Expressing a Cytosolic Pyruvate Dehydrogenase Complex to Increase Free Fatty Acid Production in Saccharomyces Cerevisiae

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Research

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Abstract

Saccharomyces cerevisiae is being exploited as a cell factory to produce fatty acids and their derivatives as biofuels. Previous studies found that both precursor supply and fatty acid metabolism deregulation are essential for enhanced fatty acid synthesis. A bacterial pyruvate dehydrogenase (PDH) complex expressed in the yeast cytosol was reported to enable production of cytosolic acetyl-CoA with lower energy cost and no toxic intermediate.

Results

Overexpression of the PDH complex significantly increased cell growth, ethanol consumption and reduced glycerol accumulation. Furthermore, to optimize the redox imbalance in production of fatty acids from glucose, two endogenous NAD⁺-dependent glycerol-3-phosphate dehydrogenases were deleted, and a heterologous NADP⁺-dependent glyceraldehyde-3-phosphate dehydrogenase was introduced. The final strain with engineering of precursor and co-factor metabolism could produce 840.5 mg/L free fatty acid in shake flask, which was 83.2% higher than the control strain. Profile analysis of free fatty acid suggested the cytosolic PDH complex mainly resulted in the increases of unsaturated fatty acids (C16:1 and C18:1).

Conclusions

We demonstrated that cytosolic PDH pathway enabled more efficient acetyl-CoA provision with the lower ATP cost, and improved FFA production. Together with engineering of the redox factor rebalance, the cytosolic PDH pathway could achieve high level of FFA production at similar levels of other best acetyl-CoA producing pathways.

Background

Bioproduction of fatty acids and their derivatives serves as a sustainable route to alleviate the dilemma between the growing demands of high-energy-density fuels and concerns on their environmental impacts. However, fatty acids are involved in several essential cellular processes, such as membrane synthesis, energy storage, and protein modification, and their levels are therefore tightly regulated to ensure in vivo homeostasis [1]. Intensive efforts have been carried out to improve the titer, rate and yield (TRY) of fatty acid-derived biofuels, focusing mainly on the deregulation of fatty acid metabolism and redirection of carbon flux [2–4].

Regarding deregulation of fatty acid metabolism, deletions of dominant acyl-CoA synthetases (FAA1, FAA4), and fatty acyl-CoA oxidase (POX1) catalyzing the first step in β-oxidation, could significantly
accumulate free fatty acids (FFAs) [5–7]. Moreover, deletions of main sterol ester formation genes (ARE1, ARE2), and phosphatidate phosphatase genes (PAH1, DPP1, LPP1) could also deregulate fatty acid synthesis and increase the production of total fatty acids [8]. Combining deletion of several lipid regulation mechanisms resulted in a reduced complexity of the fatty acid metabolic network, leading to enhanced FFA accumulation [8, 9].

Besides engineering lipid metabolic networks, engineering of precursor supply and redox cofactor regeneration was also essential for carbon flux redirection towards FFA production. Each elongation cycle in fatty acid synthesis extends the backbone with a C2 unit and requires 2 molecules of NADPH as cofactors. Endogenous synthesis of cytosolic acetyl-CoA, the main C2 metabolite, is often limited and several heterologous enzymes have therefore been expressed to increase acetyl-CoA supply for FFA synthesis, including NAD\(^{+}\)-dependent alcohol dehydrogenase (AdhE), NAD\(^{+}\)-dependent acylating acetaldehyde dehydrogenase (EutE) [10], ATP citrate lyases (ACLs) [6, 7, 11–13], xylulose-5-phosphate phosphoketolase (XpkA) and phosphotransacetylase (Pta) [14]. Compared to these strategies, expression of cytosolic pyruvate dehydrogenase (PDH) is attractive because of its lower energy cost [15]. The PDH complex consists of three catalytic subunits, including pyruvate dehydrogenase (E1), dihydrolipoyl transacetylase (E2), and dihydrolipoyl dehydrogenase (E3), and requires four kinds of cofactors TPP, lipoate, FAD and NAD\(^{+}\). Among three PDH complexes that have been expressed in the cytosol of yeast, PDH from Enterococcus faecalis was demonstrated to enable complete replacement of the native pathway for cytosolic acetyl-CoA synthesis, and was less sensitive to high NADH/NAD\(^{+}\) ratios compared with other PDH complexes [16–18]. However, one concern with PDH expression for FFA production is the redox imbalance between redox couples of NAD\(^{+}/\)NADH and NADP\(^{+}/\)NADPH.

