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Biosensor-assisted titratable CRISPRi high-throughput (BATCH) screening for over-production phenotypes

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Abstract

With rapid advances in the development of metabolic pathways and synthetic biology toolkits, a persisting challenge in microbial bioproduction is how to optimally rewire metabolic fluxes and accelerate the concomitant high-throughput phenotype screening. Here we developed a biosensor-assisted titratable CRISPRi high-throughput (BATCH) screening approach that combines a titratable mismatch CRISPR interference and a biosensor mediated screening for high-production phenotypes in *Escherichia coli*. We first developed a programmable mismatch CRISPRi that could afford multiple levels of interference efficacy with a one-pot sgRNA pool (a total of 16 variants for each target gene) harboring two consecutive random mismatches in the seed region of sgRNA spacers. The mismatch CRISPRi was demonstrated to enable almost a full range of gene knockdown when targeting different positions on genes. As a proof-of-principle demonstration of the BATCH screening system, we designed doubly mismatched sgRNA pools targeting 20 relevant genes in *E. coli* and optimized a PadR-based *p*-coumaric acid biosensor with broad dynamic range for the eGFP fluorescence guided high-production screening. Using sgRNA variants for the combinatorial knockdown of *ptkA* and *ptsI*, the *p*-coumaric acid titer was increased by 40.6% to 1308.6 mg/l from glycerol in shake flasks. To further demonstrate the general applicability of the BATCH screening system, we recruited a HpdR-based butyrate biosensor that facilitated the screening of *E. coli* strains achieving 19.0% and 25.2% increase of butyrate titer in shake flasks with sgRNA variants targeting *sucA* and *ldhA*, respectively. This work reported the establishment of a plug-and-play approach that enables multilevel modulation of metabolic fluxes and high-throughput screening of high-production phenotypes.

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Jian Wang and Chenyi Li contributed equally to this work.

Author contributions

J.W., C.L. and Y.Y. conceived the study; J.W., C.L., and T.J. conducted the experiments and associated analyses; Y.Y. supervised the research; J.W., C.L., and Y.Y. wrote and revised the manuscript.

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Competing interests

The authors declare that they have no conflicts of interest.

Keywords

Mismatch CRISPRi; Titratable repression; Biosensor; *p*-Coumaric acid; Butyrate

1. Introduction

Metabolic engineering harnesses the ability of living microorganisms to degrade renewable and low-cost carbon feedstocks in order to produce pharmaceutically or industrially important chemicals (Keasling, 2010; Khosla and Keasling, 2003; Lee et al., 2019; Nielsen and Keasling, 2011). However, the sophisticatedly regulated carbon metabolisms naturally coined in the genomes of most microbes always devote to cell survival and produce target chemicals non-efficiently (Montaño López et al., 2022; Zha et al., 2009; Zu et al., 2020). To obtain high-production strains that satisfy commercial relevance, traditional approaches rely on either domestication of microbes or deletion of specific non-essential competing genes (Gallone et al., 2016; Park et al., 2007; Steensels et al., 2019). With increasingly available genomic and transcriptomic data and better understanding of cellular metabolisms, increasing efforts have been made to systematically interrogate and rewire metabolic network in both “localized” and “global” scales (Andreozzi et al., 2016; Feist and Palsson, 2008; Lee et al., 2021; Park et al., 2009).

To that end, programmable synthetic biology toolkits have been developed to expedite genome-scale perturbation of metabolic fluxes, such as synthetic DNA oligonucleotides and CRISPR mediated multiplex genome engineering (Cong et al., 2013; Nyerges et al., 2016; Wang et al., 2009), retron-enabled genome editing (Lopez et al., 2022; Schubert et al., 2021; Simon et al., 2019), synthetic regulatory RNA mediated post-transcriptional interference (RNAi) (Na et al., 2013; Yang et al., 2015), and CRISPR-based transcriptional interference (CRISPRi) (Bikard et al., 2013; Gilbert et al., 2014; Hawkins et al., 2015). Among those, CRISPRi has been extensively used in systematic metabolic engineering due to its programmability and multiplexity, robust activity, and host flexibility (Banerjee et al., 2020; Liu et al., 2022; Schultenkämper et al., 2020; Wang et al., 2018b). However, CRISPRi mediated stringent gene repression may cause cell fitness defects especially when targeting growth-essential genes (Cui et al., 2018; de Bakker et al., 2022; Wang et al., 2021a), rendering this approach unable to thoroughly investigate the impact of expression levels of individual genes on desired high-production phenotypes.

To address this limitation, one ideal strategy would be to design massively pooled sgRNA libraries that guide dCas9 only to their respective gene targets while simultaneously affording parallel interrogation of individual genes on microbial production when repressed to different extents. CRISPRi repression efficiency could be modulated by engineered sgRNAs with random single mismatches (Jost et al., 2020; Qi et al., 2013), with interspersed double or multiple mismatches (Feng et al., 2021; Hawkins et al., 2020), with inducible hairpin or aptazyme-embedded structures (Ferry et al., 2017; Kocak et al., 2019; Siu and Chen, 2019; Tang et al., 2017), or with truncated spacers (Qi et al., 2013; Wang et al., 2021b). All these strategies require lab-intensive synthesis of large sgRNA libraries, empirically tested sgRNA structures, or specialized molecule inducers that have prevented

their practical applications in metabolic engineering practices. Moreover, concomitant with large-scale sgRNA libraries, one recurring challenge would be the lack of efficient and cost-effective high-throughput CRISPRi screening methods for desired production phenotypes.

In this work, we developed a biosensor-assisted titratable CRISPRi high-throughput (BATCH) screening approach that combines a titratable mismatch CRISPR interference and a biosensor mediated screening for high-production phenotypes in *Escherichia coli*. Based on the fact that either mismatch position, type or combination could impair CRISPRi activity, we first defined the rule for a cost-effective one-pot mismatch sgRNA design that enabled a full range of CRISPRi repression efficiencies. Then, to use *p*-coumaric acid and butyrate as two proof-of-principle applications, we recruited their respective PadR and HpdR based biosensor-reporter system with broad dynamic properties. With mismatch sgRNA libraries targeting genes involved in central carbon metabolisms, the BATCH screening system identified multiple target genes that enhanced titers of both products. This work established a readily applicable approach of high-throughput and multilevel CRISPRi screen in metabolic engineering to obtaining desired high-production phenotypes with programmable and customizable mismatch sgRNA libraries.

2. Materials and Methods

2.1 Strains, medium and chemicals

All strains and plasmids used in this study are listed in Table 1. *E. coli* XL1-Blue (Stratagene, La Jolla, CA) was used for plasmid construction and storage. *E. coli*:dCas9 was created by integrating the *P_{LacOI}*-dCas9 cassette into the genome at *dkgB* locus and was used as the host strain for CRISPRi test and butyrate production. Further deletion of *pheA* and *tyrR* in *E. coli*:dCas9 resulted in strain *E. coli*:dCas9 *pheA tyrR*, which was used as the host strain for *p*-coumaric acid production. Gene knockouts were conducted via P1 phage transduction method according to standard protocols (Thomason et al., 2007). Luria-Bertani (LB) medium (10 g/l tryptone, 5 g/l yeast extract and 10 g/l NaCl) was used for seed culture preparation. M9Y medium, prepared from M9 minimal medium (6 g/l Na₂HPO₄, 0.5 g/l NaCl, 3 g/l KH₂PO₄, 1 g/l NH₄Cl, 1 mM MgSO₄, 0.1 mM CaCl₂) containing 5 g/l yeast extract and 20 g/l glycerol or 20 g/l glucose, was used for shake flask experiments for producing *p*-coumaric acid or butyrate, respectively. Antibiotics including ampicillin (100 mg/ml) and kanamycin (50 mg/ml) were added to the medium at a ratio of 0.1% when necessary. Butyrate and *p*-coumaric acid standards were purchased from Sigma-Aldrich and MP Biomedical LLC, respectively.

