Yeast transcriptional device libraries enable precise synthesis of value-added chemicals from methanol

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ABSTRACT

Natural methylothrophs are attractive methanol utilization hosts, but lack flexible expression tools. In this study, we developed yeast transcriptional device libraries for precise synthesis of value-added chemicals from methanol. We synthesized transcriptional devices by fusing bacterial DNA-binding proteins (DBPs) with yeast transactivation domains, and linking bacterial binding sequences (BSs) with the yeast core promoter. Three DBP–BS pairs showed good activity when working with transactivation domains and the core promoter of PAOX1 in the methylothrophic yeast, Pichia pastoris. Fine-tuning of the tandem BSs, spacers and differentiated input promoters further enabled a constitutive transcriptional device library (cTRDL) composed of 126 transcriptional devices with an expression strength of 16–520% and an inducible TRDL (iTRDL) composed of 162 methanol-inducible transcriptional devices with an expression strength of 30–500%, compared with PAOX1. Selected devices from iTRDL were adapted to the dihydromonacolin L biosynthetic pathway by orthogonal experimental design, reaching 5.5-fold the production from the PAOX1-driven pathway. The full factorial design of the selected devices from the cTRDL was adapted to the downstream pathway of dihydromonacolin L to monacolin J. Monacolin J production from methanol reached 3.0-fold the production from the PAOX1-driven pathway. Our engineered toolsets ensured multilevel pathway control of chemical synthesis in methylothrophic yeasts.

GRAPHICAL ABSTRACT

INTRODUCTION

In view of its quantitative, predictive and engineering characteristics, synthetic biology has pushed forward the conversion of life cognition to life design. Recently, the development of theories and methods in synthetic and molecular biology has allowed the sophisticated rewiring of non-natural life. Most successes have been achieved on model microbial hosts, such as Escherichia coli (1,2) and Saccharomyces cerevisiae (3,4), whereas other non-conventional hosts have not been explored. As different strains have different genetic backgrounds, they may have specific biological components that adapt to different application scenarios. Therefore, exploration of individualized set-ups for different strains is required.

For instance, the bio-utilization of sustainable one-carbon (C1) substrates such as methanol, methane and CO2 has attracted widespread attention in both academia and industry (5). Methanol is a major byproduct of the fossil industry and a promising product of methane oxidation or CO2 reduction. It represents a probable C1 substrate for industrial bio-utilization, with its liquid state being compatible for transportation and fermentation con-

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control (6–8). Moreover, methanol has a 50% higher degree of reduction per mole of carbon than sugar substrates such as glucose, and thus provides more surplus electrons for compound synthesis (9). With mature genetic manipulation tools, some non-methylotrophic strains, such as E. coli (10–12), Corynebacterium glutamicum (13–15) and S. cerevisiae (16–18), have been engineered to utilize methanol by the re-assembly of exogenous methanol assimilation pathways in cells. Nevertheless, their weak methanol utilization ability still lags far behind their production requirements (19). Instead, natural methylotrophs are known to efficiently utilize methanol, but lack sufficient genetic tools to enable multigene pathways (20–22).

In the past two decades, the methylotrophic yeast, *Pichia pastoris* (syn. *Komagataella phaffii*), has been shown to be an excellent workhorse for protein production (20,23–25) and a potential cell factory for chemical synthesis (26–29). This organism can efficiently utilize methanol with high-level expression of endogenous alcohol oxidase 1 (Aox1) and thus grows well on methanol as the sole carbon source (22). Most recently, it was further developed as an autotrophic strain capable of growth on CO2 with methanol (syn. *P. pastoris* gene pathways (20–22). Methanol-inducible promoters with gradient strength were used to construct a synthetic methanol-inducible TRDL (iTRDL) composed of 162 devices with an intensity range of 30–520% (P_{AOX1} as 100%). Methanol-inducible input promoters and gradient strength were used to construct a synthetic methanol-inducible TRDL (iTRDL) composed of 162 devices with an intensity range of 30–520% (P_{AOX1} as 100%).

The functions of iTSAD inspired us to explore the synthetic transcriptional components of the full coverage intensity. This could be realized by combinatorial assembly of various cis- and trans-acting elements from bacteria and yeast, and by manipulation of the spacers between the binding and core sequences of the cis-acting elements. Constitutive control of these elements enabled a synthetic constitutive transcriptional device library (cTRDL) composed of 126 devices with an intensity range of 16–520% (P_{AOX1} as 100%). Methanol-inducible input promoters and gradient strength were used to construct a synthetic methanol-inducible TRDL (iTRDL) composed of 162 devices with an intensity range of 30–520% (P_{AOX1} as 100%). We subsequently tested the applicability of the iTRDL and cTRDL in multigene pathway expression and balance. We selected devices from the iTRDL to adapt to dihydromonacolin L (DML) biosynthesis from methanol using an orthogonal experimental design, which led to a 5.5-fold DML titre compared with that from the P_{AOX1}-driven pathway. Furthermore, the selected devices from the cTRDL were adapted to the downstream pathway of DML to monacolin J (MJ) using a full factorial design. It finally achieved a 3.0-fold MJ titre compared with that from the P_{AOX1}-driven pathway with methanol as the substrate. Our study provides an alternative expression toolbox and a strategy for methanol bio-utilization in natural methylotrophs.