Previous studies showed that high glycerol production was often observed in FFA production strains, indicating a high NADH pressure [7]. Glycerol synthesis from dihydroxyacetone phosphate consists of two steps, catalyzed by NAD\(^{+}\)-dependent glycerol-3-phosphate dehydrogenase (encoded by GPD1 and GPD2) and glycerol-3-phosphate phosphatase (encoded by GPP1 and GPP2), respectively (Fig. 1). Glycerol accumulation is also an issue in bioethanol production and has therefore been studied for years. Several studies found that glycerol formation could be decreased with altered redox-cofactor specificity [19–22], and could be eliminated through inactivation of both GPD1 and GPD2 with the expense of a dramatic reduction in cell growth [23]. As alternatives, an NADP\(^{+}\)-dependent glyceraldehyde-3-phosphate dehydrogenase (GapN) was demonstrated to enable rescuing the negative effects and improve ethanol yield [24, 25]. Moreover, since GapN irreversibly converts glyceraldehyde-3-phosphate to 3-phosphoglycerate and produces NADPH (Fig. 1), it was reported to increase the synthesis of NADPH demanding bioproducts, including 3-hydroxypropionic acid (3-HP) and polyhydroxybutyrate (PHB) [26, 27]. Therefore, it could potentially eliminate glycerol accumulation and increase FFA production by interruption of glycerol synthesis and expression of GapN.

The goal of this study is to investigate the effect of expressing the cytosolic PDH complex on FFA production in yeast strains with deregulated fatty acid metabolism and optimized cytosolic redox balance.
The strategies reported here may shed lights for production of other acetyl-CoA and NADPH demanding chemicals as well.

**Results And Discussion**

**Expression of a bacterial PDH complex to increase the acetyl-CoA supply**

The three subunits of PDH complex, encoded by \textit{pdhA}, \textit{pdhB}, \textit{aceF} and \textit{lpd}, and lipoate ligases from \textit{E. faecalis}, encoded by \textit{lplA} and \textit{lplA2} [16], were integrated into the two genome sites Chr XII-5 and Chr XI-3 [28] of the FFA production strain YJZ08 (\textit{faa1A faa4A pox1A hfd1A}) [6], yielding strain PDH1. As shown in Fig. 2, YJZ08 with deregulated FA synthesis produced FFA at 458.9 mg/L in 72 h in minimal medium with 20 g/L glucose, which was consistent with previous studies [6, 9]. Meanwhile, the constructed strain PDH1 produced 512.7 mg/L FFA at 72 h (Fig. 2A). From the growth data, we found that \textit{E. faecalis} PDH reduced the maximum specific growth rate from 0.33 h\(^{-1}\) to 0.28 h\(^{-1}\) but didn't result in much difference in the final biomass. PDH1 accumulated 1.90 g/L glycerol, compared with 1.90 g/L glycerol with YJZ08 (Fig. 2A), suggesting that cytosolic PDH did not cause an NADH burden in YJZ08.

Theoretically, a combination of the PDH route with the phosphoketolase (PK) pathway can result in a higher maximum FFA yield [15], since the PDH route could produce acetyl-CoA with low energy cost and the PK pathway could produce acetyl-CoA with low carbon loss. Moreover, the PK pathway utilizes xylulose-5-phosphate from the pentose phosphate (PP) pathway as the substrate, and it may thermodynamically help to pull the carbon flux towards the PP pathway and increase NADPH generation. However, we tested the integration of xylulose-5-phosphate phosphoketolase from \textit{Leuconostoc mesenteroides} [29] and phosphotransacetylase from \textit{Clostridium kluyveri} [30] into both YJZ08 and PDH1, and did not found significant improvement in cell growth and FFA synthesis (Supplementary Fig. 1), probably because of the strict control of carbon flux through the PP pathway [31] or inefficient phosphoketolase enzymes [29].