2.2 Plasmid and strain construction

All plasmid constructions were carried out following standard molecular cloning protocols (Sambrook et al., 1989). Phusion high-fidelity DNA polymerase, restriction DNA endonucleases and Quick Ligation kit were purchased from New England Biolabs (Beverly, MA, USA). The high-copy plasmid pHA-MCS was derived from pZE12-luc by replacing *luc* gene with a multi-cloning site (MCS) and was applied as a negative control when needed. Fully matched sgRNA and rationally designed mismatch sgRNA variants targeting the eGFP coding sequence (starting from +39 bp) were constructed into the pCS-sgRNA

scaffold plasmid using with *Apa*LI and *Bam*HI, resulting in plasmids pCS-sgegfp-cds1, and pCS-sg-m7 to m16. Inverse PCR was applied to introduce random mismatches in sgRNA variants targeting genes of interest. To create mismatch sgRNA libraries targeting eGFP, the forward primer sgegfp-rbs-F (*Apa*LI) or sgegfp-cds2-F (*Apa*LI) was individually paired with the reverse primer sgRNA-R (*Apa*LI) (Table S1) to amplify the whole template plasmid pCS-sgRNA, resulting in linear plasmids containing sgRNA variants targeting the RBS region or the coding sequence (+378 bp) of *eGFP*, respectively. The resultant linear plasmids were then digested by *Apa*LI and *Dpn*I, and subjected to ligation. These ligation products were co-transferred into *E. coli*::dCas9 with the reporter plasmid pZE-eGFP. When plated on LB plates, transformants were randomly picked for eGFP fluorescence assay by cultivating in LB tubes and plasmids were extracted for gene sequencing analysis. To construct mismatch sgRNA variants targeting genes in *E. coli* genome for improved production of *p*-coumaric acid or butyrate, the inverse PCR templates were pCS-sgRNA and pZE-sgRNA, respectively. Selected targets and corresponding primers were listed in Table S1. Spacers of selected sgRNA variants that enabled high-producing phenotypes were later sequenced and included in Table S2.

For constructing *p*-coumaric acid responsive biosensor systems, the eGFP expression cassettes controlled by hybrid promoters (P1, P2, P7, and P9) were amplified from plasmids pZE-P1-eGFP, pZE-P2-eGFP, pZE-P7-eGFP, and pZE-P9-eGFP, respectively (Jiang et al., 2021). These cassettes were then inserted into plasmid pZE-Plpp0.2-PadR (Li et al., 2022) using *Aad*I and *Xho*I, resulting in plasmids pZE-Plpp0.2-PadR-P1-eGFP, pZE-Plpp0.2-PadR-P2-eGFP, pZE-Plpp0.2-PadR-P7-eGFP, and pZE-Plpp0.2-PadR-P9-eGFP. The *P_LJacOI*-controlled RgTAL expression cassette was amplified from plasmid pZE-RgTAL (Huang et al., 2013) and was constructed to plasmid pZE-Plpp0.2-PadR-P9-eGFP using *Bam*HI and *Avr*II, yielding plasmid pZE-CA. The biosynthetic pathway of butyrate consisting of *thl*, *hbd*, *crt*, *ter*, and *atoDA*, were sequentially constructed to plasmid pCS27 to form plasmid pCS-THCTA using *Acc*65I, *Nde*I, *Sal*I, *Eco*RI, *Bam*HI, and *Mlu*I. The Plpp1-HpdR-P_{hpdH}-eGFP (HPE) biosensor cassette was amplified from plasmid pCS-V2 (Wang et al., 2021b) and constructed to plasmid pCS-THCTA using *Xho*I and *Xba*I, yielding pCS-HPE-THCTA.

2.3 Biosensor dynamic range test and eGFP repression assay

To test the dynamic range or substrate scope of engineered biosensor systems, plasmids harboring the biosensor systems were transferred into *E. coli* BW25113(F'). Three independent colonies of each testing group were randomly picked and inoculated into 3.5 ml fresh LB medium (with appropriate antibiotics) as seed cultures. After overnight cultivation (10–12 h) in a 37°C rotatory shaker at a speed of 270 rpm, seed cultures were transferred into fresh 3.5 ml LB medium with a ratio of 5% (175 µl), and new cultures were placed in the shaker under 37°C with a speed of 270 rpm. After 1 h of cultivation, gradient concentrations of corresponding substrates (0–1000 mg/L for *p*-coumaric acid, and 0–2.5 g/L for butyrate) were fed into the cultures. After 12 h of cultivation post substrate induction, cultures were sampled and diluted by 10-folds before subjected to measurement of cell density (OD₆₀₀) and green fluorescence intensities using the 96-well plate reader (Synergy HT, BioTek). The eGFP repression assay was conducted with *E.*

coli::dCas9 co-transformed with pZE-eGFP and pCS27 harboring sgRNAs. *E. coli*::dCas9 co-transformed with pZE-eGFP and pCS27 empty plasmid was used as a control. The relative eGFP expression is calculated as the ratio of normalized eGFP fluorescence per OD₆₀₀ (RFU/OD₆₀₀) of cells with sgRNAs to those without sgRNAs. Green fluorescence intensities were determined using an excitation filter of 485/20 nm and an emission filter of 528/20 nm.

2.4 Screening high-production phenotypes in test tubes

For screening of *p*-coumaric acid high-production phenotypes, constructed mismatch sgRNA libraries on pCS27 for each target gene were separately co-transferred into *E. coli*::dCas9 *pheA tyrR* with plasmid pZE-CA. For screening of butyrate high-production phenotypes, constructed mismatched sgRNA libraries on pZE12-luc for each target gene were separately co-transferred into *E. coli*::dCas9 with plasmid pCS-HPE-THCTA. All transformants were plated on LB plates with appropriate antibiotics and 0.5 mM IPTG. Colonies with high green fluorescence intensities under blue light exposure were picked and inoculated in 3.5 ml M9Y medium containing 20 g/l glycerol (for *p*-coumaric acid production) or 20 g/l glucose (for butyrate production). IPTG was added in the medium to a final concentration of 0.5 mM at the beginning to induce expression of CRISPRi system and biosynthesis pathways. Samples were taken after 24 h of induction and were subjected to measurement of cell densities, green fluorescence intensities, and product accumulations. Cell density and green fluorescence intensity were quantitatively analyzed using the 96-well plate reader (Synergy HT, BioTek) with the same protocol described earlier. Production of *p*-coumaric acid or butyrate was also analyzed by high-performance liquid chromatography (HPLC). Selected high-production strains were streaked in LB plates (with appropriate antibiotics) and plasmids were extracted for DNA sequencing of variant sgRNA spacers.

2.5 Validation of selected sgRNA variants via shake flasks fermentation

To validate the production performance of selected colonies from test tubes, three colonies of the strains streaked on LB plates were inoculated in 3.5 ml LB medium and grown at 37 °C for 8–10 h. Then, 1 ml of each seed culture was transferred to 20 ml fresh M9Y medium (containing 20 g/l glycerol or glucose) in 125-ml shake flasks (Chemglass Life Sciences LLC) and grown at 30 °C for 48 h. Shake flask experiments were performed in a 30 °C rotary shaker with a speed of 270 rpm. Glycerol was used as carbon source for *p*-coumaric acid production while glucose was used for butyrate production. IPTG was added in the medium with a final concentration of 0.5 mM. Samples were collected after 48 h post induction and were subjected for measurement of cell density (OD₆₀₀) and product accumulation. Cell density (OD₆₀₀) was measured using the spectrophotometer (VWR), and the products *p*-coumaric acid or butyrate were analyzed by HPLC.

