Engineering *Yarrowia lipolytica* as a chassis for *de novo* synthesis of aromatic-derived natural products and chemicals

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Abstracts

Yarrowia lipolytica is an attractive host to upgrade renewable low-cost carbon feedstocks to high-value

commodity chemicals and natural products. In this work, we systematically characterized and removed the

rate-limiting steps of the shikimate pathway and achieved de novo synthesis of five aromatic chemicals in Y.

lipolytica. We determined that engineering feedback-insensitive DAHP synthases and eliminating amino acids

formation are critical steps to mitigate the feedback regulation of shikimate pathway. Further overexpression

of heterologous phosphoketolase and deletion of pyruvate kinase provided a sustained metabolic driving force

that channels E4P (erythrose 4-phosphate) and PEP (phosphoenolpyruvate) precursors through the shikimate

pathway. Precursor competing pathways and byproduct formation pathways were also blocked by inactivating

chromosomal genes. To demonstrate the utility of our engineered chassis strain, three natural products, 2-PE,

p-coumaric acid and violacein, which were derived from phenylalanine, tyrosine and tryptophan, respectively,

were chosen to test the chassis performance. We obtained 2426.22 mg/L of 2-phenylethanol (2-PE), 593.53

mg/L of p-coumaric acid, 366.30 mg/L of violacein and 55.12 mg/L of deoxyviolacein from glucose in shake

flask. The 2-PE production represents a 286-fold increase over the initial strain (8.48 mg/L). Specifically, we

obtained the highest 2-PE, violacein and deoxyviolacein titer ever reported from the de novo shikimate

pathway in yeast. These results set up a new stage of engineering Y. lipolytica as a sustainable biorefinery

chassis strain for de novo synthesis of aromatics with economic values.

Keywords: shikimate pathway, metabolic bottlenecks, *Y. lipolytica*, aromatic metabolism, feedback regulation,

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metabolic engineering

1. Introduction

Y. lipolytica, as a 'generally regarded as safe' (GRAS) yeast ¹, has been extensively engineered for the production of oleochemicals, fuels and commodity chemicals ²⁻⁵. The abundant acetyl-CoA and malonyl-CoA precursors in *Y. lipolytica* have been harnessed for synthesizing plant secondary metabolites in recent years, including flavonoids ⁶⁻⁸, polyketides ^{9, 10}, polyunsaturated fatty acids ^{11, 12} and isoprenoids ¹³⁻¹⁵. A large collection of customized genetic toolkits, including Golden-gate cloning ¹⁶⁻¹⁸, genome integration ^{6, 19, 20}, CRISPR-Cas9/Cpf1 genome editing ²¹⁻²³, transposons ²⁴, auxotrophic markers ²⁵ and promoter libraries ²⁶⁻²⁸ have accelerated our ability to perform targeted genetic manipulations. Different from *S. cerevisiae*, *Y. lipolytica* lacks Crabtree effects generating no overflowed metabolism toward ethanol under high glucose conditions ²⁹, which might be more suitable for high gravity fermentation and process control. *Y. lipolytica* is reported to valorize a broad range of cheap and renewable feedstocks ^{30, 31}, including sugars, volatile fatty acids, alkanes and municipal organic wastes. This substrate flexibility provides us an environmentally friendly approach to upgrade low-value carbons to high value chemicals with reduced carbon footprint and improved process economics.

Aromatic compounds have wide applications ranging from health care to food industry, nutraceutical supplements and pharmaceutical intermediates, which collectively represent a multi-billion-dollar global market ^{32,34}. Although metabolically-engineered *E. coli* has achieved gram-per-liter levels of aromatic compounds, primarily phenylpropanoids ^{35,36}, yeast proves to be a more attractive host, due to its GRAS status, robust cell growth, tolerance of harsh conditions (such as low pH and high osmolarity), and the spatially-organized subcellular compartment for regio- or stereo- activity of cytochrome P450 enzymes ^{7,32,37}. Up to date, various metabolic engineering strategies have been implemented in *E. coli* and *S. cerevisiae* to improve aromatics production, including expression of the feedback-insensitive DAHP synthases ³⁸ and chorismite synthase, enhancing the shikimate pathway ³⁹ and modular microbial coculture ^{40,42}. A number of studies have been focused on optimizing the endogenous Ehrlich pathway to produce 2-phenylethanol up to 3-6 g/L ^{43,45}, however, the expensive precursor L-phenylalanine was fed into the bi-phasic bioreactors ^{46,47}, restricting its economic potential for large-scale production. The highest titer of 2-phenylethanol production from the *de novo* shikimate pathway was reported recently as 1.58 g/L (13 mM) from *S. cerevisiae* ³⁸. There is a pressing need to explore the metabolic potential of alternative yeast for aromatics production.

In this work, we attempted to engineer the nonconventional oleaginous yeast (Y. lipolytica) as a

competitive platform host to produce aromatic derivatives (Figure 1). With 2-phenylethanol (2-PE) as the

testbed molecule, we systematically characterized and removed the bottlenecks of the endogenous shikimate

pathway, resulting in the production of 2426.22 mg/L (19.9 mM) of 2-PE from glucose, the highest titer ever

reported from the de novo shikimate pathway. Using this yeast as chassis, we further redirected the shikimate

flux toward other aromatics and achieved high tier of p-coumaric acid (593.53 mg/L), violacein (366.30 mg/L),

highest titer reported in yeast) and deoxyviolacein (55.12 mg/L, highest titer reported in yeast) in shaking

flasks, indicating the superior metabolic potential of Y. lipolytica as an aromatics-producing host. This report

highlights the prominent metabolic characteristics of Y. lipolytica as chassis for production of various

aromatics and natural products with economic values.

2. Material and Methods

2.1. Strains, plasmid, primers, and chemicals

All stains of engineered Y. lipolytica, including the genotypes, recombinant plasmids, and primers have

been listed in Supplementary Table1 and 2. Chemicals used in this study were all purchased from

Sigma-Aldrich. Codon-optimized heterologous synthetic genes, including genes BbxfpK, AcxpkA, RgTAL,

VvSTS1, and Pc4CL2, were ordered from GENEWIZ (Suzhou, China). Codon-optimization was performed

with the IDT website.