**MATERIALS AND METHODS**

**Strains and growth conditions**

The plasmids and strains used in this study are listed in Supplementary Table S5 and S6, respectively. *Pichia pastoris* strains GS115, GS_P_{AOX1G}, and GS_C_{AOX1G} and the yeast expression vectors pPIC3.5K, pPICZ_B and pGAPZ B were stored in our laboratory. *Escherichia coli* Top10 was purchased from Invitrogen. The plasmids BB1_{12}, BB1_{23}, BB1_{34}, BB2_AB, BB2_BC and BB3eN_{14} for GoldenPcCS cloning were purchased from Addgene (#98549). The plasmids pP_{AOX1G} (pP-GFP), pPCag, pPcAG, and pGAPZ were stored in our laboratory. *Escherichia coli* Top10 was cultivated in LB medium (0.5% yeast extract, 1% tryptone and 0.5% NaCl). Antibiotics (100 μg/ml ampicillin or 50 μg/ml zeocin) were added when required. *Pichia pastoris* was incubated at 30°C in YPD (1% yeast extract, 2% tryptone and 2% glucose) or MGY (0.67% YNB and 1% glycerol) medium for cell growth. To screen *P. pastoris* transformants, YPD or MGY medium was supplemented with the appropriate antibiotics (100 μg/ml zeocin, 750 μg/ml hygromycin B and 100 μg/ml nourseothricin) or histidine (20 μg/ml). *Pichia pastoris* was cultured in YNM medium (1.34% YNB and 0.5% methanol) for reporter protein expression or in YNMB medium (YNM with 0.3% K_{2}HPO4 and 1.18% KH2PO4) for compound production. A solid medium was obtained by adding 2% agar.
Plasmid and strain construction

Oligonucleotides were synthesized by Suzhou Genewiz Biotech Co., Ltd, China and are listed in Supplementary Table S7. The plasmids used in this study were constructed by seamless cloning and/or Gibson assembly (ClonExpress™ II one-step cloning kit, Vazyme Biotech Co., Ltd, China). Molecular genetic analyses of *E. coli* and *P. pastoris* were performed as previously described (29,54). The construction details of the plasmids and strains are described in supplementary Materials and methods.

Fluorescence and transcriptional analysis

The strain stored at −80°C (20 μl) was transferred into 2 ml of fresh YPD medium in a 20 ml serum bottle and cultivated for 2 days for strain activation. Yeast broth was inoculated into fresh YPD medium for pre-culturing to an optical density (OD600) of 6.0. The cells were harvested by centrifugation (5000 × g, 5 min), washed twice with sterile water and then distributed to the required medium (YNM) to a final OD600 of 1.0 for culture in a 24-well plate (<4 ml of liquid in each 10 ml well). The cell broth was collected at 12, 18 and 24 h and washed twice with sterile water for fluorescence determination. Reporter enhanced green fluorescent protein (eGFP) fluorescence (normalized to OD600) of the cultured samples was analysed using a multimode microplate reader (Synergy 2, BioTek Instruments, USA) at an excitation wavelength of 485 nm and an emission wavelength of 525 nm (gain, 60 nm). Transcriptional levels of biosynthetic genes were tested to evaluate their relationships with MJ and DML production among the different strains. The details of transcriptional analysis were described in supplementary Materials and methods.

Cell growth analysis

The growth of strains carrying synthetic TFs (sTFs) and eGFP expression cassettes driven by various promoters was analysed in a shake-flask culture. *Pichia pastoris* GS115 was used as the control strain (wild-type, WT). Cells were precultivated overnight in 50 ml of YPD medium after activation from storage (−80°C). Then, 200 μl of culture broth was transferred into 50 ml of YPD medium in a 250 ml shake flask and grown to OD600. Cells were harvested by centrifugation (5000 × g, 5 min), washed twice with sterile water and resuspended in YNB medium. Finally, the cells were inoculated into YNM medium at a final OD600 of 1.0 for growth measurement. Samples were collected and measured every 4 h. Three biological replicates were used for each strain.

Production and detection of MJ and intermediates

Cells resuspended in YNB medium were then inoculated into YNMB medium to produce the desired compounds at a suitable cell density. During culture, methanol was added separately to an appropriate level every 24 h. Culture broth was collected at certain time intervals, extracted using a suitable volume of ethyl acetate and vortexed for 3 min. The organic phase was evaporated under reduced pressure and dissolved in methanol. The extracts were analysed by high-performance liquid chromatography (HPLC) equipped with a C18 column (Kromasil™, Sweden; 250 mm × 4.6 mm × 5 mm, 100 A spherical silica) using a gradient elution strategy at 1 ml/min. The mobile phase consisted of HPLC-grade H2O containing 0.1% acetic acid (A) and HPLC-grade acetonitrile (B).

Statistical analysis

Data were obtained from three biological replicates assayed in at least two experimental batches and are presented as the mean ± standard deviation. Data were analysed using GraphPad Prism (version 7.04). The unpaired two-tailed Student’s *t*-test was used to assess differences among the grouped data. Statistical significance was set at *P* < 0.05.