2.6 Product analysis

Collected samples (1 ml) were centrifuged at 12,000 rpm for 15 minutes, and supernatants were filtered through 0.22 µm film before applied to HPLC analysis. Concentrations of *p*-coumaric acid were quantitatively analyzed by Agilent HPLC 1260 Infinity II (1260 Infinity II Diode Array Detector WR) with a reverse-phase ZORBAX SB-C18 column. A methanol-water (containing 0.1% trifluoroacetic acid) gradient system with a flow rate of 1

ml/min was used. The analyzing method, adopted from our previous study (Li et al., 2022), was set as follows: 5% methanol from 0 to 2 min, 5% to 80% methanol from 2 to 10 min, 80% methanol from 10 to 14 min, 80% to 5% methanol from 14 to 18 min, and 5% methanol from 18 to 20 min. A standard curve was drawn using gradient concentrations of *p*-coumaric acid solution prepared from *p*-coumaric acid standard (MP Biomedical LLC). The retention time of *p*-coumaric acid was 8.89 min and the detection wavelength was set at 280 nm.

Butyrate was analyzed by a Dionex Ultimate 3000 HPLC equipped with a Coregel-64H column (Transgenomic, Omaha, NE). The collected samples (1 ml) were centrifuged at 12,000 rpm for 15 minutes, and the supernatants were filtered through 0.22 μ m film before applied to HPLC analysis. The HPLC method was modified from our previous research (Wang et al., 2018a). Briefly, 4 mN sulfuric acid was used as the mobile phase with a flow rate of 0.4 ml/min. The column temperature was set at 45 °C. The retention time of butyrate was 38.897 min.

3. Results

3.1 Design of BATCH screening system

The BATCH screening system is composed of two critical parts: (1) a titratable CRISPRi for customizable and multilevel gene repression in microbial chassis, and (2) chemical inducible biosensors with broad dynamic ranges for phenotype screening (Fig. 1). The titratable CRISPRi represses all relevant gene targets within the metabolic network or competing genes of target biosynthetic pathways to different extents. Biosensors are committed to sensing intracellular product chemicals and inducing the reporter gene eGFP expression, thus linking the product concentration with fluorescence intensity. Implementation of the BATCH screening system involves two rounds of selection: the first round selection of fluorescent single colonies on plates for test tube cultivation and the second round selection of highly fluorescent transformants from test tube cultures for shake flask cultivation validation. A combination of selected beneficial repressions could be iteratively launched to screen hyperproduction phenotypes.

3.2 Establishing one-pot mismatch sgRNA libraries for titratable CRISPRi

First, in order to establish a titratable CRISPRi, we sought to alter the repression strengths of target genes simply by engineering customizable sgRNA libraries. Since CRISPRi activity is highly sensitive to mismatch positions, types and combinations between sgRNA and target DNA (Feng et al., 2021; Gilbert et al., 2014), we herein endeavored to establish a titratable CRISPRi method with a facile one-pot mismatch sgRNA library design.

To facilitate CRISPRi system construction and also avoid a potential toxicity of SpdCas9 overexpression in *E. coli* cells (Calvo-Villamañán et al., 2020; Zhang and Voigt, 2018), we constructed a strain *E. coli*:dCas9 carrying SpdCas9 under the control of *P_{LacO1}* promoter integrated into the genome at the low-expression *dkgB* locus (Bryant et al., 2014; Wang et al., 2021b). To generate mismatch sgRNAs, inverse PCR was applied for introducing mutations in sgRNA spacers with degenerate primers (Fig. 2A). We first

tested the single mismatch sensitivity of sgRNA on eGFP repression, by transforming *E. coli*:dCas9 with pCS-eGFP and pZE12-sgRNAs harboring mismatch sgRNAs targeting 5' coding sequence (+39 bp) of eGFP (Fig. 2B and C). The eGFP repression assay indicated that single mismatches outside a 7bp PAM-proximal seed position are tolerated without significantly affecting CRISPRi repression efficiencies (Fig. 2C). This was consistent with previous findings that mismatches within the generally recognized ~8bp seed region greatly compromised or abolished the CRISPR/(d)Cas9 functionality, at least in *E. coli* (Datsenko et al., 2012; Gilbert et al., 2014; Qi et al., 2013; Sternberg et al., 2015). To further investigate the sensitivity of double mismatches, we narrowed down to construct sgRNA variants harboring saturated double substitutions (NN) in PAM-proximal 7–8th bp seed region. The repression assay showed that the 16 possible sgRNA variants almost enabled a broad repression profile of eGFP from 14.6% to 95.6%, suggesting that mismatching the 7–8th bp seed region could afford a multilevel gene repression (Fig. 2D). To examine the impact of target positions on the repression profile, we therefore designed double mismatch sgRNAs with random mutations of the 7–8th bp seed region that targeted both ribosome binding site (RBS) and alternate coding sequence (+378 bp) of eGFP with degenerate primers. In both cases, out of 32 randomly picked transformants, 12 different mismatch sgRNAs were obtained, accounting for a coverage of 75% of total variants. Both sgRNA variants enabled a full range of knockdown levels of eGFP (Fig. 2E and F), further confirming the generable applicability of one-pot mismatch sgRNA libraries for programmable and titratable CRISPRi.

3.3 Optimizing the *p*-coumaric acid biosensor PadR with broad dynamic behavior

For a proof-of-principle demonstration, we chose to target the *p*-coumaric acid biosynthetic pathway and optimized a PadR-based *p*-coumaric acid biosensor system for its high-production screening from glycerol. *p*-Coumaric acid has extensive applications in food, pharmaceutical and cosmetic industry, and also serves as an important precursor for more complex natural products such as flavonoids and polyphenols (Trantas et al., 2015; Wang et al., 2016).

To assist high-production phenotype screening, natural or engineered chemical-inducible biosensors with broad dynamic range would be critical. An engineered *p*-coumaric acid-responsive biosensor system PadR- P_{padC} from *Bacillus subtilis* was established in our previous study (Jiang et al., 2021). The transcriptional repressor PadR could repress the expression of downstream gene controlled by P_{padC} promoter, while the presence of *p*-coumaric acid could activate its transcription initiation (Jiang et al., 2021). To enable biosensor-assisted screening, four previously engineered hybrid P_{padC} promoters (P1, P2, P7, and P9) with optimized dynamic ranges and strong output strengths were selected to pair with PadR (Table S3). The hybrid promoters were constructed by inserting PadR binding boxes into the strong constitutive pL promoter (Jiang et al., 2021). PadR under control of the constitutive Plpp0.2 promoter and eGFP under control of the hybrid promoters were assembled into pZE12-luc, resulting pZE-Plpp0.2-PadR-P1-eGFP, pZE-Plpp0.2-PadR-P2-eGFP, pZE-Plpp0.2-PadR-P7-eGFP, and pZE-Plpp0.2-PadR-P9-eGFP (Fig. 3A). To test the dynamic performance, all plasmids harboring biosensor systems were transferred into *E. coli* BW25113(F'), and transformants were cultivated with supplementation of

gradient concentrations of *p*-coumaric acid (0, 100, 200, 400, 600, and 1000 mg/l). The PadR-P9 biosensor system exhibited the best performance in the selected concentration range, resulting in a 14.2-fold eGFP induction with 1000 mg/l *p*-coumaric acid (Fig. 3B). Therefore, this combination was selected for the following BATCH screening of *p*-coumaric acid high-production phenotypes.

3.4 BATCH screening of *p*-coumaric acid over-production phenotypes

The *p*-coumaric acid pathway initiates from the aromatic amino acid (AAA) pathway and ends up with deamination of tyrosine to *p*-coumaric acid via tyrosine ammonia-lyase (TAL) (Fig. 4A). Extending from the central carbon metabolism, the *p*-coumaric acid pathway suffers from multiple potential competing pathways that divert carbon fluxes. To redirect carbon flux to the *p*-coumaric acid pathway, we designed random double mismatch sgRNAs targeting 20 relevant genes in the glycolysis pathway (*ptsI*, *pgi*, *pfkA*, *fbaA*, *phoA*, *gapA*, *pgk*, *gpmA*, *gpmM*, *eno*, *pykA*, *pykF*, and *csrA*), the TCA cycle (*aceE*, *gltA*, and *ppc*), the fatty acid biosynthesis pathway (*accA*, and *fabD*), and the AAA pathway (*ubiC*, and *trpE*) (Fig. 4A). To minimize deleterious effects on cell growth and customize sgRNA design, sgRNAs targeted available NGG PAMs near the start codon for each gene target to block transcription elongation.