2.2. Shake flask cultivations

For performing shake flask cultivations, seed culture was carried out in the shaking tube with 2 mL seed

culture medium at 30 °C and 250 r.p.m. for 48 h. Then, 0.8 mL of seed culture was inoculated into the 250 mL

flask containing 30 mL of fermentation medium and grown under the conditions of 30 °C and 250 r.p.m. for

120 h. One milliliter of cell suspension was sampled every 24h for OD₆₀₀, glucose, and desired metabolisms

measurements.

Seed culture medium used in this study included the yeast complete synthetic media regular media (CSM,

containing glucose 20.0 g/L, yeast nitrogen base without ammonium sulfate 1.7 g/L, ammonium sulfate 5.0

g/L, and CSM-Leu 0.74 g/L) and complex medium (YPD, containing glucose 20.0 g/L, yeast extract 10.0 g/L,

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and peptone 20.0 g/L). Fermentation medium used in this study also included the yeast complete synthetic media regular media (CSM, containing glucose 40.0 g/L, yeast nitrogen base without ammonium sulfate 1.7 g/L, ammonium sulfate 5.0 g/L, and CSM-Leu 0.74 g/L) and complex acetate medium (YPA, containing glucose 40.0 g/L, yeast extract 10.0 g/L, peptone 20.0 g/L, and sodium acetate 5.0 g/L).

2.3. Yeast transformation and screening of high-producing strains

The standard protocols of *Y. lipolytica* transformation by the lithium acetate method were described as previously reported ^{48, 49}. In brief, one milliliter cells was harvested during the exponential growth phase (16-24 h) from 2 mL YPD medium (yeast extract 10 g/L, peptone 20 g/L, and glucose 20 g/L) in the 14-mL shake tube, and washed twice with 100 mM phosphate buffer (pH 7.0). Then, cells were resuspended in 105 uL transformation solution, containing 90 uL 50% PEG4000, 5 uL lithium acetate (2M), 5 uL boiled single stand DNA (salmon sperm, denatured) and 5 uL DNA products (including 200-500 ng of plasmids, lined plasmids or DNA fragments), and incubated at 39 °C for 1 h, then spread on selected plates. It should be noted that the transformation mixtures needed to be vortexed for 15 seconds every 15 minutes during the process of 39 °C incubation. The selected markers, including leucine, uracil and hygromycin, were used in this study. All engineering strains after genetic manipulations were performed optimized screening by the shaking tube cultivations, and the optimal strain was used to perform shaking flask (these data have been shown in Supplementary Materials).

2.4. Single-gene and multi-genes expression vectors construction

In this work, the YaliBrick plasmid pYLXP' was used as the expression vector ²⁸. The process of plasmid constructions also has been reported ⁵⁰. In brief, recombinant plasmids of pYLXP'-xx (a single gene) were built by Gibson Assembly of linearized pYLXP' (digested by *SnaBI* and *KpnI*) and the appropriate PCR-amplified or synthetic DNA fragments. Multi-genes expression plasmids were constructed based on restriction enzyme subcloning with the isocaudamers *AvrII* and *NheI* ^{51, 52}. All genes were respectively expressed by the TEF promoter with intron sequence and XPR2 terminator, and the modified DNA fragments and plasmids were sequenced by Quintarabio.

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2.5. Gene knockout

A marker-free gene knockout method based on Cre-lox recombination system was used as previously reported ⁵³. For performing gene knockout, the upstream and downstream sequences (both 1000 bp) flanking the deletion targets were PCR-amplified. These two fragments, the loxP-Ura/Hyr-loxP cassette (digested from plasmid pYLXP'-loxP-Ura/Hyr by AvrII and salI), and the gel-purified plasmid backbone of pYLXP'(linearized by AvrII and salI) were joined by Gibson Assembly, giving the knockout plasmids pYLXP'-loxP-Ura/Hyr-xx (xx is the deletion target). Next, the knockout plasmids were sequence-verified by Quintarabio. Then, the gene knockout cassettes were PCR-amplified from the knockout plasmids pYLXP'-loxP-Ura/Hyr-xx, and further transformed into Y. lipolytica. The positive transformants were determined by colony PCR. Ku70 (Po1f background) was knocked out by screening more than 200 yeast colonies. Other knockout strains were built on top of the Ku70-deficient strains. Subsequently, plasmid pYLXP'-Cre was introduced into the positive transformants and promoted the recombination of loxP sites, which recycle the selected marker. Finally, the intracellular plasmid pYLXP'-Cre was evicted by incubation at 30 °C in YPD media for 48h. Here, Ura is the uracil marker, and Hyr is hygromycin marker. Specifically, for deletion of gene ylPYK, final concentration of 0.5, 1, 1.5, 2, 2.5, 5 g/L acetate was added into the selective CSM-Ura plate to rescue the cell growth. Besides integration of linearized plasmid, other genomic manipulations in platform strain YL33, including gene knockout and integration of desired genes, hygromycin resistance gene was used as the selective marker grown on YPD plate containing 200 mg/L hygromycin.

2.6. Genomic integration of desired genes

In this work, genomic integration of desired genes were performed in two different ways: site-specific genomic integration plasmids or application of pBR docking platform by linearizing the plasmid pYLXP' with digested enzyme *NotI*. Here, to meet specific requirements, we constructed three genomic integration plasmids pURLK, pURLA, and pHyLD, corresponding to the *Ku70*, *YALI0E30965g* (encoding acetyl-CoA hydrolase, named as Ace), and *YALI0E03212g* (encoding lactate dehydrogenase, named as LDH) genomic sites, respectively. The procedure of using these three plasmids was similar as that of gene knockout protocol. Plasmid maps and sequences for pURLK, pURLA, and pHyLD have been uploaded in the Supplementary Materials. Specifically, desired genes could be assembled into pURLX based on restriction enzyme subcloning by using the multiple cloning sites (see plasmid maps in SI files), and the integration cassettes of desired genes were retrieved by digesting plasmid pURLX-xx with enzyme *AvrII*. Additionally, the standard

protocol of 26s rDNA genomic integration by plasmid prDNAloxP was used in this work, with detailed protocol described in previous work ⁶.

