for obtaining better cell growth use. The FM bases (8 g/L KH2PO4, 15 g/L (NH4)2SO4, and 6.2 g/L MgSO4.7H2O) and glucose stock solution (667 g/L) were sterilized using an autoclave. Vitamin solution, trace metal solution, and amino acid solution were sterilized by filtration. All the components of the FM were mixed together after sterilization. Glucose stock solution (667 g/L), filtrated 95% (v/v) ethanol solution and filtrated feed stock solution (80 g/L KH2PO4, 150 g/L (NH4)2SO4,
containing 25 ml FM medium at initial OD600 of 0.05 and grown for another 16 h cultivation at 30°C. The seed culture was transferred into 250 ml flask containing 25 ml FM medium at initial OD600 of 0.2. The cells were grown at 30°C with shaking at 200 rpm. After 24 h, 5 ml IPM (Isopropyl myristate) and 20 g/L ethanol was added to each flask. The whole fermentation process continued for 120 h until harvest.

5 L Bioreactor Fermentation
Seed culture preparation was the same as in-flask fermentation. The seed culture (200 mL) was inoculated into a 5 L bioreactor containing a 2 L batch FM medium with 20 g/L glucose. The pH was controlled at 5 by adding 5 mol/L NaOH. The gas flow and the temperature were maintained at 1.5 vvm and 30°C. During the fermentation, glucose solution was fed into the bioreactor at a speed of 0.3 mL/min after the glucose was depleted at the running time of about 6 h. During the glucose feeding stage, the dissolved oxygen level (DO) was kept at 40% by cascading agitation from 400 to 600 rpm. When OD600 increased to 50 (about 30–36 h), the feeding of glucose was switched to ethanol feeding. The concentration of ethanol was maintained at 10 g/L to guarantee no starvation and no excess carbon accumulation. During the ethanol feeding process, DO was automatically maintained at about 30% by cascading stirring.

The Measurement of DHAA and AA and Other Intermediates
After harvest, the fermentation broth was centrifuged at 12,000 g for 2 min and the IPM phase was collected. And then 50 μL organic phase was mixed with 950 μL methanol. After filtrated with 0.22 μm Nylon66 filter, the sample was ready for HPLC analysis. A 10 μL aliquot was injected into waters e2695 HPLC with ultraviolet detection at 194 nm. A thermoHypersil BDS C18 column (4.6 mm × 150 mm × 5 μm) was used for separation, with the following gradient (channel A: acetonitrile, channel B: water plus 0.1% formic acid): 0–3 min 65% A, gradually increased to 100% A from 3 to 10 min, held at 100% A from 10 to 13 min, decreased to 65% A from 13 to 15 min, kept 65% A from 15 to 18 min. The column was held at 25°C during the separation. Under this condition, DHAA and AA were found to elute at 6.67 and 7.43 min, respectively (see Figure 2B). The concentrations of both products in the samples were calculated using the calibration curves of standards (HPLC ≥ 98%) which was purchased from Chengdu Pufe Di Biotech, Co., Ltd. Standards of DHAO, AO, AOH, and AD were purchased from TRC (Toronto Research Chemicals).

Homology Modeling, Molecular Docking, and Structural Analysis
Three-dimensional structure models of ALDH1 were constructed using the program Swiss-Model2 to get the structure information accordingly. The structures were modeled using the high sequence homology (~41% identified) and high resolution crystal structure of the aldehyde dehydrogenase family protein (indole-3-acetaldehyde dehydrogenase) combined with NAD+ from tomato as the template (pdb id:5iuw-A). The structure models were subjected to energy minimization using the Swiss-Pdb Viewer. Afterward the docking of enzyme and ligand were performed using the AutoDockVina program (Oleg and Olson, 2010). The docking cluster analysis was performed in the AutoDockVina program environment, and clusters were characterized by binding energy (in kilocalories per mole). Establishment of dative bonds between ligand, NAD+ and the corresponding amino acids were followed by energy minimization. The built complex structural analysis was done using Pymol software (Delano, 2010). The mutation at the specific amino acid site was also introduced using this software, which allowed exploration of the spatial and molecular interactions among amino acids.