To implement the BATCH screening for *p*-coumaric acid over-production phenotype, *E. coli*:dCas9 *tyrR pheA* with knockouts of the tyrosine repressor (*tyrR*) and the chorismate mutase/prephenate dehydratase (*pheA*) was applied as the host. *E. coli*:dCas9 *tyrR pheA* was co-transformed with plasmids pZE-CA (harboring RgTAL and PadR biosensor) and pCS-sgRNAs harboring mismatch sgRNA libraries targeting each gene (Fig. 4B). After transformation and growing on LB plates supplemented with 20 g/l glycerol and 0.5 mM IPTG, a total of 284 colonies with notably high green fluorescence intensities were picked and cultivated in test tubes with M9 minimal medium containing 20 g/l glycerol (Fig. S1). Determination of eGFP fluorescence intensities and *p*-coumaric acid concentrations of all selected transformants exhibited a moderate positive correlation (Pearson correlation coefficient $r = 0.34$) (Fig. 4B). For each target gene, the difference of eGFP fluorescence intensities and *p*-coumaric acid concentration among mismatch sgRNA variants underlined the impact of gene repression level on *p*-coumaric acid production (Fig. S1). Among all targets, sgRNA variants towards *csrA*, *fbaA*, *gapA*, *pfkA*, *phoA*, *ppc*, *ptsI*, and *pykF* produced more hits with increased *p*-coumaric acid titers (Fig. S1). Shake flask based production tests with selected colonies from the test tube screening validated that mismatch sgRNAs towards *csrA*, *ptsI*, *pykF*, *ppc*, and *pfkA* afforded more than 25% increase of *p*-coumaric acid titers (Fig. 4C, and Fig. S1). The overall yields of *p*-coumaric acid per biomass were generally improved in accordance with *p*-coumaric acid titers (Fig. S2A). Especially, compared with the control strain (930.6 mg/l and an overall yield of 0.09 g/l/OD600), sgpfkA-3 permitted the highest production of *p*-coumaric acid, reaching a peak titer of 1232.0 mg/l and an overall yield of 0.13 g/l/OD600 (Fig. 4C and Fig. S2A).

One notable advantage of mismatch sgRNAs in BATCH screening over authentic sgRNAs in conventional CRISPRi is its non-deleterious effect to cell fitness especially when targeting essential genes. To corroborate that, we chose to compare the functionality of

mismatch sgRNAs and authentic sgRNAs towards growth-essential genes such as *ppc*, *ptsI* and *csrA*. As expected, while authentic sgRNAs significantly impaired both cell growth (approximately 50% decrease) and *p*-coumaric acid titer, selected sgRNA variants improved *p*-coumaric acid production respectively by 28.4%, 25.0% and 24.6%, without sacrificing cell growth (Fig. 4D). This further underlined that the conditional and optimal gene repression with the BATCH screening system would expedite the rational high-production phenotype screening. One other advantage of BATCH screening is applying it for easy testing of combinatorial effects of beneficial repressions from initial rounds. Therefore, we combined selected mismatch sgRNAs to further boost *p*-coumaric acid production. The combination of *sgpfkA-3* and *sgptsI-5* increased *p*-coumaric acid titer to 1308.6 mg/l, accounting for a 40.6% increase over the control strain (Fig. 4E).

3.5 Application of the BATCH system for screening of butyrate over-production phenotypes

To explore the applicability of the BATCH screening system to other production scenarios, we then implemented it to the screening of butyrate over-production phenotypes. Butyrate is a C4 fatty acid and a commodity chemical with wide applications in chemical industry, food manufacturing, pharmaceuticals, perfume, and animal feed supplements (Guo et al., 2021; Jang et al., 2014). The butyrate pathway initiates from condensation of two acetyl-CoA to acetoacetyl-CoA followed by its reduction to 3-hydroxybutyryl-CoA with the enzyme thiolase (*thl*) and hydroxybutyryl-CoA dehydrogenase (*hbd*) (Fig. 5A). 3-Hydroxybutyryl-CoA was then reduced to butyryl-CoA by crotonase (*crt*) and transenoyl-CoA reductase (*ter*), followed by release of butyrate from butyryl-CoA with acetyl-CoA:acetoacetyl-CoA synthase (*atoDA*) (Choi et al., 2012; Saini et al., 2014). As the direct precursor for butyrate and also a critical central metabolite, increasing acetyl-CoA supply or reducing its consumption could enhance butyrate titers.

To deploy the BATCH screening, we first sought to identify and characterize a butyrate biosensor with broad dynamic range. We previously engineered a promiscuous 3-hydroxybutyrate responsive HpdR-P_{hpdH} system on pCS-V2 plasmid, which harbors a deletion of a palindromic sequence (−209 to −118 bp) on the P_{hpdH} promoter and expresses HpdR from *Pseudomonas putida* under control of Plpp1.0 promoter (Hanko et al., 2017; Wang et al., 2021b). To investigate the ligand scope of the engineered HpdR-P_{hpdH} system, *E. coli* BW25113(F′) transformed with pCS-V2 was subjected to C3-C5 fatty acids, hydroxy acids, and diacids with gradient concentrations (0–2.5 g/l). The eGFP fluorescence assay showed that the engineered HpdR-P_{hpdH} system exerted high sensitivity and broad dynamic range towards all C3-C5 fatty acids (propionate, butyrate, and valerate) and hydroxy acids (3-hydroxypropionate, 4-hydroxybutyrate, and 5-hydroxyvalerate) but not diacids (Fig. 5B). Among all ligands, butyrate afforded the highest eGFP induction fold (41-fold at 2.5 g/l) (Fig. 5B), indicating that the engineered HpdR-P_{hpdH} system could serve as an ideal butyrate biosensor.

To experimentally implement the BATCH screening, the engineered HpdR-P_{hpdH} system was assembled with the butyrate pathway, resulting in plasmid pCS-HPE-THCTA (Fig. 5C). To reserve acetyl-CoA for butyrate biosynthesis, 14 mismatch sgRNA libraries

involved in pyruvate consumption (*ldhA*), acetyl-CoA consumption (*ackA*, and *adhE*), fatty acid pathway (*accA* and *fabD*), TCA cycle (*gltA*, *acnA*, *acnB*, *icd*, *sucA*, *sucC*, *sdhC*, *sdhD*, *fumA* and *mdh*) were constructed into pZE12-luc (resultant pZE-sgRNAs). All mismatch sgRNAs harboring 7–8th random mutations targeted available NGG PAMs near the start codon for each gene target to block transcription elongation. *E. coli*:dCas9 was co-transformed with pCS-HPE-THCTA and pZE-sgRNAs and subjected to BATCH screening. For over 150 variants screened in test tubes, eGFP fluorescence intensities and butyrate titers were strongly correlated (Pearson correlation coefficient $r = 0.69$), confirming the applicability of biosensor-based screening of butyrate over-production phenotypes (Fig. 5D, and Fig. S3). The initial screening underlined that sgRNA variants targeting each gene enabled differential butyrate titers, and sgRNAs targeting *acnB*, *ldhA*, *sucA* and *sucC* generally afforded more sgRNA variants with enhanced butyrate titers (Fig. S3). Further validation of 27 selected over-production colonies in shake flask cultivations showed that the best sgRNA variants *sgsucA-7* and *sgldhA-10* increased butyrate titer by 19.0% and 25.2% over the control strain (2.90 g/l), reaching 3.45 and 3.63 g/l, respectively (Fig. 5E). Interestingly, the overall yields of butyrate per biomass were not necessarily improved, as we observed comparable overall yields per biomass between *sgldhA-10* (0.49 g/l/OD600) and the control strain (0.48 g/l/OD600) (Fig. S2B). These results together demonstrated the general applicability of BATCH screening in different production scenarios.