The application of pBR docking platform was achieved by linearizing the plasmid pYLXP' with *NotI* restriction enzymes. It should be noted that, plasmids pYLXP'-ylPAR4-ylAR010-ylAR07-ylPHA2-scAR07^{G141S} and pYLXP'-VioDCBAEI have more than 2 *NotI* digestion sites, thus, we performed site-directed mutagenesis to change the *NotI* site to *SnaBI* site in plasmid pYLXP', generating plasmid pYLXPs'. Subsequently, gene fragments in pYLXPs'-VioDCBAE, pYLXPs'-ylPAR4-ylAR010-ylAR07-ylPHA2-scAR07^{G141S} and its derivates were linearized by *SnaBI* digestion, and then the linearized fragments were integrated at genomic pBR docking site.

2.7. Quantification of biomass, glucose, 2-PE, *p*-coumaric acid, resveratrol, violacein and deoxyviolacein

Cell densities were monitored by measuring the optical density at 600 nm (OD₆₀₀). The concentrations of 2-PE, penylacetate, glucose, p-coumaric acid, resveratrol, violacein, and deoxyviolacein were all measured by high-performance liquid chromatography (HPLC) through Agilent HPLC 1220. In detail, 2-PE and penylacetate were measured at 215 nm under 40 °C (column oven temperature) with a mobile phase containing 50% (v/v) methanol in water at a flow rate of 0.5 mL/min equipped with a ZORBAX Eclipse Plus C18 column (4.6 × 100 mm, 3.5 μ m, Agilent) and the VWD detector. The concentrations of glucose were measured by a SupelcogelTM Carbohydrate column (Sigma, USA) and a refractive index detector with H₂SO₄ (5 mM) as the mobile phase at a flow rate of 0.6 mL/min at 40 °C.

To quantify the concentration of p-coumaric acid, 0.1 mL of fermentation culture was mixed with 9-fold volume of absolute methanol (100% v/v), vortexed thoroughly, and centrifuged at 12000 r.p.m. for 10 min. The supernatants were analyzed at 304 nm under 40 °C (column oven temperature) with a mobile phase containing 45% (v/v) methanol in water at a flow rate of 0.5 mL/min equipped with a ZORBAX Eclipse Plus C18 column (4.6 × 100 mm, 3.5 μ m, Agilent) and the VWD detector. To quantify the concentration of resveratrol, 0.25 mL of fermentation culture was mixed with an equal volume of ethyl acetate and appropriate glass beads, vortexed at 30 °C for 24 h, and centrifuged at 12000 r.p.m. for 10 min. Then, 100 uL supernatants of top organic layer was transferred to glass vial and evaporated to dryness, then resolubilized with 100 uL of methanol. Resveratrol samples were analyzed with the same HPLC protocols as p-coumaric acid.

To quantify the concentration of violacein and deoxyviolacein, 0.20 mL of fermentation culture was mixed with 5-fold volume of ethyl acetate and appropriate glass beads, vortexed at 30 °C for 24 h, and centrifuged at 12000 r.p.m. for 10 min. The supernatants of top organic layer were analyzed at 570 nm under 40 °C (column oven temperature) with a gradient method with two solvents, water (A) and methanol (B), at a flow rate of 0.4 mL/min equipped with a ZORBAX Eclipse Plus C18 column (4.6 × 100 mm, 3.5 μm, Agilent) and the VWD detector. The elution started with 100% of solvent A, the fraction of solvent A was decreased linearly from 100% to 20% (0-5 min), and maintained at 20% for 3 min (5-8 min), then the fraction of solvent A was increased from 20% to 100% (8-12 min), and maintained at 100% for 1 min (12-13 min).

3. Results and discussions

3.1. De novo synthesis of 2-PE from shikimate pathway

In this work, the Y. lipolytica endogenous metabolite 2-phenylethnol (2-PE, an aromatic alcohol that is widely used as flavor and fragrance agent) was chosen as a target molecule to determine the bottlenecks of the shikimate pathway. 2-PE synthesis is a metabolic branch derived from shikimate pathway, in which 2-PE is synthesized through four enzyme-dependent cascade reactions starting from chorismate (Figure 2a). To identify the potential limiting-steps in 2-PE synthesis, we firstly systematically overexpressed all genes that are involved in 2-PE synthesis under the control of strong constitutive pTEF-intron promoter ²⁸, including genes ylPAR4 (YALI0D07062g, encoding phenylacetaldehyde reductase), ylAR010 (YALI0D06930g, encoding phenylpyruvate decarboxylase), ylPHA2 (YALI0B17336g, encoding prephenate dehydratase) and ylARO7 (YALI0E17479g, encoding chorismate mutase). Besides gene ylARO10, individual overexpression of genes ylPAR4, ylPHA2, and ylAR07 did not result in a significant increase in 2-PE titer (Supplementary Figure S2), however, combining all four genes (strain YL5) led the strain to produce 55.53 mg/L of 2-PE from glucose (Supplementary Figure S2), a 6.55-fold increase in comparison with the control strain (without any gene overexpression, 8.48 mg/L), indicating that the four enzymes in the Ehrlich pathway are essential to channel the carbon flux toward 2-PE synthesis. In addition, chorismate mutase (ARO7) is known to be strictly feedback inhibited by aromatic acids ³⁹. To resolve this issue, introduction of a feedback-resistant chorismate mutase ScARO7^{G141S} originating from Saccharomyces cerevisiae further improved 2-PE production up to 87.63 mg/L (strain **YL6**, Figure 2b), which was consistent with the previous report ⁵⁴. We have configured the Ehrlich pathway to achieve *de novo* 2-PE synthesis from glucose with enhanced titer.