RESULTS

Engineering of the cTRDL with constitutively varied expression strength

The transcription of protein-coding genes in eukaryotes begins with the assembly of RNA polymerase II and general TFs on promoter DNA (55). A transcriptional activator harbouring a separate N-terminal DNA-binding domain and C-terminal activation domain is essential for this process (55). For example, the most typical yeast transcription factor, Gal4, binds to regulatory sites on the *GAL1* promoter naturally. The activation domain of Gal4 fused to the bacterial regulator LexA (DNA-binding protein) was then expressed in yeast together with a reporter plasmid bearing LexA-binding sites upstream of the *GAL1* promoter. The chimeric protein successfully activated reporter transcription (55). This method further allows different applications of domain-swapped and function-extended biological TFs (53,56). We previously identified a Zn(II)2Cy5s6-type transactivator, Mit1, with a separable DNA-binding domain interacting with P*AOX1* (a strong and widely used methanol-inducible promoter) and an activation domain that helps recruit RNA polymerase II in *P. pastoris* (57). Using the domain-swapping strategy, we constructed TSAD using an sTF of LacI-Mit1AD (*E. coli lac* operator linked to the core region of P*AOX1*) (29). The improvement of the functional components further generated an iTSAD with high expression capacity (54). These preliminary results provide the basic structure of the transcriptional device libraries that we aimed to develop in this study.

A synthetic transcriptional device mainly carries four components: a DNA-binding protein (DBP), binding sequence (BS), TF activation domain (TFAD) and core promoter (CP). Therefore, the combination of diverse DBP–BS and TFAD–CP may lead to the development of various engineered devices (Figure 1A). We first selected different bacterial regulators, LacI (29), AraC (54), LexA (58), AcrR and BetI (59), as DBPs to fuse with Mit1AD using a GGGGS linker (Supplementary Figure S1; Supplementary Table S1). An SV40 nuclear localization sequence (NLS) was fused at the N-terminus of DBPs (except LacI) to allow them to enter the nucleus of *P. pastoris*. LacI was able
Figure 1. Design and characterization of the cTRDL. (A) Genetic circuit scheme of the cTRDL with increased regulator BSs. A chimeric transactivator composed of a DBP and a TFAD is driven by an input promoter. It targets the BSs located upstream of cP\textsubscript{AOX1} and recruits RNA polymerase to activate transcription of the gene of interest. (B) Heatmaps showing the fluorescence intensity of the diverse devices. Three DBPs fused with three TFADs (Supplementary Figure S1) to generate nine chimeric transactivators, i.e. LacI–Mit1AD (IM), AraC–Mit1AD (CM), LexA–Mit1AD (AM), LacI–Mxr1AD (IX), AraC–Mxr1AD (IX), LexA–Mxr1AD (AX), LacI–Prm1AD (IP), AraC–Prm1AD (CP) and LexA–Prm1AD (AP). Tandem copies (n = 1–9) of BSs (lacO, araI and lexO) were linked with cP\textsubscript{AOX1} to obtain synthetic promoters. cA represents cP\textsubscript{AOX1} and G indicates eGFP. The eGFPs expressed by the cTRDL were tested in methanol medium (YNM). The detected fluorescence data for each subset of cTRDLs are shown in Supplementary Figure S2. (C) Genetic circuit scheme of the cTRDL with increased spacers between BSs and cP\textsubscript{AOX1}. Tandem copies (m = 1–9) of HA spacer were inserted between one BS and cP\textsubscript{AOX1} to fine-tune the output strength of the cTRDL. (D) Heatmaps showing the fluorescence intensity of the diverse devices. Nine chimeric transactivators were described in (B). Each activator was paired with a single copy of a specific BS which was spaced apart from cP\textsubscript{AOX1} by 1–9 copies of HA spacers. cA and G were the same as in (B). The detected fluorescence data for each subset of cTRDLs are shown in Supplementary Figure S3. (E) Full functional devices in the cTRDL with smoothly increasing output strength. A total of 126 devices are included (exclusion of 18 inactive ones) from all the constructs in (B) and (D). Chimeric transactivators were controlled by the constitutive promoter P\textsubscript{GAP}. The serial number (Supplementary Table S2) for each column is marked sequentially below the figure, which corresponds to region I, II and III, respectively, on the horizontal axis.
to localize to the P. pastoris nucleus without an NLS (29).