4. Discussion

Identification of competing gene targets and optimal control of their expression are critical in achieving high production of desired biochemicals in metabolic engineering (Ko et al., 2020; Na et al., 2013). Due to the intricately regulated nature and the essentiality in cellular maintenance of metabolic network, conventional gene knockout strategies are often labor intensive and would miss identification of potential targets. RNAi or CRISPRi based screening has expedited large-scale profiling and repression of competing gene targets without modifying genomes. However, the static gene repression imposed by RNAi or CRISPRi is sometimes suboptimal and even detrimental to metabolic pathways in production scenarios (Crook et al., 2016; Rousset et al., 2018). This work established a superior BATCH screening system that renders (1) titratable CRISPRi using customizable mismatch sgRNA libraries for full-range gene repression, and (2) pinpointing the optimal gene repression with product biosensor based high-production screening.

The sensitivity of CRISPRi towards mismatches enabled fine-tuning of CRISPRi repression efficiency. In this work, we first located the mismatching window to the 7–8th bp within the seed region of sgRNA, whose random mutation almost afforded a full-scale of eGFP repression regardless of targeting positions (Fig. 2). With designable degenerate primers, mismatches could be easily created with one-pot PCR and incorporated into sgRNA plasmid libraries, as indicated by the 75% coverage from 32 randomly picked colonies (Fig. 2). These two features add greatly to the general applicability and manipulation simplicity of the BATCH screening system in metabolic engineering practices. Deployment of the BATCH system in *p*-coumaric acid production validated our hypothesis that repressing gene targets at different extents could dynamically affect cell fitness and product formation. As demonstrated by repressing growth-essential genes including *ppc*, *ptsI* and *csrA*, repression

of all three genes with authentic sgRNAs significantly impaired cell growth and *p*-coumaric acid production, while mismatch sgRNAs marginally affected cell growth but significantly increased *p*-coumaric acid production (Fig. 4D). Thus, the BATCH screening system allows the production based screening of competing gene targets and their optimal repressions in production scenarios.

The BATCH screening system relies on the biosensors of target chemicals with broad dynamic ranges. Chemical-inducible biosensors have been widely used to accelerate high-production screening and implemented into dynamic control paradigms for carbon metabolism rewiring (Seok et al., 2021; Zhang et al., 2015). However, the lack of biosensors or poor dynamic performance always limited their wide applications. In this study, we recruited two engineered biosensors, the PadR- P_{padC} system for *p*-coumaric acid and the HpdR- P_{hpdH} system for butyrate. When placed on high-copy plasmid, while the native PadR- P_{padC} (P1) showed low-induction fold (3.9-fold) and eGFP expression even in the presence of 1 g/l *p*-coumaric acid, the engineered PadR- P_{padC} (P9) with mutations in P_{padC} promoter showed markedly increased induction fold (14.2-fold) and elevated eGFP expression (Fig. 3). Biosensors for short-chain fatty acids (C3-C5) are rarely documented. We previously engineered a promiscuous 3-hydroxypropionate-responsive HpdR- P_{hpdH} system with significantly improved sensitivity and dynamic range towards 3-hydroxybutyrate (Wang et al., 2021b). Ligand profiling substantiated that the engineered HpdR- P_{hpdH} system could actually accept all C3-C5 fatty acids and counterpart hydroxy acids including butyrate (Fig. 5B). With broad dynamic ranges, both the engineered PadR- P_{padC} and HpdR- P_{hpdH} system rendered modest to strong positive correlations between eGFP fluorescence and product titers, which successfully guided the screening of their high-production *E. coli* cells. These chemical-inducible biosensors demonstrated here in phenotype screening could also be readily translated into other applications like sensor-mediated dynamic gene regulations.

It is noteworthy that mismatch positions, types or numbers within sgRNA spacers would have an impact on CRISPRi repression efficiencies. Thus, it is feasible to create diverse mismatch sgRNAs to achieve more fine-tuned gene repression. For instance, introducing more mismatches would create larger sgRNA libraries (4^n , n is the number of mismatches) for multilevel gene repression. Targeting different locations on genes (such as promoter, RBS, or coding sequence) by mismatched sgRNAs would also influence the CRISPRi repression range. Thus, it is possible to expand sgRNA libraries by targeting different locations on gene and potentially enable more beneficial variants. However, expanded sgRNA libraries would require screening of larger colony libraries to cover most of the potential sgRNA variants and would be labor-intensive without biosensor-based or other high-throughput screening technologies. To customize sgRNA design, we herein demonstrated the applicability of doubly mismatched sgRNA libraries targeting locations near start codons of gene targets to titrate gene expression and improve production performance.

Combination of multiple beneficial sgRNAs did not always result in the expected synergetic effect in production improvement. This underlined that individually modulated transcription level of each target gene may not be optimal in production scenario when combined. A

potential solution to address this issue would be to screen multiple gene targets in a single round, which will be feasible with proper designs and appropriate DNA assembly methods. Given the multiplexity nature of CRISPRi, BATCH screening of multiple gene targets can be performed to expand the library size and potentially obtain combinatorial sgRNA variants with more desired high-producing phenotypes. We also noticed the variations of production improvement between test tube screenings and shake flask validations, which were mostly due to different cultivation conditions. This underscored that all obtained over-production strains from first round screening needed to be further confirmed in shake flask cultivations.

In conclusion, we developed a new BATCH screening system that could be readily applied for large-scale target identification and multilevel gene repression with one-pot mismatch sgRNA libraries. We demonstrated its applicability to identify multiple gene targets for production improvement of *p*-coumaric acid and butyrate within a single round of selection. Ideally, efficient assembly of sgRNA variants targeting multiple potential genes could also allow for the investigation of the combinatorial effects with the BATCH screening system. In summary, the screening system presented here would expedite the development of high-production microbial strains without genome intervention.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Data and materials availability

All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Additional data or materials related to this paper may be requested from the corresponding author.

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Highlights

- A biosensor-assisted titratable CRISPRi high-throughput screening was established.
- One-pot mismatch sgRNA libraries could enable a full range of CRISPRi efficacy.
- Biosensors with broad dynamic ranges allowed reporter-based phenotype screening.
- Multiple gene targets were identified to enhance *p*-coumaric acid and butyrate titers.