Moreover, the yeast ZS01 strain with simplified lipid metabolic networks and redirected flux towards FFA synthesis was constructed by deletion of \textit{PAH1}, \textit{DPP1}, \textit{LPP1} and \textit{ARE1} in YJZ08. Then, the genes encoding the PDH complex, composing of \textit{pdhA}, \textit{pdhB}, \textit{aceF}, \textit{lpd}, \textit{lplA} and \textit{lplA2}, were integrated into the chromosome of ZS01, yielding strain PDH2. As shown in Fig. 3, when cultivated in minimal medium with 20 g/L glucose, ZS01 accumulated 497.9 mg/L FFAs at 72 h with a high productivity of 133.7 mg/L/OD, which was consistent with previous studies [8, 9]. After 72 h, ZS01 accumulated more FFAs since there was ethanol remaining and the FFA titer reached 734.2 mg/L at 144 h when ethanol was exhausted (Fig. 2A). Interestingly, PDH2 produced 660.2 mg/L FFAs at 72 h and 756.3 mg/L FFAs at 96 h when ethanol was completely exhausted. In PDH2, the faster FFA production rate accompanied with faster ethanol consumption rated and lower glycerol levels, indicating that cytosolic PDH may alleviate NADH burden and speed up cell growth and FFA production (Fig. 2B), probably due to the lower energy cost.
Due to the possible mismatch between acetyl-CoA supply, NADH production and NADPH consumption for FFA synthesis, another hypothesis for these effects is that PDH from *E. faecalis* might utilize NADP\(^+\) as well as NAD\(^+\) as redox cofactors. Therefore, we evaluated PDH in a zwf1Δ ald6Δ mutant, which is inviable on glucose medium due to lack of ability to produce cytosolic NADPH [32]. Indeed, the results showed that cytosolic PDH expression could rescue cell growth of the zwf1Δ ald6Δ mutant on synthetic glucose medium, and disruption of E3 encoding gene *lpd* suspended the growth recovery (Fig. 2D). The growth assay suggested that the PDH complex, possibly the E3 domain, might utilize NADP\(^+\) as redox cofactor as well. However, growth assay on minimal glucose medium didn't show similar results, which might be caused by its poor activity. We therefore hypothesize that protein engineering on E3 (Lpd) could enhance its cofactor factor preference on NADP\(^+\). A previous study showed that cofactor specificity of *E.coli* PDH could be converted from NAD\(^+\) to NADP\(^+\) via seven amino acid mutations in E3 [33, 34]. The alignment result of E3 proteins from *E. faecalis* and *E. coli* suggested that the mutated amino acids are highly conserved, as shown in Supplementary Fig. 2A. When these E3 mutations were introduced into the strain PDH1, FFA titer didn't increase as expected, but decreased to 439.5 mg/L, comparable with YJZ08 (Supplementary Fig. 2B). The FFA results indicated that the mutations resulted in enzymatic activity lost instead of cofactor specificity conversion, probably due to unsuccessful protein assembly. To improve its enzymatic activity towards NADP\(^+\), rational design based on PDH structures or directed evolution will be required for further investigation. As *E. faecalis* PDH has been demonstrated to be a functional enzyme in *E. coli* and yeast at both aerobic and anaerobic conditions [16, 35], a novel PDH complex with NADP\(^+\) preference will clearly be valuable for bioproduction of acetyl-CoA derived chemicals.

It was also observed that, PDH expression altered the distribution of saturated and unsaturated FFAs (C16 and C18) in PDH2, but not in PDH1 (Fig. 2C). The double bonds in unsaturated FFAs were formed by Delta-9 fatty acid desaturase, encoded by *OLE1*, through an oxygen-dependent mechanism that requires reducing equivalents from NADH. Thus, the increase in unsaturated FFA ratio might also indicate that the functional PDH resulted in cytosolic redox changes without imposing an increased NADH burden, due to its insensitive feature to high NADH/NAD\(^+\) ratios [35–37].

**Balance of cytosolic redox factors for FFA production**

NAD and NADP redox couples are crucial to maintain cellular redox hemostasis, with lower cytosolic NADH/NAD\(^+\) ratios (range from 0.001 to 0.01) and higher NADPH/NADP\(^+\) ratios (range from 15 to 60) due to their different functions [38]. In FFA production strains, over-production of NADH resulted in glycerol accumulation, while NADPH as the cofactor required for fatty acid elongation was limited. Thus, the balance between NAD and NADP redox couples would be important for FFA production.