These five engineered sTFs, driven by the constitutive promoter P\textsubscript{GAP}, were tested to determine whether the constitutive eGFP expression with interacting with their specific operators lexO, araI, lexO, acrAB and bet linked by cP\textsubscript{AOX1} (Supplementary Figure S1; Supplementary Table S1). Almost no cross-talk was observed between the different sTFs and the hybrid promoters (Supplementary Figure S1). LacI, AraC and LexA functioned well; therefore, they were selected for fusing with the activation domains of the transcription activators Mit1, Mxr1 and Prm1 (57, 60, 61). Nine cross combinations were constructed to work with the corresponding promoters Mit1AD, Mxr1AD and Prm1AD (57, 60, 61). Therefore, our engineering cTRDL output reached 100%. Moreover, the strength of the commonly used promoters Mit1, Mxr1 and Prm1 (57, 60, 61). Nine cross combinations were constructed to work with the corresponding promoters Mit1AD, Mxr1AD and Prm1AD (57, 60, 61). Therefore, our engineering cTRDL output reached 100%. Moreover, the strength of the commonly used promoter P\textsubscript{GAP} in P. pastoris was approximately one-third that of methanol-inducible P\textsubscript{AOX1} (51, verified in our lab). Therefore, our engineering cTRDL output reached \( \sim 60\)–1650% of the intensity of P\textsubscript{GAP}. Overall, we provide a constitutive expression toolbox that far exceeds conventional P\textsubscript{GAP} promoter variants (42, 43).

Engineering of the iTRDL with inducibly varied expression strength

Genes can be expressed in the induction mode using inducible promoters. By inducible expression, the cell growth and production phases are separated to relieve the metabolic burden from recombinant proteins or their catalytic products (63, 64). Therefore, we designed an iTRDL for flexible expression control of product synthesis from methanol.

The overexpression of sTFs may impair cell growth and metabolism in methanol (65). Thus, we first selected various methanol-inducible promoters, such as P\textsubscript{AOX2}, P\textsubscript{FGHI}, P\textsubscript{PEX5}, P\textsubscript{TAL2}, P\textsubscript{FBG}, P\textsubscript{FLD1}, P\textsubscript{CAT7}, P\textsubscript{DAS1} and P\textsubscript{AOX1} (51, 66), to drive sTFs (specifically LacI–TFADs) and analysed their effects on cell growth. These promoters were first tested in a methanol medium (YNM) to verify their differential expression activities (Supplementary Figure S4). The growth of strains containing LacI–TFADs driven by these promoters was tested correspondingly (Supplementary Figure S5). We found that the growth of strains carrying LacI–Mit1AD was severely impaired by all promoters except for the weakest P\textsubscript{AOX2}. Impairment of cell growth was also observed in strains harbouring LacI–Mxr1AD with promoters stronger than P\textsubscript{FLD1}. In contrast, strains harbouring LacI–Prm1AD grew normally, except those driven by the strongest P\textsubscript{AOX1}. To retain Mit1AD-derived synthetic devices, we used the methanol-inducible promoter P\textsubscript{P374} (66) to drive LexA–Mit1AD, which did not impair cell growth (Supplementary Figure S5).

Subsequently, various sTFs were expressed by their adaptive inducible promoters to work with hybrid promoters harbouring tandem operators and spacers, as described in the cTRDL (Figure 2A). As anticipated, increased copies of BS improved the output strength, and increased copies of the spacer permitted a decrease in the output strength of devices with various sTFs (Figure 2B; Supplementary Figure S6A–I). Generally, synthetic devices harbouring Mxr1AD or Mit1AD are stronger than those harbouring Prm1AD. We obtained the strongest device LexA–Mit1AD/lexO3–cP\textsubscript{AOX1} with the input promoter P\textsubscript{P374} (Figure 2B; Supplementary Figure S6C) and the weakest device LacI–Prm1AD/\textsubscript{lacO1HA8}–cP\textsubscript{AOX1} with the input promoter P\textsubscript{DAS1} (Figure 2B; Supplementary Figure S6G). Finally, they constituted a fine-tuned iTRDL containing 162 methanol-inducible synthetic devices (Figure 2C; Supplementary Table S2). The iTRDL covers an expression strength of 30–500% (tunable range of 16.7-fold) compared with methanol-inducible P\textsubscript{AOX1} (referred to as 100%). Therefore, we provided a methanol-responsive expression toolbox that far exceeds that of the P\textsubscript{AOX1} promoter variants (42, 43).

iTRDL devices adapted for upstream pathway improved synthesis of the intermediate product dihydroxymalonolactone L from methanol