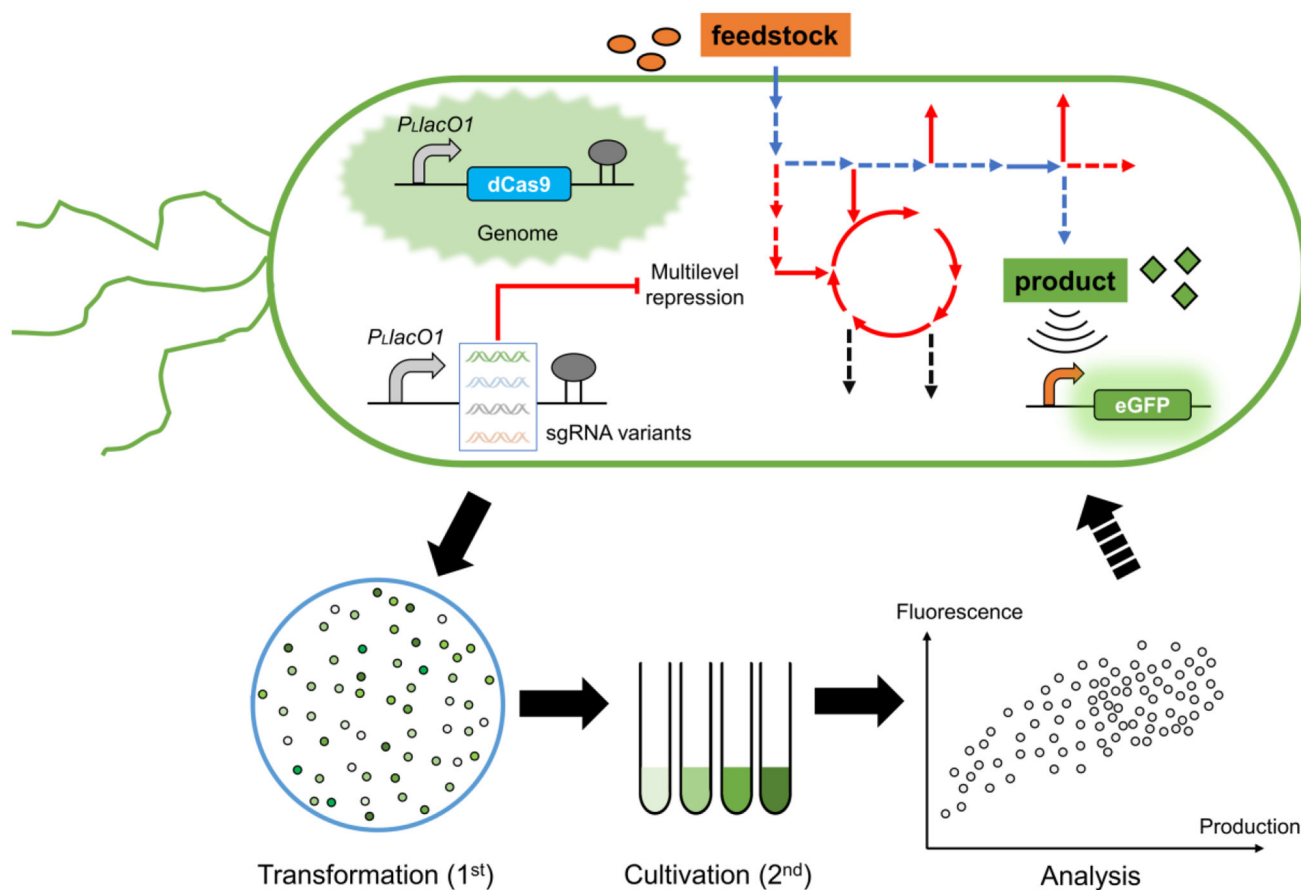


Figure 1. Schematic diagram of the BATCH screening workflow.

The BATCH screening involves *sgRNA* variants mediated multilevel CRISPRi of genes of interest within metabolic network to redirect carbon flux to target biosynthetic pathways. Improved production phenotypes will be reflected by the target chemical inducible reporter GFP fluorescence. The BATCH screening consists of a first round plate based selection and a second round test tube based selection followed by final shake flask based validation. Blue lines indicate biosynthetic pathways and red lines indicate relevant or potential competing pathways.

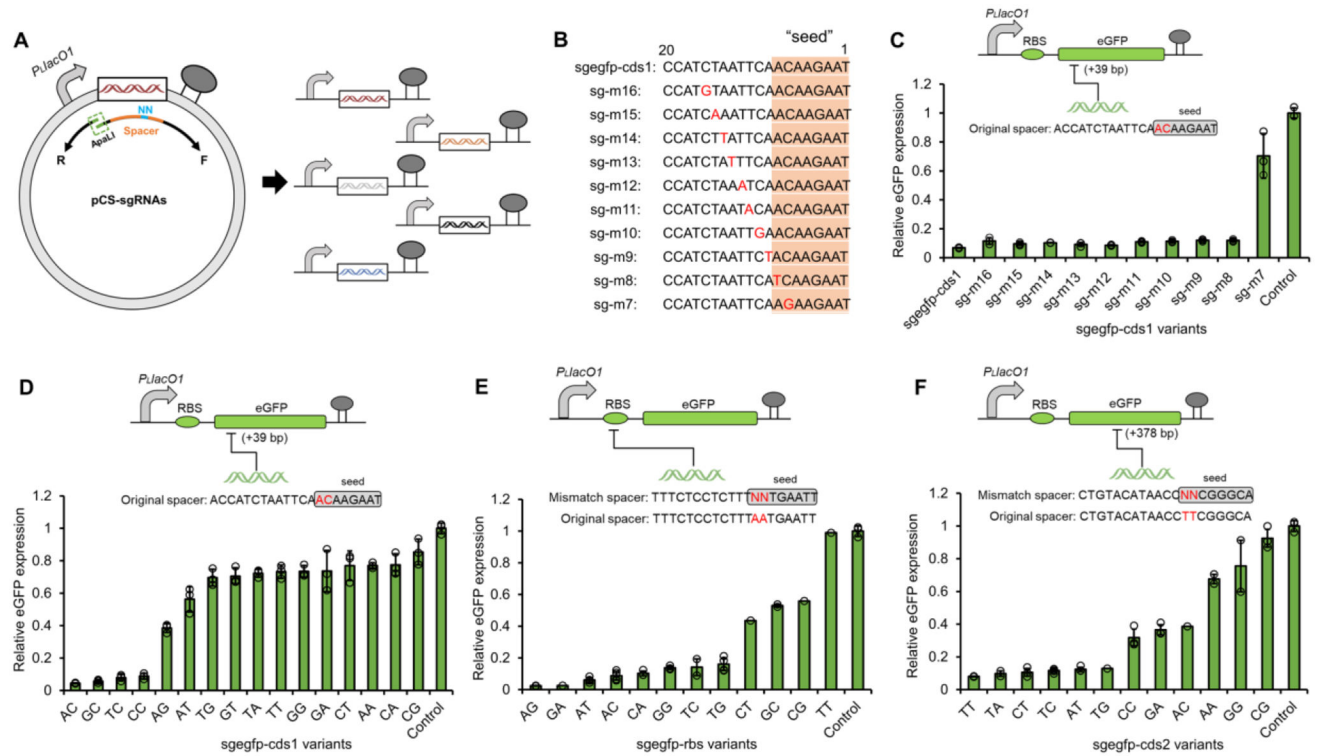


Figure 2. Establishing titratable CRISPRi with mismatch sgRNA libraries.

(A) The design of inverse PCR for one-pot construction of mismatch sgRNAs. (B) Spacer sequences (5'-3') of single mismatch sgegfp-cds1 variants targeting the coding sequence (+39 bp) of eGFP. (C) The CRISPRi efficiencies of sgRNAs with single mismatch at different positions in the sgegfp-cds1 spacer (c), with rationally designed double mismatches in the 7–8th bp seed region of sgegfp-cds1 spacer (D), with randomly designed double mismatches in the 7–8th bp seed region of sgegfp-rbs spacer (targeting the RBS region eGFP promoter) (E), and with randomly designed double mismatches in the 7–8th bp seed region of sgegfp-cds2 spacer (targeting the coding sequence (+378 bp) of eGFP) (F). Control, *E. coli*::dCas9 co-transformed with pZE-eGFP and pCS27 empty plasmid. In E and F, 32 single colonies were randomly picked from each plate and were subjected to both eGFP fluorescence assay and spacer determination via plasmid sequencing. All data in C and D represent the mean of three biologically independent samples and error bars show standard deviation.