We subsequently optimized the shikimate pathway to unlock the potential of Y. lipolytica for production of aromatic compounds. The entire Ehrlich pathway for 2-PE synthesis contains multiple genes with total size more than 16,000 bp. To simplify the genetic manipulations, we integrated the 2-PE pathway at the genomic *YALI0E30965g* loci (encoding acetyl-CoA hydrolase) with the integration plasmid pURLA-ylPAR4-ylARO10-ylARO7-ylPHA2-scARO7^{G141S}. The resultant strain YL7 produced 50.13 mg/L of 2-PE from glucose (Figure 2b). In Y. lipolytica, shikimate pathway is consisted of seven enzymatic steps, which starts from precursors erythrose-4-phosphate (E4P) and phosphoenolpyruvate (PEP) and ends up with chorismate (Figure 1). By overexpressing genes ylARO1 (YALI0F12639g, encoding pentafunctional aromatic protein), ylARO2 (YALIOD17930g, encoding bifunctional chorismate synthase), ylARO3 (YALIOB20020g, encoding DAHP synthase), ylARO4 (YALI0B22440g, encoding DAHP synthase), and ylARO5 (YALI0C06952g, encoding DAHP synthase) in strain YL7, we generated strains YL8, YL9, YL10, YL11, and YL12 that were able to produce 2-PE with titers at 62.53, 52.80, 59.49, 54.71, and 57.16 mg/L (Supplementary Figure S3), respectively. Strains YL8, YL10, and YL12 showed slight increase of 2-PE titer compared with strain YL7. However, by simultaneous overexpression of genes ylARO1 and ylARO2, 2-PE production was increased to 84.64 mg/L (strain YL13, Supplementary Figure S3). Further overexpression of ylARO3, ylARO4, and ylARO5 did not have significant effect on 2-PE titer (strain YL14, 78.28 mg/L, Supplementary Figure S3), which is possibly due to the allosteric regulation of DAHP synthase by aromatic amino acids ^{32, 39}.

3.2. Relieving metabolic bottlenecks in the shikimate pathway

Heterologous DAHP synthases scARO4^{K229L} (originating from *S. cerevisiae*), AroG^{L175D} and AroG^{S180F} (originating from *E. coli*) previously have been shown to be feedback-resistant of aromatic amino acids regulation ³⁹. We next introduced three variants of the feedback-resistant DAHP synthases into the 2-PE chassis strain, and generated strains **YL15**, **YL16**, and **YL17**. Shake flask cultivation of strains **YL15**, **YL16**, and **YL17** led to the production of 313.13, 48.70, and 430.11 mg/L of 2-PE (Figure 3a, b), respectively. This result clearly demonstrates that overexpression of scARO4^{K229L} and AroG^{S180F} are effective to remove the shikimate pathway bottleneck in *Y. lipolytica*. Likewise, the results indicated the importance of relieving allosteric regulation of DAHP synthase to increase carbon flux toward shikimate pathway.

On the other hand, we sought to eliminate the synthesis of aromatic amino acids byproducts, including phenylalanine, tryptophan, and tyrosine. For this purpose, five genes (Figure 3c) were chosen as knockout targets, including ylTYR1 (YALI0F17644g, encoding prephenate dehydrogenase), ylTRP2 (YALI0D11110g, encoding anthranilate synthase), ylTRP3 (YALI0E14751g, encoding anthranilate synthase), ylARO8 (YALI0E20977g, encoding aromatic amino acid aminotransferase), and ylARO9 (YALI0C05258g, encoding aromatic amino acid aminotransferase). By sequentially deleting ylTYR1, ylTRP2, ylTRP3, ylARO8, and ylARO9 in strain po1fk (which is Ku70-deficient), we got strains YL18 (auxotroph of tryptophan), YL19 (auxotroph of tryptophan and tyrosine), and YL20 (auxotroph of tryptophan, tyrosine and phenylalanine). Then the de novo 2-PE pathway (ylPAR4-ylARO10-ylARO7-ylPHA2-scARO7^{G141S}) was introduced into these auxotrophic strains (YL18, 19 and 20), leading to strains YL21, YL22 and YL23. In comparison to the parental strain YL6 (87.63 mg/L of 2-PE titer), strains YL21, YL22, and YL23 led to 2-PE production (Figure 3d) increased by 2.65-fold (231.84 mg/L), 3.49-fold (306.30 mg/L), and 3.86-fold (338.35 mg/L), respectively, despite that negative effect was observed in the cell growth. This result suggests that blocking of aromatic amino acids formation may further relieve allosteric regulation of DAHP synthase by L-Phe, L-Tyr and L-Trp.

Taken together, we have proved that introduction of feedback-resistant heterologous DAHP synthases and blocking aromatic amino acids formation are both effective to debottlenecking the shikimate pathway in *Y. lipolytica*. We next combined these two strategies to boost the 2-PE yield. Genes encoding *ylARO1* and *ylARO2-ylARO3-ylARO4-ylARO5-scARO4*^{K229L}-*aroG*^{S180F} were sequentially integrated at genome loci of *ku70* and *YALI0E30965g*, to generate the chassis strain **YL24**. By expression of the 2-PE pathway (*ylPAR4-ylARO10-ylARO7-ylPHA2-scARO7*^{G141S}) in strain **YL24**, the engineered strain produced 508.58 mg/L of 2-PE (strain **YL25**, Figure 3e), which was 1.18-fold, 1.50-fold, and 59.97-fold increase over the **YL17**, **YL23** and the starting strain **YL0**, respectively.

On the other hand, the operation of shikimate pathway is driven by the upstream precursors erythrose-4-phosphate (E4P, that is derived from pentose phosphate pathway) and phosphoenolpyruvate (PEP, that is derived from glycolysis). To further unlock the shikimate pathway, we next turned to rewire the carbon flows in the central carbon metabolism to optimize the supply of both E4P and PEP.