RESULTS AND DISCUSSION
Strain Construction for DHAA Production
Referring to previous work (Westfall et al., 2012), CEN.PK2.1C was selected as the host strain for DHAA production. In order to decouple cell growth and product production, inducible promotor GAL (Pgal1, Pgal7, Pgal10) was explored to control all the overexpressed genes (Peng et al., 2017). GAL80 was deleted to eliminate the demand of galactose for de-repressing the promotor GAL. To increase the metabolic flux toward FPP, all the genes of MVA pathway (Westfall et al., 2012) were overexpressed in the genome. Among these genes, tHMG1 was integrated into the genome with three copies (Supplementary Figure S1). Three key heterologous genes for DHAA biosynthesis including ADS, CYP71AV1 and DBR2 were constructed in a multi-copy plasmid pRS425 for higher level expression. Meanwhile, four other heterologous genes, including ADH1, ALDH1, CYB5 and CPR1, were integrated in the genome at one copy to enhance the metabolic flux to DHAA (Figure 2A). The strain for DHAA production (Sc085) was characterized in shake flask fermentation and the products were detected by HPLC at a wavelength of 194 nm. The results showed the presence of DHAA and AA, indicated by the observation that the retention time of DHAA and AA peaks was same as the mix standard (6.67 min for DHAA, 7.43 min for AA) (Figure 2B). And 327 mg/L of DHAA and 129 mg/L of AA were successfully detected. The DHAA/AA ratio of 2.53 was much higher than the reported highest ratio (DHAA/AA = 1.67) (Figure 2C) (Chen et al., 2017).

2http://swissmodel.expasy.org/
Nevertheless, there was still about 30% of the metabolic flux flowing toward AA (129 mg/L), probably indicating increasing expression level of DBR2 was needed to enhance the transformation from AO to DHAO. However, because of the reportedly slightly higher affinity of ALDH1 to the joint intermediate AO [Km(ALDH1) = 2.58 μM vs. Km(DBR2) = 19 μM] (Zhang et al., 2008; Teoh et al., 2009), individual over-expression of DBR2 might not be sufficient to tackle this issue. When Sc470 held another copy of DBR2 within the chromosome, the DHAA/AA ratio was reduced as well as the DHAO titer and no AOH, AO or DHAO was accumulated (Figure 3 and Supplementary Figure S4). To summarize, it was difficult to directly control the reaction route toward DHAA rather than AA by simply increasing the expression of these genes. Considering that ALDH1 is a promiscuous enzyme that can catalyze AO and DHAO simultaneously, we need to reconfigure the enzymes (ADH1, DBR2, and ALDH1) in a desired order and switch the catalysis preference of ALDH1 to tailor the ratio of DHAA/AA.

Switch of the Biosynthesis Pathway to DHAA by Fusion Proteins

To increase the substrate accessibility of AO by DBR2, the biosynthetic route from AOH to DHAO was re-built via protein fusion of ADH1 and DBR2 (Figure 1). As known, linker type

![Figure 2](image-url)
the fusion proteins DBR2-ADH1 and DBR2-ALDH1 were co-expressed to form the strain Sc361 which resulted in a lower production of AA with no concomitant increase in DHAA (323.2 mg/L) and raised the DHAA/AA to 6.76 (Figure 3).

Since there is a high accumulation of AD (about 1.05–1.20 g/L) and no accumulation of any other intermediates (including AO, DHAO, AOH) in all strains described above (Figure 3 and Supplementary Figures S2, S3), the oxidation of AD was also the final step of biosynthesis pathway for all strains, and AOH could be converted to the end product AA or DHAA in time. Additionally, if comparing the effect of the fusion proteins DBR2-ADH1 and DBR2-ALDH1 on the DHAA/AA ratio, DBR2-ALDH1 exhibited a more important role in tailoring DHAA/AA ratio (from 2.53 to 4.58), while the other only increased DHAA/AA ratio to 2.95. Although the sum of the product was reduced, combining two fusion proteins would slightly increase the conversion yield of DHAA transformed from AD [proportion of DHAA (DHAA%) increased from 19.7 to 22.8%]. All of the above data demonstrated that engineering minimal conversion of AO to AA is crucial to increase the DHAA/AA ratio and slightly increase the DHAA%. Thus, in order to get optimal ratio of DHAA to AA, reducing the catalysis preference of the shared enzyme ALDH1 to AO or enhancing the catalysis preference to that of DHAO to smooth flux toward DHAA seems to be especially important, in addition to protein fusion.