To balance the redox state of the cell and further improve FFA synthesis, NADP\(^+\)-dependent glyceraldehyde-3-phosphate dehydrogenase (*GapN*) from *Streptococcus mutans* [39] was expressed on a multi-copy plasmid pGapN in the FFA production strain YJZ08, yielding strain ZS02. As shown in Fig. 3, *GapN* expression resulted in reduced glycerol accumulation and increased FFA production.
We further balanced cytosolic redox conditions through interruption of glycerol synthesis via glycerol-3-phosphate dehydrogenase (GPD) deletions. Since \( GPD1 \) was regulated by osmotic stress [23], \( GPD2 \) was first deleted in YJZ08, and the resulting strain ZS03 showed decreased glycerol accumulation, and increased FFA titer and productivity (Fig. 3A). When both \( GPD1 \) and \( GPD2 \) were deleted, the specific growth rate of the resulted strain ZS04 decreased dramatically, which was consistent with previous studies [23]. The final FFA titer of ZS03 was 556.0 mg/L, 23.6% higher than YZJ08. When \( GapN \) was introduced into ZS04, the impaired growth was improved, and FFA production at 72 h improved significantly. The final FFA titer of ZS05 reached 578.7 mg/L at 96 h, which was 28.6% higher than YJZ08 and the productivity was comparable with ZS04.

Similarly, the distributions of saturated and unsaturated FFA also varied in the constructed strains (Fig. 3B). \( GPD \) deletion did not alter the distributions, whereas \( GapN \) expression increased the ratios of saturated FFAs. With \( GapN \) expressed in YJZ08, the percentage of saturated FFAs increased slightly from 50–53%, with the increases in the titers of C16:0 and C18:0 but no increases of C16:1 and C18:1. When \( GapN \) expressed in \( gpd1 \Delta gpd2 \Delta \) mutant, in ZS05 the percentage of saturated FFAs further increased to 59%, indicating the possible redox changes between NAD and NADP redox couples.

**Increased provision of acetyl-CoA and NADPH improved FFA production**

Plasmid pGapN was transformed into PDH1 and PDH2, yielding PDH3 and PDH6, respectively. As shown in Fig. 4, the FFA production in PDH3 and PDH6 increased to 534.0 mg/L and 797.6 mg/L, respectively. Also, glycerol accumulation levels in PDH3 and PDH6 reduced to 0.93 g/L and 0.58 g/L, respectively. The decrease in glycerol level and increase in FFA titer in both PDH3 and PDH6 were similar with those observed in ZS02.

Similarly, when \( GPD1 \) and \( GPD2 \) were deleted in PDH1 and PDH2, the resulting strains PDH4 and PDH7 exhibited decreased growth rates, but increased FFA production to 664.0 mg/L and 840.5 mg/L, respectively (Fig. 4). Finally, when \( GapN \) was introduced into PDH3 and PDH6, yielding PDH5 and PDH8, the impaired cell growth improved, whereas the FFA production decreased to 614.0 mg/L and 715.3 mg/L, respectively (Fig. 4). The improved cell growth and decreased FFA titer upon \( GapN \) expression indicated a possible competition for the supply of the precursors and redox factors.

Regarding the distribution of saturated and unsaturated FFAs (C16 and C18) of engineered PDH strains, it was found that PDH expression significantly increased the ratios of unsaturated FFAs (Fig. 5), even with efforts for redox cofactor rebalance. These changes might be because of the PDH complex with excess NADH produced and its consistent activity under high NADH/NAD\(^+\) ratios. The PDH complex may be a potential target to increase the unsaturation of the FFAs and their derivatives, besides manipulations on Delta-9 fatty acid desaturase OLE1 reported to control the fatty acid saturation in previous studies [40, 41].
Conclusions

The expression of the PDH complex from *E. faecalis* in the yeast cytosol enabled efficient acetyl-CoA provision with low ATP cost, and improved FFA production in the titer, productivity and rate. Rebalancing NAD and NADP redox couples further improved FFA production in PDH expressing strains. Briefly, the *GapN* expression seemed to increase FFA production via the growth improvement, and *GPD* deletion resulted in dramatically decreased in cell growth probably due to the high cytosolic NADH burden. Moreover, the increase in FFA unsaturation in PDH strains suggested that the PDH complex can be further engineered for the production of unsaturated fatty acids and their derivatives.