Currently, heterologous synthesis of complicated compounds via microbial systems is often inefficient because of low yield and intermediate accumulation. Pathway balancing has been used as a prioritized strategy for solving such problems (41). Our synthetic transcriptional de-
Design and characterization of the methanol-inducible iTRDL. (A) Genetic circuit scheme of iTRDL. Four available methanol-inducible promoters were identified from 10 candidates, which were of different strength (Supplementary Figure S4) and favoured cell growth when expressing specific chimeric transactivators (Supplementary Figure S5). These four promoters, i.e. $P_{AOX1}$, $P_{AOX2}$, $P_{DAS1}$ and $P_{FBA2}$, were used to drive the diverse chimeric transactivators verified in Figure 1, which can target the BSs of various hybrid promoters. Increased tandem copies of BSs and HA spacer were also adopted in this design to adjust device strength. (B) Heatmaps showing the fluorescence intensity of the diverse devices. Three DBPs fused with three TFADs (Supplementary Figure S1) to generate nine chimeric transactivators as described in Figure 1. Different chimeric transactivators were controlled by $P_{AOX1}$, $P_{AOX2}$, $P_{DAS1}$ and $P_{FBA2}$ correspondingly as determined in Supplementary Figure S6. Tandem copies ($n = 1–9$) of BSs ($lacO$, $araI$ and $lexO$) were linked with $cP_{AOX1}$ to obtain synthetic promoters. Also, tandem copies ($m = 1–9$) of HA spacer were inserted between one BS and $cP_{AOX1}$ to further fine-tune the output strength of the iTRDL. $cA$ represents $cP_{AOX1}$ and $G$ indicates eGFP. eGFPs expressed by the iTRDL were tested in methanol medium (YNM). The detected fluorescence data for each subset of the iTRDL are shown in Supplementary Figure S6. (C) Full functional devices in the iTRDL with smoothly increasing output strength. A total of 162 devices are included from all the constructs in (B). The serial number (Supplementary Table S2) for each column is marked sequentially below the figure, which corresponds to region I, II and III, respectively, on the horizontal axis.
ervices offer new choices for fine-tuning pathway control in methylotrophic yeast. We then tested the applicability of the cTRDL and iTRLD in combinatorial control of biosynthetic pathways. We used MJ, which is a critical substrate for the semi-synthesis of the commercial hypolipidemic drug simvastatin, as a reporter compound. MJ is synthesized via the fungal polyketide pathway and involves six enzymes: nonaketide synthase (LovB), enoyl reductase (LovC), thioesterase (LovG), phosphopantetheinyl transferase (NpgA), cytochrome P450 (LovA) and cytochrome P450 oxidoreductase (CPR) (28).

We split the full pathway into an upstream polyketone synthesis module and a downstream oxidation module. The methanol-inducible iTRLD was first applied to the upstream module to separate the cell growth and production phases (Figure 3A). This module contains four enzymes, LovB, LovC, NpgA and LovG, which synthesize the intermediate DML. To precisely balance the DML pathway, we used a statistical method, an orthogonal experimental design, involving four factors (enzyme-coding genes) and three expression levels (selected devices from the iTRLD).

Two previously constructed strains, \(P_{Pp}P_{AOX1}/BCGN (D1 and D9)\), were used as control groups, in which biosynthetic genes were recognized as single-copy (D1) or multicopy (D9), driven by \(P_{AOX1}\) (28, 67). We selected devices with expression strengths not lower than that of \(P_{AOX1}\) for this experimental design. Moreover, methanol-inducible promoters with an expression intensity of \(\geq 200\%\) relative to \(P_{AOX1}\) have seldom been reported. To test the effectiveness of our devices, we set the three expression levels to \(\sim 100, 250\) and \(450\%\) (Supplementary Table S3) to adapt to the four biosynthetic genes, thus generating nine experimental groups (A1–A9) (Supplementary Table S4; Figure 3B).

We then selected suitable devices from the iTRLD, i.e. \(P_{FB2\Lambda}LacI-\text{Mxr1AD}\) paired with \(HA2/lacO1cP_{AOX1}\) (~100%; Device No. 162), \(lacO3cP_{AOX1}\) (~250%; Device No. 166) and \(lacO9cP_{AOX1}\) (~450%; Device No. 172) (Supplementary Table S2), to regulate the expression of biosynthetic genes (Figure 3C). Finally, we successfully obtained strains of \(P_{Pp}/iTRDL-BCGN (A1–A9)\) with the desired genotypes, all of which were identified as carrying single-copy gene cassettes. Subsequently, the strains were cultured for DML synthesis in methanol (YNMB medium). Notably, the resulting DML titre varied greatly among the nine experimental groups (Figure 3C), illustrating that the expression levels of biosynthetic enzymes significantly affected pathway balancing. These results also proved that our methanol-inducible iTRLD effectively regulates various pathway nodes. The A3 strain with LovB expression by the strongest device (~450%), LovC (~250%), LovG (~100%) and NpgA (~250%). Therefore, we constructed this strain (Opt), which further improved the DML titre (by 6.0%) and productivity (by 11.4%) compared with strain A3. The growth of these strains was also determined (Figure 3D). The overexpression of biosynthetic enzymes imposes a metabolic burden on cells, thus affecting cell growth. Compared with strain D1 with single-copy genes expressed by \(P_{AOX1}\), the growth of strains with either multicopy genes driven by \(P_{AOX1}\) (D9) or single-copy genes driven by iTRLD devices (A1–A9, Opt) was relatively weak.

**Adaptation of iTRLD and cTRDL devices for downstream pathway improved synthesis of the final product MJ from methanol**

DML was oxidized to the final product MJ by the downstream module with LovA and its partner CPR (Figure 4A). We further tested the adaptation of the iTRLD and cTRDL devices to the downstream pathway for MJ synthesis. Independent DBP–BS pairs are necessary to reduce transcriptional cross-talk between the upstream and downstream modules. Thus, we chose LexA–Mit1AD, but not LacI–Mit1AD, as the downstream module (Figure 4B), considering the non-interactive relationship between LacI–lacO and LexA–lexO (Supplementary Figure S1), and non-interaction between Mit1 and Mxr1 (57).