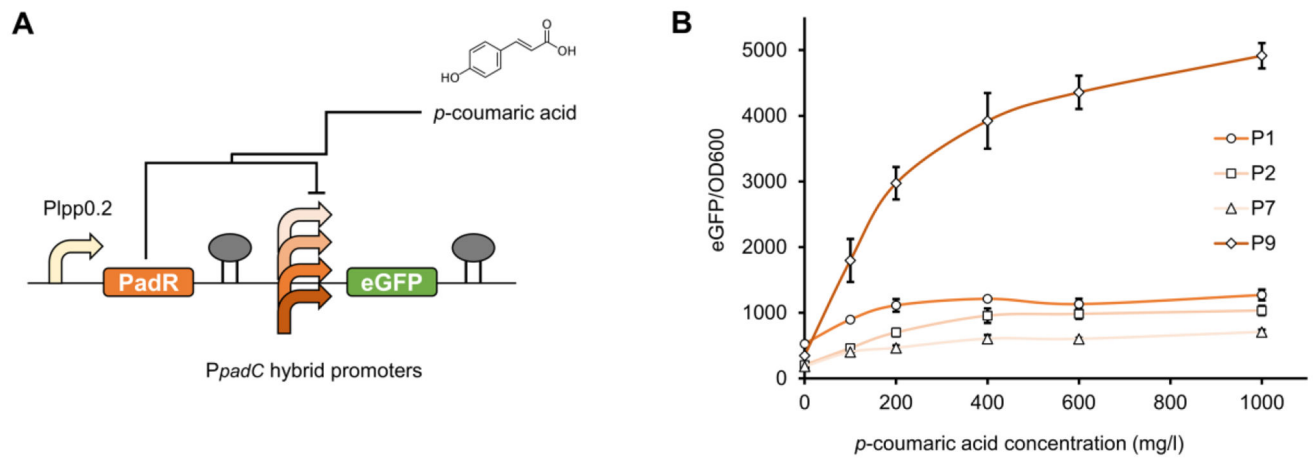


Figure 3. Optimizing the *p*-coumaric acid biosensor PadR with broad dynamic behavior.

(A) The plasmid configuration for the PadR-based biosensor system. PadR was constitutively expressed under the control of the Plpp0.2 promoter and the reporter gene eGFP was controlled by the hybrid P_{padC} promoters (P1, P2, P7, and P9). (B) The dynamic range of the PadR- P_{padC} biosensor system harboring different hybrid promoters against different concentrations of *p*-coumaric acid (0, 100, 200, 400, 600, and 1000 mg/l). All data represent the mean of three biologically independent samples and error bars show standard deviation.

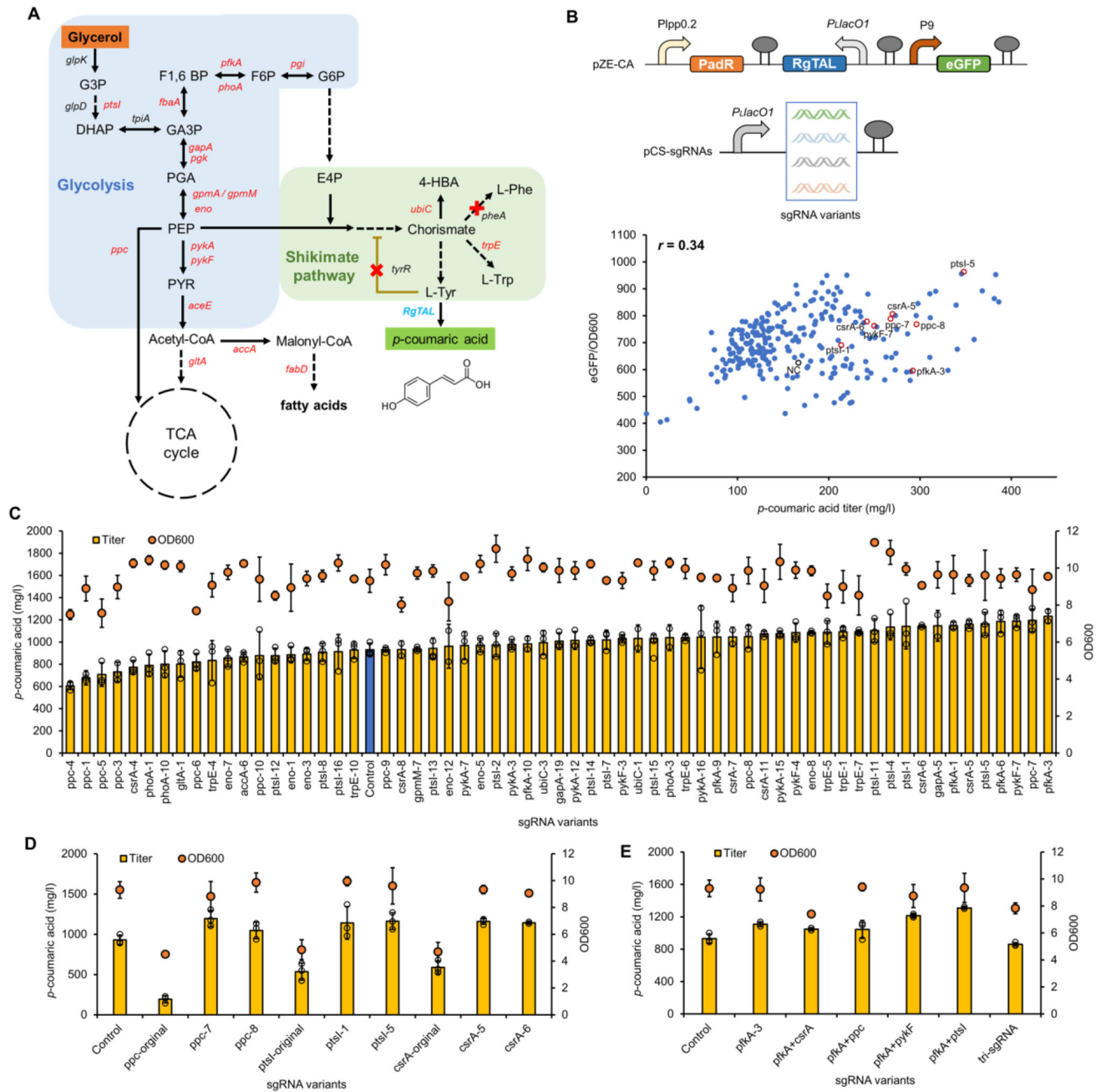


Figure 4. BATCH screening of *p*-coumaric acid over-production phenotypes.

(A) The biosynthetic pathway of *p*-coumaric acid from glycerol. Genes marked in red are CRISPRi targets and red crosses are genes to be knocked out. (B) The correlation between *p*-coumaric acid titers (in test tubes) and green fluorescence intensities in *E. coli*:dCas9 *pheA tyrR* with pZE-CA and pCS-sgRNA mismatch variants. *p*-Coumaric acid titers and normalized fluorescence intensities of the negative control strain with no sgRNA and strains with representative beneficial sgRNA variants were highlighted in black and red circles, respectively. (C) Titers of *p*-coumaric acid and cell densities (OD₆₀₀) of *p*-coumaric acid producers with selected sgRNA variants in shake flasks. The blue bar represents the

control strain without any sgRNA. **(D)** Comparison of *p*-coumaric acid production between authentic (fully matched) sgRNAs targeting *ppc*, *ptsI* and *csrA*, and their counterpart mismatch sgRNA variants in shake flasks. **(E)** The *p*-coumaric acid production test with combination of selected sgRNA variants in shake flasks. The number behind each gene name represented the order of selected colonies harboring counterpart mismatch sgRNA variants. Negative control (NC), *E. coli*::dCas9 *pheA tyrR* with pZE-CA and pCS27. All data represent the mean of three biologically independent samples and error bars show standard deviation.