3.3. Removing pentose phosphate pathway bottleneck to boost precursor E4P

Metabolic flux analysis indicates that the available carbon flux toward E4P is as least one order of magnitude lower than the flux to PEP in yeast ³², which suggested the availability of E4P is the key step to maximizing flux to shikimate pathway. Overexpression of gene *ylTKT* (*YALI0E06479g*, encoding transketolase) improved 2-PE titer to 598.36 mg/L (strain **YL26**, Supplementary Figure S4), a 1.18-fold increase over the strain **YL25**. This result indicates that the supply of E4P indeed was a bottleneck.

In the pentose phosphate pathway, phosphoketolase splits fructose-6-phosphate into E4P and acetyl-phosphate (Figure 4a) 32, 55-57. In a recent study, overexpression of heterologous phosphoketolase BbxfpK from Bifidobacterium breve led to a 5.4-fold increase of intracellular E4P concentration in S. cerevisiae⁵⁵. We therefore introduced the codon-optimized phosphoketolase BbxfpK and AcxpkA (originating from Acidobacterium capsulatum) into our yeast chassis to generate strains YL27 and YL28, respectively. However, we observed declined 2-PE production (Supplementary Figure S4) in both strains YL27 (191.72 mg/L of 2PE) and YL28 (146.76 mg/L of 2PE). As plasmids genetic instability (i.e. unequal distribution/propagation of plasmid) was observed in our previous research, we speculated that a similar effect was occurred in strains YL27 and YL28. To resolve this issue, linearized gene fragments (see Methods) pYLXP'-BbxfpK-ylPAR4-ylARO10-ylARO7-ylPHA2-scARO7^{G141S} containing and pYLXP'-AcxpkA-ylPAR4-ylARO10-ylARO7-ylPHA2-scARO7^{G141S}, were integrated at the pBR docking site of strain YL24, generating strains YL29 and YL30, respectively. As expected, shake flask cultivation of strains YL29 and YL30 led to a 133.64% (679.66 mg/L) and 139.33% (708.58 mg/L) improvement in 2-PE titer (Supplementary Figure S4), compared to the parental strain YL25. This result clearly demonstrates that manipulation of the phosphoketolase pathway is a feasible strategy to channeling carbon flux toward E4P. After this, we integrated genes ylTKT, BbxfpK and AcxpkA at 26s rDNA site in strain YL24 with the helper plasmid prDNAloxP-ylTKT-BbxfpK-Acxpk, obtaining strain YL31. However, expression of the 2-PE pathway (ylPAR4-ylARO10-ylARO7-ylPHA2-scARO7^{G141S}) in strain YL31 gave almost same amount of 2-PE (strain YL32, 719.67 mg/L, Figure 4b) compared to strain YL30. We speculate that the engineered yeast chassis has sufficient E4P flux to maintain the shikimate pathway.

3.4. Removing glycolytic pathway bottleneck to boost precursor PEP

To shift the metabolic equilibrium and further improve carbon flux toward shikimate pathway, we moved to boost the precursor availability of PEP. Previous study ⁵⁸ found that blocking the reaction of PEP to pyruvate by deleting pyruvate kinase could dramatically increase the intracellular pools of PEP. To validate

this strategy in Y. lipolytica, we attempted to delete pyruvate kinase encoding gene ylPYK (YALI0F09185g). Reports found that yeast with pyruvate kinase deficiency could not grow in the complete synthetic media (CSM) with glucose as sole carbon source ⁵⁹. To successfully delete *ylPYK* (*YALI0F09185g*) in YL31 background chassis, we replenished 0.5 g/L of acetate in the selective CSM-plate (see Methods) to rescue the growth phenotype and obtained stain YL33. However, strain YL33 carrying the 2-PE pathway (ylPAR4-ylARO10-ylARO7-ylPHA2-scARO7^{G141S}) showed a remarkable decrease in both biomass and 2-PE production in the synthetic minimal media, even with the feeding of acetate ⁶⁰ (stain YL34, Supplementary Figure S5). Interestingly, when the same strain (YL33) was cultivated in YPD medium with 5 g/L sodium acetate, we observed robust cell growth (YL33 can also grow in YPD medium without acetate, as shown in Supplementary **Figure** S6). To this integrated 2-PE pathways end, we the (ylPAR4-ylARO10-ylARO7-ylPHA2-scARO7^{G141S}) into chassis **YL33** and cultivated YL33 with YPD medium containing 5 g/L sodium acetate (YPA medium). The generated strain YL35 significantly improved 2-PE production to 1572.70 mg/L in YPA medium (Figure 4c). The control strain YL36 (integration of 2-PE pathway ylPAR4-ylAR010-ylAR07-ylPHA2-scAR07^{G141S} in strain YL31) produced 1252.41 mg/L of 2-PE with YPA media under the same cultivation conditions. This result confirmed that blocking PEP consumption by deletion of ylPYK is a promising strategy for enlarging flux toward shikimate pathway.

Interestingly, the highest 2-PE titer of strain YL35 (1572.70 mg/L) was obtained at 72 h, and subsequently, 2-PE was gradually oxidized to PEA (phenylacetic acid), which reached 1481.8 mg/L at 120 h (Figure 4c). We speculate that the metabolic bottleneck may be shifted from E4P/PEP to NADH due to the exhaustion of glucose (Figure 4d): the engineered cells need to oxidize 2-PE to generate NADH and maintain metabolite homeostasis. To resolve this issue, feeding with 40 g/L of glucose effectively mitigated the synthesis of PEA (Figure 5e). As a result, we obtained a final 2-PE titer at 2426.22 mg/L (Figure 5e). To further improve 2-PE titer, we next sought to eliminate byproducts formation in strain YL35 by knocking out ylALD2, ylALD3, and ylHPD (encoding aldehyde dehydrogenases), resulting in strains YL37 and YL38. Deletion of genes ylALD2 and ylALD3 (strain YL37) did not significantly improve 2-PE production (2214.14 mg/L, Figure 4e), in addition, further deletion of gene ylHPD (strain YL38) led a remarkable decrease in both cell growth and 2-PE production. This is possibly due to the reversibility of ylHPD that contributes to the reduction of PEA to 2-PE.