### Improving the Ratio of DHAA/AA by Changing ALDH1 Specificity

ALDH1 can employ AO or DHAO as the substrate and may have a slightly stronger binding affinity to AO, which will potentially restrain the production of DHAA (Supplementary Table S1). Herein, we attempted to change the substrate specificity of ALDH1 by rationally modifying its structure to improve the preference to DHAO.

Resorting to the molecular structure, DHAO and AO structures are very similar. They both have the aldehyde group which will then be oxidized to the carboxyl group. Nevertheless, the AO molecule has an additional vinyl group adjacent to the aldehyde group (Supplementary Figure S5) that forms a conjugated structure. Such conjugated structure in AO will affect the electronic cloud distribution of the catalyzed aldehyde group. Therefore, the structure difference of the catalyzed site of substrates would cause distinct binding force of the shared ALDH1 to DHAO or to AO. By analyzing the complex structure of ALDH1 with its cofactor NAD$^+$ and substrates, it was found that the nicotinamide group of NAD$^+$ just bound to the catalytic site of substrates in the active pocket of ALDH1 (Supplementary Figure S5). Consequently, a hypothesis was proposed that the binding characteristic of NAD$^+$ to DHAO or to AO potentially made the crucial contributions to the differences in ALDH1’s affinities to its substrates. With this regard, the structural modifications of ALDH1 were carried out in two aspects. Firstly, the binding pocket, which for both NAD$^+$ and substrate, was downsized to shorten the distance between the nicotinamide group of NAD$^+$ and the catalyzed aldehyde group of DHAO. Secondly, the binding force of ALDH1 and NAD$^+$...
was enhanced to improve the complex structural stability and enhance the enzymatic catalytic activity to DHAO, since NAD$^+$ is loosely bound to ALDH1 (Butterworth, 2010).

Accordingly, an attempt was made to replace the residues G223 and G227 with branch chains of a larger size (like V or F) by squeezing them through the space within the binding pocket. However, by further comparing the complex structures between the wild type ALDH1/NAD$^+$ and mutant (G223V/F, G227V/F)/NAD$^+$, it was found that although the binding pocket structure of the mutant complex was downsized, the adenine group of NAD$^+$ was extruded out of the original position, due to the steric effect of large branch chains of the mutated residues (Supplementary Figure S6). This structure weakened the binding force between NAD$^+$ and ALDH1 mutants (Supplementary Figure S6), probably damaging or even destroying the enzymatic activity of ALDH1 mutants. As expected, either the ratio of DHAA/AA or the DHAA titer were dramatically decreased by mutagenesis of G223F and G227V/F (Supplementary Figure S6F), compared to that of the wild type ALDH1. Although G223V increased the ratio of DHAA/AA, the actual production of DHAA was reduced significantly (Supplementary Figure S6F).

The significant accumulation of DHAO was found in all mutants at G223 and G227, showing that the activity of these ALDH1 mutants were reduced. Although CYP71AV1 was reported to transform AO to AA (Teoh et al., 2009), its activity might not be strong enough to produce more AA or DHAA.

Alternatively, we sought to enhance the binding between ALDH1, NAD$^+$ and the substrate. The site-directed mutant H194R was constructed to enhance the binding of enzyme to NAD$^+$. Comparing with the complex structures of wild type ALDH1/NAD$^+$, the mutant H194R formed an additional electrostatic interaction with the phosphate group of NAD$^+$ (Figure 4A). Such electrostatic interactions would improve the structural stability of the mutant complex, which would significantly regulate the enzymatic catalysis activity to substrate (AO and DHAO).

Consequently, H194R significantly increased the ratio of DHAA/AA to 3.73 without reducing the production of DHAA (Figure 4). Different from other mutants in G223 or G227, the strain with mutant H194R (Sc429) didn’t accumulate DHAO and any other intermediates (AO, AOH) in the product. H194R seemed to affect the activity of ALDH1 with AO and reduced the production of AA (see Supplementary Figure S6F). In summary, our experimental results were well-consistent with our modeling based on structural analysis (Figure 4).