We hypothesized that the accumulation of downstream enzymes prior to the induction of the upstream module might accelerate the conversion of the intermediate to the final product. Therefore, the devices selected from cTRDL were first used for downstream pathway assembly. The Opt strain producing DML (Figure 3C) was used as the parent strain. Similar to the upstream module, devices with expression levels of ~100, 250 and 450% (referred to as \(P_{AOX1}\)) were adapted for codon-optimized LovA (sLovA) and CPR. We then selected appropriate constitutive devices from cTRDL, i.e. \(P_{Gap1}LexA–Mit1AD\) paired with \(lexO1HA8cP_{AOX1}\) (~100%; Device No. C128), \(lexO1HA3cP_{AOX1}\) (~250%; Device No. C133) and \(lexO3cP_{AOX1}\) (~500%; Device No. C57) (Supplementary Table S2). A full factorial design of these devices adapted with sLovA and CPR was performed, and nine recombinant strains were generated and designated as \(P_{Pp}/iTRDL-BCGN_cTRDL-sAR (OC1–OC9)\). Similarly, our previously constructed \(P_{Pp}/P_{AOX1}-BCGN_sAR\) strains, J1 (single copy of sLovA and CPR expressed in D9) and J9 (multicopy of sLovA and CPR expressed in D9) (28, 67) were used as controls. We grew 11 strains directly in methanol medium (YNMB) with an initial OD\(_{600}\) of 1.7, similar to the culture of strains harbouring the upstream module. However, an extremely low MJ titre was observed with the accumulation of intermediates (Supplementary Figure S7A), as well as severe growth impairment (Supplementary Figure S8A). We suspected that the continued expression and accumulation of sLovA and CPR by cTRDL devices resulted in a metabolic burden on the cells. Then, inducible devices from iTRLD with comparative strengths were used, i.e. \(P_{O_{FB2}}LexA–Mit1AD\) paired with \(lexO1HA8cP_{AOX1}\) (~100%; Device No. I38), \(lexO1HA3cP_{AOX1}\) (~250%; Device No. I143) and \(lexO3cP_{AOX1}\) (~500%; Device No. I148). A full factorial de-
Figure 3. Orthogonal experimental design of iTRDL devices matching upstream pathway nodes for the fine-tuning synthesis of DML from methanol. (A) Biosynthetic pathway of DML used for evaluation of the iTRDL. It involves four enzymes: LovB, LovC, LovG and NpgA (28). We expressed these genes as an upstream module for the full MJ pathway. (B) Design scheme for fine-tuning production of DML by iTRDL devices. The LacI–Mxr1AD-derived devices from iTRDL were used. The corresponding hybrid promoters of lacOnHAcmA (various numbers of lacO and H4 linked with pAOX1) were used for differential expression of biosynthetic genes, respectively. (C) DML biosynthesis by separate control of pathway nodes with iTRDL devices. Devices were selected from the library subset described in (B). We employed the orthogonal experimental design method with four factors (genes) and three levels (output strengths of devices) to evaluate the fine-tuning functions of the iTRDL. The three expression levels were set as ∼100% (1.0-fold), ∼250% (2.5-fold) and ∼450% (4.5-fold) relative to PAOX1 to adapt to the four pathway genes separately (Supplementary Table S3). Thus, devices Nos I62, I66 and I72 in Figure 2C were used. Nine recombinant single-copy strains, Pp/iTRDL-BCGN (A1–A9), were obtained with correct genotypes. K-value analysis from the experimental data predicted an optimal expression combination of LovB (4.5-fold), LovC (2.5-fold), LovG (1.0-fold) and NpgA (2.5-fold). It then led to a strain with this optimal combination (designated as Opt). In contrast, two previously constructed high-yield strains, i.e. D1 (single-copy strain) and D9 (optimum multicopy strain), were used as control groups, in which each biosynthetic gene was driven by the PAOX1 (28,67). These strains were grown in glucose medium (YPD) and then shifted to methanol medium (YNMB) to the final density at an OD600 of 1.0 for incubation. N.D., not detected. (D) Cell growth of various DML-producing strains from (C). Error bars represent the standard deviation of three biological replicates assayed in duplicate or triplicate. Statistical significance of the DML titre and cell growth of Opt relative to other strains is shown at different time points (*P < 0.05 at 72, 96 and 120 h, respectively).
Figure 4. Full factorial design of iTRDL and cTRDL devices matching downstream pathway nodes to adapt to the optimal upstream module for the fine-tuning synthesis of MJ from methanol. (A) The biosynthetic pathway of MJ consisting of the upstream DML module and the downstream DML to MJ module was reassembled. The upstream module is described in Figure 3. The defined downstream module carries sLovA and CPR that mediate double oxidation steps to transform DML to the final product MJ (28). (B) Design scheme for fine-tuning production of MJ by iTRDL and cTRDL devices. The Opt strain from Figure 3 was used as the parent host. We employed the LexA–Mit1AD-derived devices in the downstream module to avoid the cross-talk with LacI–Mxr1AD in the upstream module. The subset of LexA–Mit1AD-derived devices from cTRDL and from iTRDL were used. The corresponding hybrid promoters of lexOnHAmcA (various numbers of lexO and the H4 spacer linked with cP\textsubscript{AOX1}) were used for differential expression of sLovA and CPR, respectively. Time profiles of cell growth of strain Opt equipped with the downstream cTRDL driving module (C) and iTRDL driving module (D). We selected constitutive devices (C128, C133 and C57 in Figure 1E) of three varying strengths [100% (1.0-fold), 250% (2.5-fold) and 520% (5.2-fold) relative to P\textsubscript{AOX1}] and methanol-inducible devices (I38, I43, I48 in Figure 2C) of three varying strengths [100% (1.0-fold), ∼250% (2.5-fold) and 500% (5.0-fold) relative to P\textsubscript{AOX1}] from the subset of LexA–Mit1AD-derived devices described in (B). Full factorial constitutive or inducible devices matching downstream pathway nodes were designed. It resulted in two groups of single-copy strains, i.e. P\textsubscript{p}iTRDL-BCGN\textsubscript{cTRDL-sAR} (OC1–OC9) and P\textsubscript{p}iTRDL-BCGN\textsubscript{iTRDL-sAR} (OI1–OI9) (E and F). These strains were grown in glucose medium (YPD) and then shifted to methanol medium (YNMB) to the final density at an OD\textsubscript{600} of 10.0 for incubation. As a control, two previously constructed high-production strains, i.e. J1 (single copy of sLovA and CPR expressed in D9) and J9 (multicopy of sLovA and CPR expressed in D9), were used, in which each biosynthetic gene was driven by the native P\textsubscript{AOX1} separately (28,67). (E and F) The titre of the final product MJ and the intermediates of DML and ML were analysed after 96 h culture of the OC strains (E) and OI strains (F). N.D., not detected. The error bars represent the standard deviation of three biological replicates assayed in duplicate or triplicate. Statistical significance of the MJ titre of OC4 and OI4, relative to other corresponding strains, is shown (*P < 0.05).
level production of MJ with minor accumulation of intermediates. However, excessive expression of sLovA had negative effects on MJ synthesis, leading to low titres of MJ in the O17, O18 and O19 strains, and almost no titre in the OC7, OC8 and OC9 strains. Finally, the O14 and OC4 strains produced 172 and 208 mg/l MJs, respectively. The constitutive expression of sLovA and CPR by the cTRDL surpassed their inducible expression by the iTRLD when adapted to the inducible expression mode of the upstream module by the iTRLD. MJ production of strain OC4 by selected devices from the cTRDL reached 3.0-fold that of the single-copy control J1 and 1.8-fold that of the multicopy control J9 by P<sub>AOX1</sub>. The results illustrate that the strengthening and balancing of the pathway nodes inside and between the two modules can be precisely controlled. We offer new toolboxes for the high-level biosynthesis of heterologous chemicals from methanol in yeast.