In summary, we obtained the *Y. lipolytica* platform strain **YL33** with optimized metabolic flux toward shikimate pathway. This chassis strain was validated for *de novo* production of 2-PE at 2426.22 mg/L (strain **YL35**), a 286-fold increase over the initial strain (8.48 mg/L). This is the highest 2-PE titer from the *de novo* pathway reported to date. To demonstrate the utility of this 2-PE-producing platform, we turned to synthesize other aromatic-derived compounds, including *p*-coumaric acid (the derivative of tyrosine), resveratrol (the derivatives of *p*-coumaric acid) and violacein (the derivative of tryptophan) in the following section.

3.5. Extend the 2-PE platform for production of aromatic derivatives

In line with the optimization of 2-PE production, we decided to extend the yeast chassis to produce other aromatic derivatives, namely *p*-coumaric acid and violacein. *p*-Coumaric acid is the deamination product of tyrosine (Figure 5a), which is the universal precursor in the synthesis of flavonoids, aromatic polyketides, and lignin polyphenols ^{34, 61}. By simply overexpressing a codon-optimized tyrosine ammonia lyase *RgTAL* (originating from *Rhodotorula toruloides*) ³²and *ylTYR1* (that has been deleted in the platform strain) in the yeast chassis **YL33**, the resulting strain **YL39** produced 398.12 mg/L of *p*-coumaric acid with glucose supplementation (Figure 5b). However, 994.02 mg/L of 2-PE was detected, indicating a strong competing flux from the endogenous Ehrlich pathway. Thus, we next blocked 2-PE synthesis by deleting gene *ylPHA2* (encoding prephenate dehydratase), the obtained strain **YL40** produced 593.53 mg/L of *p*-coumaric acid with yield at 7.42 mg/g glucose (Figure 5b), a 1.49-fold increase compared to the control strain. We further extend the *p*-coumaric acid pathway for resveratrol synthesis by introducing genes *pc4CL* (4-coumarate-CoA ligase from *Petroselinum crispum*) and *VvSTS1* (resveratrol synthase from *Vitis vinifera*) ⁶². Unexpected, only 12.67 mg/L of resveratrol was detected in shake flasks (Supplementary Figure S7), indicating the existence of other rate-limiting precursors such as malonyl-CoA in the chassis strain.

Violacein is a naturally-occurring purple pigment with proven chemotherapeutic activity against tumors and cancers ^{63, 64}. Both violacein and its derivate deoxyviolacein have demonstrated a broad range of biological activities ⁶³. By introducing violacein biosynthetic genes (VioA, VioB, VioC, VioD, and VioE, Figure 5c), we previously obtained violacein production around 31 mg/L in *Y. lipolytica* ^{28, 65}. Surprisingly, overexpression of genes *ylTRP2*, *ylTRP3*, *VioA*, *VioB*, *VioC*, *VioD*, and *VioE* in strain **YL33** led to the accumulation of black pigment in the fermentation broth (Supplementary Figure S8). Violacein and deoxyviolacein reached 366.30 mg/L and 55.12 mg/L (Figure 5d), which were 2.88-fold and 2.51-fold

increase over the control strain (**po1fkV**, 126.99 mg/L of violacein and 21.98 mg/L of deoxyviolacein) and 5.23-fold and 10.44-fold higher than the production reported in our previous work ⁶⁵. To further improve the violacein and deoxyviolacein production, we enhanced tryptophan availability by overexpressing genes *ylTRP5*, *ylTRP4*, *ylTRP3*, *ylTRP2*, and *ylTRP1*. However, no significant improvements of violacein (348.43 mg/L) and deoxyviolacein (58.21 mg/L) titer were observed in strain **YL42**, suggesting that the supply of tryptophan was sufficient in strain **YL41** and there might be other limiting steps in the violacein pathway. Besides, we also attempted to block the chorismite-competing pathway by deleting *ylARO7*. No positive colonies were obtained in the background of strain **YL41** with several attempts, but *ylARO7* could be deleted in po1fk. The specific reason of our inability to delete *ylARO7* is not clear, which might be due to the metabolic burden of violacein accumulated in the cell negatively impacting cell fitness.

Future investigations of strains YL25, YL40, and YL42 should be performed under fed-batch cultivation with bench-top bioreactors and pilot-scale tests to improve the titer, yield and productivity of 2-PE, p-coumaric acid, and violacein. Major bioprocess considerations should be targeted at optimizing the culture medium, relieving 2-PE toxicity and promoting cell fitness, as well as reducing the separation and purification cost for 2-PE, p-coumaric acid, and violacein. Biphasic fermentation with end-product stripping might be an effective way to remove product inhibition. Considering that lignin is widely distributed in nature and it is made up with a variety of aromatic monomers, co-culture of Y. lipolytica with lignin-degradation microbes, including Pseudomonas putida and Rhodococcus jostii et al, with solid-state or liquid co-cultivation, might provide an attractive approach to valorize lignin and produce other complex aromatic derivatives.

In summary, by circumventing the intrinsic limitations of the endogenous shikimate pathway, we successfully engineered an oleaginous yeast chassis that produces 2426.22 mg/L of 2-PE, 593.53 mg/L of *p*-coumaric acid, 366.30 mg/L of violacein, and 55.12 mg/L of deoxyviolacein. To the best of our knowledge, this result represents the highest 2-PE, violacein and deoxyviolacein titer from the *de novo* shikimate pathway. This report highlights the prominent metabolic characteristics of *Y. lipolytica* as chassis for production of various aromatic derivatives.