**DHAA Enrichment by 5 L Fed-Batch Fermentation**

Mutated ALDH1(H194R) was introduced into strain Sc361 to replace the wild type of ALDH1 existing in fusion protein DBR2-ALDH1, obtaining strain Sc457. As shown in Figure 5A, by further reducing the production of AA, the highest ratio of DHAA/AA at 10.05 was achieved at shake flask level. The DHAA titer was maintained at 311 mg/L without accumulation of AOH, AO, or DHAO (Figure 5 and Supplementary Figure S4). Taking the DHAA/AA ratio and the DHAA titer together into considerations, strain Sc457 was chosen for the following fermentation process.

In order to promote DHAA production and investigate the performance of strain Sc457 in fermenters, 5 L bioreactor fermentation was conducted using the carbon restriction strategy. As shown in Figure 5B, OD$_{600}$ increased sharply before 32 h when glucose was utilized as a carbon source. When the carbon source was switched to ethanol, the OD$_{600}$ was gradually promoted to 142 until harvest. Under above control process, the highest reported DHAA titer at 1.70 g/L in microbes was accomplished after 132 h fermentation (Figure 5B). A slight accumulation of AA was also obtained. It was also observed that the average ratio of DHAA to AA during the fermentation process was quite steady, demonstrating that the genetic stability of strain Sc457 was quite stable and the process was scalable to mimic the strain performance in shake flask to some extent. As the current DHAA production is not as high as that of AA in Amyris, Inc. (Paddon et al., 2013), further optimization of the fermentation efficiency via integrating medium and feed strategy (Westfält et al., 2012) as well as off-gas analysis feedback control could be tried.
During the fermentation process, a significant accumulation of glycerol was also found at about 5 g/L in glucose consumption stage. Ho and colleagues reported both UBR2 and GUT1 with single point mutation could be regarded as targets for establishing glycerol utilization in strains of the CEN.PK family (Ho et al., 2017). It was also found that regulatory and metabolic trade-offs of glycerol utilization in *S. cerevisiae* were revealed by laboratory evolution, which could guide us to enhance glycerol consumption rationally in our study (Strucko et al., 2018). Furthermore, no acetate was detected during the whole process due to carbon restriction strategy (data not shown).

**CONCLUSION**

The combination of metabolic engineering and protein engineering has shown great performances in constituting microbial cell factories for the production of natural products with complex structures. In this study, fusion proteins and modifying ALDH1 catalysis preference strategies have been adopted and conducted to successfully tailor the ratio of DHAA/AA in *S. cerevisiae*. Promoted flux toward DHAA via ADH1, DBR2 and ALDH1 was firstly reconstituted by pairing fusion proteins DBR2-ADH1 and DBR2-ALDH1. The theoretical model of enhancing the stability of the enzyme/cofactor complex was assumed and executed to switch the catalysis preference of ALDH1 toward DHAA. Consequently, the ratio of DHAA/AA was elevated from 2.53 to 10.05 with the highest DHAA titer reaching 1.70 g/L (DHAA/AA ratio of 9.84) in 5 L bioreactor fermentation. This study shows the potential of oriented arrangement of consecutive heterologous enzymes in the reconstitution of microbial cell factories. On the other hand, the reduced sum flux of the biosynthesis pathway might be enhanced by other strategies of metabolic engineering which will be studied in future work.
DATA AVAILABILITY STATEMENT

The datasets generated for this study can be found in the ADS (ACCESSION Q9AR04), CYP71AV1 (ACCESSION Q1PS23), DBR2 (ACCESSION KC50370.1), ALDH1 (ACCESSION JQ609276.1), ADH1 (ACCESSION JF910157.1), CBY5 (ACCESSION JQ582481.1), and CPR1 (ACCESSION DQ318192.1).

AUTHOR CONTRIBUTIONS

B-XZ and W-HX conceived of the study and participated in fed-batch fermentation. B-XZ and YW participated in strain construction and carried out the molecular genetic studies. M-DY carried out the protein analysis. M-DY, YW, and W-HX participated in design and coordination of the study as well as helped to draft the manuscript. W-HX supervised the whole research and revised the manuscript. All the authors read and approved the final manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fbioe.2020.00152/full#supplementary-material

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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