Materials And Methods

2.1 Construction of plasmids and strains

The plasmids, strains, and primers used and constructed in this study are listed in Dataset S1. All yeast strains were constructed using GTR-CRISPR system as described in [8]. For gene integration and deletion, donor DNA containing 50 bp of homology arms and Cas9 plasmid containing gRNA targeting sequence were co-transformed using the electroporation method as described in [42].

Six genes for PDH pathway from *Enterococcus faecalis* [16] were synthesized by Sangon Company (Shanghai, China) and then cloned in pUC57 vectors, yielding plasmids pPdhB, pPdhA, pLpd, pAceF, pLplA and pLplA2, respectively. *lplA* and *lplA2* were cloned using Golden Gate into the vector pAMPori with primers PP1-5, yielding pLplA_A2. These six genes were integrated into two sites XI-3 and XII-5 as described in [28]. *lplA* and *lplA2* were integrated at XI-3 using the Cas9 plasmid pCas9_XI-3.3 and donor DNA amplified from pLplA_A2 with primers PP6 and PP7. *pdhB*, *pdhA*, *lpd* and *aceF* were integrated at the XII-5 site simultaneously using pCas9_XII-5 and donor DNA amplified from the corresponding plasmids with primers PP8-15.

Codon-optimized *GapN* from *Streptococcus mutans* [39] was synthesized by Sangon Company (Shanghai, China) and cloned in pAMPori vector using Golden Gate assembly with primers PP16-23, yielding pGapN. *GapN* was then integrated into XI-1 using pCas9_XI-1 and donor DNA amplified with primers PP24 and PP25 from pGapN.

*GPD2* was deleted using pCas9_gpd2 and donor DNA amplified with primers PP26 and PP27. *GPD1* was deleted using pCas9_gpd1 and donor DNA amplified with primers PP28 and PP29. *PAH1*, *DPP1*, *LPP1* and *ARE1* were simultaneously deleted using pCas9_PDLA and donor DNA fragments with primers PP30-PP37. *ZWF1* was deleted using pCas9_zwf1 and donor DNA amplified with primers PP38 and PP39. *ALD6* was deleted using pCas9_ald6 and donor DNA amplified with primers PP40 and PP41. *LPD* was deleted using pCas9_lpd and donor DNA amplified with primer PP42 and PP43.

All Cas9 plasmids containing gRNAs for target genes and were constructed using Golden Gate assembly. Take pCas9_gpd2 as an example, PCR fragments amplified from pScURA with primer PP47, containing
20 bp gRNA targeting sequence, were assembled into pCas plasmid [8].

2.2 Media and cultivation conditions

*E. coli* strains were cultured at 37 °C in LB medium, composed of 5 g/L yeast extract, 10 g/L peptone, and 10 g/L NaCl, and 80 mg/ml ampicillin was added when needed.

Yeast strains were cultured either in YPD medium composed of 10 g/L yeast extract, 20 g/L peptone and 20 g/L glucose, or SC-URA medium for the selection of strains containing URA3-based plasmids, composed of 5 g/L (NH₄)₂SO₄, 1.7 g/L yeast nitrogen base without amino acids, 1.914 g/L amino acid mixture without uracil, and 20 g/L glucose. 5-FOA plates were used to lose URA3-based plasmid, composed of 1.7 g/L yeast nitrogen base without amino acids, 5 g/L (NH₄)₂SO₄, 1.914 g/L amino acid mixture without uracil, 0.05 g/L uracil, 1 g/L 5-FOA, 25 g/L agar. All solid plates contained 15 g/L agar except for 5-FOA plates.

Minimal medium was used for cultivation in shake flasks for FFA production, composed of 5 g/L (NH₄)₂SO₄, 14.4 g/L KH₂PO₄, 0.5 g/L MgSO₄·7H₂O, trace metal solution and vitamin solution and 20 g/L glucose, and 40 mg/L histidine or 60 mg/L uracil was added when needed. For all PDH strains, minimal medium was supplemented with 100 mg/mL lipoic acid.