4. Conclusions

Y. lipolytica is an oleaginous yeast with superior metabolic capability to produce a large portfolio of fuels, oleochemicals and natural products. In this work, we systematically overcame the rate-limiting steps of the

endogenous shikimate pathway and build a sustainable biorefinery chassis strain for de novo synthesis of

aromatics. We determined that relieving the feedback-inhibition of DAHP synthases is critical to channel flux

to shikimate pathway. We also demonstrated that eliminating by-product formation (L-phe, L-Tyr and L-Trp)

plays an important role in mitigating the feedback regulation of DAHP synthase. Overexpression of

phosphoketolase and deletion of pyruvate kinase provided a sustained metabolic source for E4P and PEP,

which forms the driving force to lead carbon flux through shikimate pathway. To demonstrate the utility of our

engineered Y. lipolytica chassis strain, three natural products, 2-PE, p-coumaric acid and violacein, which

were respectively derived from phenylalanine, tyrosine and tryptophan, were chosen to test the chassis

performance. With the engineered chassis Y. lipolytica, we obtained 2426.22 mg/L of 2-PE, 593.53 mg/L of

p-coumaric acid, 12.67 mg/L of resveratrol, 366.30 mg/L of violacein and 55.12 mg/L of deoxyviolacein.

These results highlight the metabolic versatility of Y. lipolytica and the native shikimate pathway could be

unlocked to produce a variety of aromatics and natural products with economic values.

Author contributions

PX conceived the topic and designed the study. YG performed genetic engineering and fermentation

experiments with input from JM and YL. GY and PX wrote the manuscript. PX revised the manuscript.

Conflicts of interests

A provisional patent has been filed based on the results of this study.

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Figure legends

Figure 1. De novo biosynthetic routes for 2-PE, p-coumaric acid, resveratrol and violacein production through the shikimate pathway in Y. lipolytica. PEP, phosphoenolpyruvate; E4P, erythrose-4P; DAHP, 3-deoxy-arabino-heptulonate-7-phosphate; 4HPP, 4-hydroxyphenylpyruvate; PAH, phenylacetaldehyde; 2-PE, 2-phenylethanol; ylTKT, transketolase; ylPYK, pyruvate kinase; ylARO3, 3-deoxy-7-phosphoheptulonate synthase; ylARO4, 3-deoxy-7-phosphoheptulonate synthase; ylARO5, 3-deoxy-7-phosphoheptulonate synthase; ylARO1, pentafunctional AROM polypeptide; ylARO2, chorismate synthase; ylARO7, chorismate anthranilate synthase; ylTRP3, anthranilate synthase; ylTRP4, phosphoribosyltransferase; ylTRP1, phosphoribosylanthranilate isomerase; ylTRP5, tryptophan synthase; ylTYP1, prephenate dehydrogenase; ylARO8, aromatic amino acid aminotransferase; ylARO9, aromatic amino acid aminotransferase; ylPHA2, prephenate dehydratase; ylARO10, phenylpyruvate decarboxylase; ylPAR4, phenylacetaldehyde reductases; ylHPD, 4-hydroxyphenylpyruvate dioxygenase; ylALD2, aldehyde dehydrogenase; ylALD3, aldehyde dehydrogenase; scARO7^{for}, the feedback-resistant chorismate mutase from S. cerevisiae; BbxfpK, phosphoketolase from Bifidobacterium breve; AcxpkA, phosphoketolase from Acidobacterium capsulatum; ecaroG^{fbr}, the feedback-resistant 3-deoxy-7-phosphoheptulonate synthase from E. coli; scARO4^{fbr}, the feedback-resistant 3-deoxy-7-phosphoheptulonate synthase from S. cerevisiae; rgTAL, tyrosineammonia-lyase from *Rhodotorula toruloides*; pc4CL, 4-coumarate-CoA ligase from *Petroselinum* crispum; VvSTS1, resveratrol synthase from Vitis vinifera; VioA, tryptophan oxidase; VioB/VioE, protodeoxyviolaceinate synthase; VioD, protodeoxyviolaceinate monooxygenase; VioC, violacein synthase. All violacein pathway genes were amplified from the genomic DNA of Chromobacterium violaceum. Right corner shows the chromosomal integration of the chosen pathways.

Figure 2. Validating and relieving bottlenecks in the Ehrlich pathway for 2-PE synthesis. (a) Enzyme cascade reactions of 2-PE production from chorismate; (b) Time profiles of 2-PE titer, cell growth, and phenylacetate titer of strains carrying the 2-PE pathway. All experiments were performed in triplicate and error bars represent standard deviations (SD).

Figure 3. Relieving bottlenecks in shikimate pathway. (a) The shikimate pathway of Y. lipolytica; (b)

Phenylacetate, 2-PE and cell growth profile of strains coexpressing ylARO1, ylARO2 and the

feedback-resistant DAHP synthases encoding genes; (c) Deletion of amino acids, including phenylalanine,

tryptophan, and tyrosine in Y. lipolytica; (d) Phenylacetate, 2-PE and cell growth profile of mutant strains

deficient in phenylalanine, tryptophan, and tyrosine formation; (e) Phenylacetate, 2-PE and cell growth profile

of mutant strains (deficient in phenylalanine, tryptophan, and tyrosine formation) expressing the shikimate

pathway. All experiments were performed in triplicate and error bars represent standard deviations (SD).

Figure 4. Rewiring carbon distribution toward precursors E4P and PEP. (a) Phosphoketolases provide

precursor E4P to drive shikimate pathway; (b) Time profiles of 2-PE, cell growth, and phenylacetate of strains

with chromosomal integration of transketolase ylTKT, phosphoketolases BbxfpK and AcxpkA at genomic

26s rDNA sites; (c) Time profile of 2-PE, cell growth, and phenylacetate of strains with pyruvate kinase

ylPYK deletion in YPD medium; (d) Time profiles of glucose consumption of strain YL35; (e) Improving

2-PE titer by feeding glucose and minimizing byproduct phenylacetate by deleting aldehyde dehydrogenase.

All experiments were performed in triplicate and error bars represent standard deviations (SD).