DISCUSSION

Although remarkable progress has been made in reprogramming bacteria and baker’s yeast into synthetic methylotrophs (10–18), breakthroughs in the methanol utilization capacity for industrial use remain a huge challenge (19,68). The production of chemicals by natural methylotrophs has provided an alternative path to methanol bio-utilization for industrial purposes (28,69–71). However, the lack of genetic tools severely prevents natural methylotrophs from synthesizing valuable chemicals via multi-gene pathways (20–22). Although the methylotrophic yeast P. pastoris has been used as an attractive protein expression host (20,23–25), its use for the production of chemicals via complicated pathways is still limited by insufficient fine-tuning expression tools (21).

In this study, we demonstrated that transcriptional device libraries with broad expression levels and smooth changes can be achieved by rational design in addition to conventional promoter variants. We engineered sTFs by fusing endogenous TFADs with different bacterial DBPs, in which hybrid cis-elements were designed by linking DBP-paired BSs and TFAD-contributing CPs. Three workable bacterial DBP-BS pairs with minimal cross-talk, LacI–lacO, LexA–lexO and AraC–araI, were selected from the five tested groups (Supplementary Figure S1). Their combinations with activation domains of three P. pastoris TFs, Mit1, Mxr1 and Prm1 (57), finally generated a cTRDL with 126 constitutive transcriptional devices when the TFs were constitutively expressed (Figure 1; Supplementary Figures S2 and S3). Subsequently, methanol-inducible promoters with gradually changing strengths were used to drive sTFs, thus generating an iTRLD with 162 methanol-inducible transcriptional devices (Figure 2; Supplementary Figure S3). With regard to the strength of the native P<sub>AOX1</sub> (100%), our engineered cTRDL and iTRLD presented expression intensity ranges of 16–520% and 30–500%, respectively. They provide diverse and powerful expression tools compared with traditional promoter variants. In addition, the minimal cross-talk among various sTF–BS pairs may extend their application scenarios, which shows a unique advantage over the reported promoter libraries (42–45,48).