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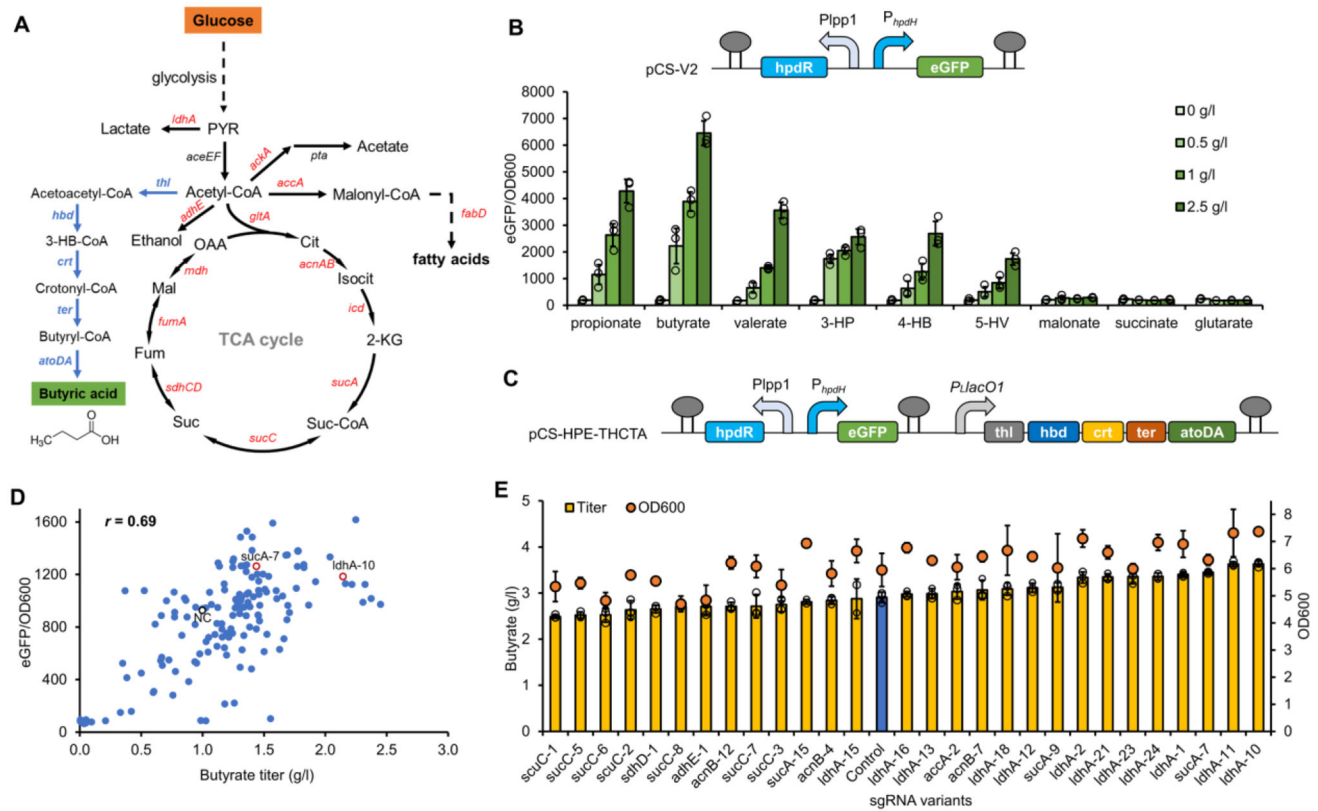


Figure 5. BATCH screening for butyrate over-production phenotypes.

(A) The biosynthetic pathway for butyrate from glucose. Genes shown in blue are butyrate pathway genes and genes shown in red are CRISPRi targets. (B) The ligand profiling of the engineered HpdR- P_{hpdH} biosensor against C3-C5 fatty acids (propionate, butyrate, and valerate), hydroxy acids (3-hydroxypropionate (3-HP), 4-hydroxybutyrate (4-HB), and 5-hydroxyvalerate (5-HV)), and diacids (malonate, succinate, and glutarate) with gradient concentrations of 0–2.5 g/l. (C) The plasmid scheme of pCS-HPE-THCTA for BATCH screening of butyrate high-production phenotypes. (D) The correlation between butyrate titers (in test tubes) and green fluorescence intensities in *E. coli*:dCas9 with pCS-HPE-THCTA and pZE-sgRNA mismatch variants. Butyrate titers and normalized fluorescence intensities of negative control strain with no sgRNA and strains with representative beneficial sgRNA variants were highlighted in black and red circles, respectively. (E) Titers of butyrate and cell densities (OD₆₀₀) of butyrate producers with selected sgRNA variants in shake flasks. The blue bar represents the control strain without any sgRNA. The number behind each gene name represented the order of selected colonies harboring counterpart mismatch sgRNA variants. Negative control (NC), *E. coli*:dCas9 with pCS-HPE-THCTA and pHA-MCS. All data represent the mean of three biologically independent samples and error bars show standard deviation.

Table 1.

Strains and plasmids

Strains	Description	Reference
<i>E. coli</i> XL1-Blue	<i>recA1 endA1gyrA96thi-1 hsdR17supE44relA1 lac</i>	Stratagene
<i>E. coli</i> BW25113(F')	<i>rrnBT14 lacZ JW16 hsdR514 araBADAH33 rhaBADLD78 F'[traD36 proAB lacIqZ M15 Tn10(Tetr)]</i>	(Atsumi et al., 2008)
<i>E. coli</i> :dCas9	<i>E. coli</i> BW25113(F') with the integration of the <i>P_LlacO1-dCas9</i> cassette into the genome at <i>dkgB</i> locus	This study
<i>E. coli</i> :dCas9 <i>pheA tyrR</i>	<i>E. coli</i> :dCas9 with the deletion of <i>pheA</i> and <i>tyrR</i>	This study
Plasmids	Description	Reference
pZE12-luc	<i>P_LlacO1</i> ; <i>luc</i> ; <i>ColE1 ori</i> ; Amp ^R	(Lutz and Bujard, 1997)
pCS27	<i>P_LlacO1</i> ; <i>p15A ori</i> ; Kan ^R	(Shen and Liao, 2008)
pZE-eGFP	pZE12luc harboring the <i>P_LlacO1</i> -controlled eGFP expression cassette	(Wang et al., 2021b)
pCS-sgRNA	pCS27 harboring sgRNA scaffold targeting eGFP	(Wang et al., 2017)
pZE-sgRNA	pZE12-luc harboring the sgRNA scaffold amplified from pCS27-sgRNA and inserted in between <i>XhoI</i> and <i>AvrII</i>	This study
pHA-MCS	pZE12-luc with the <i>luc</i> gene replaced by a newly designed multi-cloning site	This study
pCS-sgegfp-cds1	pCS27 harboring the <i>P_LlacO1</i> -controlled transcriptional cassette of a fully matched sgRNA targeting the +39 bp of <i>eGFP</i>	This study
pCS-sgegfp-rbs	pCS-sgegfp targeting the RBS region of the <i>eGFP</i>	This study
pCS-sgegfp-cds2	pCS-sgegfp targeting the coding sequence (starting from + 378 bp)	This study
pZE-Plpp0.2-PadR-P1-eGFP	pZE-Plpp0.2-PadR harboring the hybrid promoter P1-controlled <i>eGFP</i> expression cassette	This study
pZE-Plpp0.2-PadR-P2-eGFP	pZE-Plpp0.2-PadR harboring the hybrid promoter P2-controlled <i>eGFP</i> expression cassette	This study
pZE-Plpp0.2-PadR-P7-eGFP	pZE-Plpp0.2-PadR harboring the hybrid promoter P7-controlled <i>eGFP</i> expression cassette	This study
pZE-Plpp0.2-PadR-P9-eGFP	pZE-Plpp0.2-PadR harboring the hybrid promoter P9-controlled <i>eGFP</i> expression cassette	This study
pZE-CA	pZE-Plpp0.2-PadR-P9-eGFP harboring the codon-optimized RgTAL from <i>Rhodotorula glutinis</i> under control of the <i>P_LlacO1</i> promoter	This study
pCS-V2	pCS27-Plpp1-HpdR- <i>P_{hpdH}</i> -eGFP with deletion of a palindromic sequence (-209 to -118 bp) on the <i>P_{hpdH}</i> promoter and HpdR under control of Plpp1.0 promoter	(Wang et al., 2021b)
pCS-THCTA	pCS27 harbors the butyrate synthetic pathway which includes the <i>thl</i> , <i>hbd</i> , <i>crt</i> , <i>ter</i> , and <i>atoDA</i> .	This study
pCS-HPE-THCTA	pCS-THCTA harboring Plpp1-HpdR- <i>P_{hpdH}</i> -eGFP (HPE) biosensor cassette from pCS-V2	This study