Figure 5. Harnessing Y. lipolytica chassis strain for p-coumaric acid and violacein production. (a) The de novo

pathway for p-coumaric acid synthesis; (b) Phenylacetate, cell growth, p-coumaric acid, 2-PE and glucose

consumption profile of the engineered chassis from the de novo pathway; (c) Violacein biosynthetic pathway

and yeast colonies with violacein accumulation; (d) Time profiles of violacein, deoxyviolacein and glucose

consumption of the violacein-producing strain. (e) Violacein cell culture harvested from synthetic minimal

media (strain po0fkV) and YPD media (YL41). All experiments were performed in triplicate and error bars

represent standard deviations (SD).

Figures

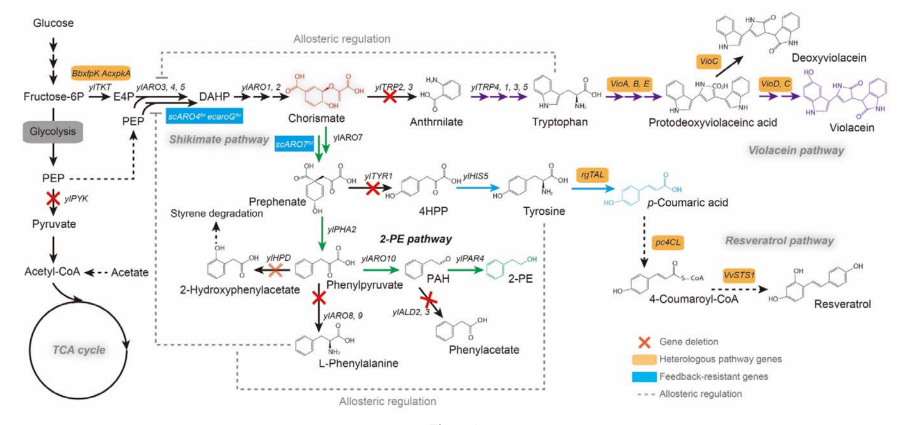


Figure 1

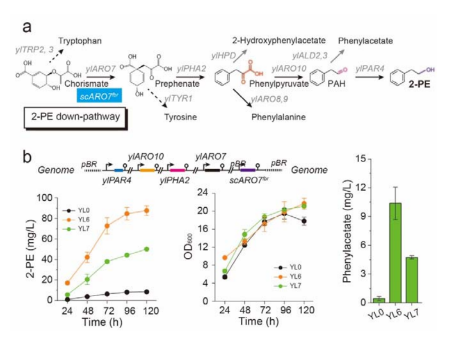


Figure 2

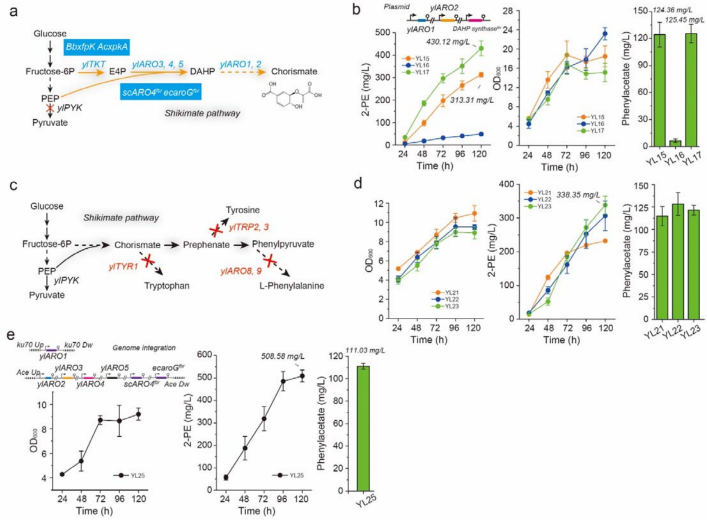


Figure 3

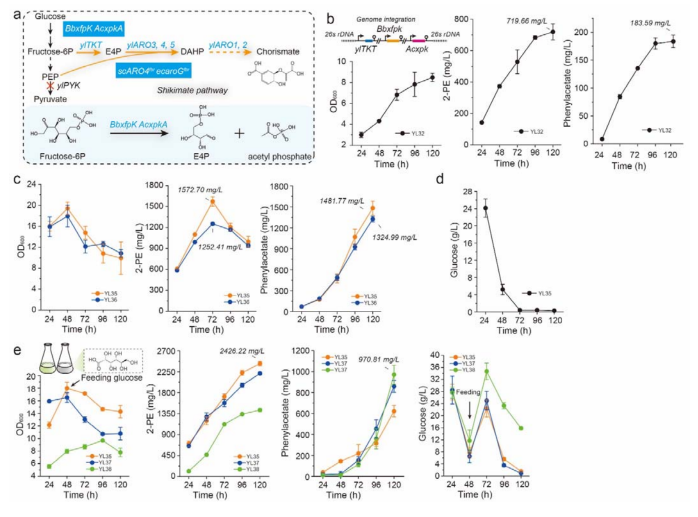


Figure 4

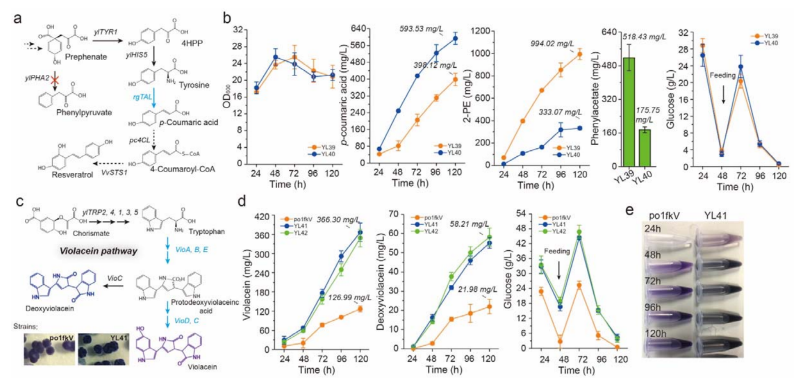


Figure 5