The synthetic cTRDL and iTRLD were then tested to fine-tune the multigene biosynthetic pathway of the hyperpolipidaemic drug intermediate, MJ. The six essential genes of the MJ biosynthetic pathway were divided into two modules. Thus, we evaluated the regulatory functions of the selected cTRDL and iTRLD devices within each module and between the two modules. The statistical method of orthogonal experimental design was employed to arrange three expression levels of devices from the iTRLD, matching the four biosynthetic genes for the upstream module. The resulting nine single-copy recombinant strains produced various levels of the target compound DML, illustrating the effectiveness of the iTRLD devices (Figure 3C). In particular, DML synthesis from methanol by the optimum strain A3 was superior to that of our previously constructed P<sub>AOX1</sub>-based single-copy-producing strain D1 and multicopy-producing strain D9. Moreover, with the predicted optimal combination of expression levels matching pathway genes, we obtained the improved strain Opt (LovB with device intensity of ∼450%, LovC and NpgA with device intensity of ∼250%, and LovG with device intensity of ∼100%). It showed a noticeable advantage over the control strains D1 (5.5-fold for titre and 9.2-fold for productivity) and D9 (2.0-fold for titre and 4.0-fold for productivity). Generally, the DML biosynthetic pathway in Opt was stronger than in the control strains, as reflected by the relative transcription levels of biosynthetic genes (Supplementary Figure S9A). These results verify the strong and flexible expression modes of the engineered iTRLD. Notably, we used only four devices that shared the same DBP–BS pair, whereas 162 iTRLD devices with various DBP–BS pairs provided sufficient optional tools for multigene pathway control. Moreover, we proved the availability of the iTRLD with a statistical experimental design, which can reduce the number of experimental trials, especially for complicated pathways.

With the success of the internal module control by the iTRLD, we next tested the iTRLD- and cTRDL-mediated pathway balancing between the two modules. A full factorial combination of three expression levels from iTRLD (or cTRDL), matching two biosynthetic genes, was designed for the downstream module (Figure 4). These downstream combinations were constructed in the Opt strain, generating nine strains that produced various levels of MJ and intermediates. The optimal expression combination, sLovA, using a device with an intensity of ∼250%, and CPR, using a device with an intensity of ∼100%, was observed for both the cTRDL and iTRLD. Moreover, the downstream module driven by cTRDL devices achieved an MJ titre which was 20.9% higher than that of iTRLD devices. This finding is in accordance with our previous study (29), which illustrated that constitutive accumulation of downstream enzymes can facilitate conversion of the generated intermediates to the final product after induction. The highest MJ production was obtained from strain OC4 by cTRDL, achieving 3.0- and 1.8-fold titres compared with the control strains J1 and J9, respectively (Figure 4). In addition, the MJ biosynthetic pathway in OC4 and O14 was stronger than that in the control strains, as reflected by the relative transcription levels of the biosynthetic genes (Supplementary Figure S9B). Generally, it represents the highest reported level of DML and MJ fermentation from methanol. Recently, we achieved an MJ production of 3.2 g/l by het-
erologous synthesis of this compound in \textit{P. pastoris} using ethanol as a substrate \cite{29}. In addition, a high MJ production of 5.5 g/l from glucose achieved by rewiring metabolic pathways has been reported in the native fungus \textit{Aspergillus terreus} \cite{72}. Therefore, there is still room for improvement in the synthesis of MJ using methanol as the sole carbon source. It is worth noting that the DBP–BS pair for devices used in the downstream module was independent of that used in the upstream module. Theoretically, this can avoid the cross-talk of transcriptional regulation control between the two modules. With the effectiveness of the statistically orthogonal experimental design and full factorial design, these synthetic TRDLs may be further applied to complicated pathway regulation using deeper algorithms, such as neural networks and machine learning, in future research. Overall, our TRDLs strengthened and balanced the pathway nodes inside and between the modules. It provides alternative yeast toolboxes for high-level biosynthesis of value-added chemicals from methanol. In addition, these TRDLs should be compatible with heterologous protein expression based on reporter protein levels.

Non-model microorganisms, such as unconventional yeast and filamentous fungi, have been extensively used in many fields, including biopharmaceuticals, novel foods, biorefining, industrial enzymes, platform chemicals, biomaterials and environmental protection. However, these strains are difficult to explore in depth because of their unclear genetic background and insufficient molecular tools. Recently, genetic manipulation tools have become possible using CRISPR–Cas-derived genetic editing technologies. However, accurate expression control systems cannot be easily achieved. Although promoter mining is a common strategy for collecting expression tools, it is difficult to obtain promoters with a high intensity and tight regulation. Ideally, promoters for expression control should be strictly inducible to separate cell growth from protein expression and to avoid a cumulative metabolic burden \cite{51,63}. Promoters covering a wide range of intensities are also indispensable for fine-tuning control from tight down-regulation to high overexpression \cite{51}. Researchers have modified promoters to breed variants of different strengths. However, the high identity of promoter sequences may cause inter-promoter recombination and transcription titration effects \cite{49–51}. Here, we propose a strategy for engineering diverse transcription machinery tools to fine-tune the expression of pathways. This represents a universal construction method of a combinatorial set-up of transactivators, CPs, DBPs and BSs (either native or heterologous), which can be reproduced in extensive methlyothrophic hosts.

**DATA AVAILABILITY**

Details of the construction steps of plasmids and strains and foundation of critical genes are provided in the Supplementary Data. All other data that support the findings of this study are available within the article and the supplementary data, and are also available from the corresponding author upon reasonable request.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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