Yeast strains were cultured at 30 °C. For shake flask cultivation, yeast cells were precultured in falcon tubes with 5 ml minimal medium for 24 h, and then inoculated into 100 mL shake flasks with 30 mL medium with initial OD₆₀₀ of 0.1.

2.3 Measurement of biomass and extracellular metabolites

Biomass was measured by optical density at 600 nm (OD₆₀₀) with a GENESYS 30 Visible Spectrophotometer (Thermo Electron Scientific, Madison, USA).

Extracellular metabolites including glucose, ethanol and glycerol were measured by HPLC (Shimadzu LC-20AT, Japan) equipped with RID and UV detectors, using 5 mM H₂SO₄ as eluent and Aminex HPX-87H column (Bio-Rad) at 65 °C with a flow rate of 0.6 ml/min.

2.4 Extraction, measurement and analysis of FFAs

FFAs in the broth were extracted and measured as described in [6]. Specifically, 200 µl of cell culture or diluted cell culture was transferred into a glass vial, and 10 µl 40% tetrabutylammonium hydroxide and 200 µl methylation reagent was added immediately. Methylation reagent contained 200 mM methyl iodide as methyl donor and 12.5 mg/L pentadecanoic acid as an internal standard. The mixture was shaken for 30 min, and then centrifuged at 5000 g for 3 minutes. Then, 100 µl of dichloromethane layer with extracted methyl esters was transferred into a GC vial, evaporated 3 h, and resuspended in 100 µl hexane. Samples were analyzed by GC-MS (QP2020, Shimadzu, Japan) with a DB-5MS column (30m × 0.250mm × 0.25 µm, Agilent). The program was set as follows: 40 °C 2 min; ramp to 130 °C at 5 °C/min;
raise to 280 °C at 10 °C/min; 280 °C 3 min. The temperatures of the inlet, mass transfer line and ion source were set at 280 °C, 300 °C, 230 °C, respectively. Flow rate of carrier gas was 3.0 mL/min and the data were collected in the full inspection mode (50–650 m/z) and analyzed using GCMS solution 4.4 software.

Samples for FFA analysis were harvested at different time points due to varied strain growth. The highest FFA titers were achieved when ethanol was completely consumed.

**Abbreviations**

PDH  
pyruvate dehydrogenase  
FFA  
free fatty acid  
GPD  
NAD⁺-dependent glycerol-3-phosphate dehydrogenase

**Declarations**

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Availability of data and materials**

All datasets generated and analyzed during this study are included in this published article and its supplementary information files.

**Competing interests**

The authors declare no competing interests.

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**Author’s contributions**
YZ, MS, JN and ZL designed the research and wrote the paper; YZ, MS, NQ performed the experiment. ZL and JN supervised the research. All authors read and approved the final manuscript.

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**Figures**

![Figure 1](image)

**Figure 1**

Schematic representation of engineering strategies for FFA production in *S. cerevisiae*. Abbreviation used: G3P, glyceraldehyde-3-phosphate; 3PG, 3-phospho-glycerate; cPDH, cytosolic pyruvate dehydrogenase.
Figure 2

The expression of PDH improved FFA production in yeast with simplified lipid networks. A) FFA production with expression of PDH complex in YJZ08. B) Growth profile with expression of PDH complex in ZS01. C) Distribution of saturated and unsaturated FFAs (C16 and C18) in engineered strains. Statistical analysis was performed using one-tailed Student’s t-test (*p < 0.05). All data represent the mean ± SD of biological triplicates. D) Growth assay of engineered strains with and without cytoplasmic PDH in glucose medium.
Figure 3

FFA production with GapN expression and GPD deletions. A). FFA production obtained with engineered strains. B) Distribution of saturated and unsaturated FFAs (C16 and C18) in the engineered strains. Statistical analysis was performed using one-tailed Student’s t test (*p < 0.05, **p< 0.01). All data represent the mean ± SD of biological triplicates.
Figure 4

FFA production with cytosolic PDH and cofactor engineering. All data represent the mean ± SD of biological triplicates.
Figure 5

Distribution of saturated and unsaturated FFAs (C16 and C18) in engineered strains. All data represent the mean ± SD of biological triplicates.

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