Original Research Article

Development of a novel platform for recombinant protein production in Corynebacterium glutamicum on ethanol

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\textbf{ABSTRACT}

Corynebacterium glutamicum represents an emerging recombinant protein expression factory due to its ideal features for protein secretion, but its applicability is harmed by the lack of an autoinduction system with tight regulation and high yield. Here, we propose a new recombinant protein manufacturing platform that leverages ethanol as both a delayed carbon source and an inducer. First, we reanalysed the native inducible promoter P\textsubscript{CPL} from the acetate uptake operon and found that its limited capacity is the result of the inadequate translation initial architecture. The two strategies of bicistronic design and ribozyme-based insulator can ensure the high activity of this promoter. Next, through transcriptional engineering that alters transcription factor binding sites (TFBSs) and the first transcribed sequence, the truncated promoter P\textsubscript{A256} with a dramatically higher transcription level was generated. When producing the superfolder green fluorescent protein (sfGFP) under 1% ethanol conditions, P\textsubscript{A256} exhibited substantially lower protein accumulation in prophase but an approximately 2.5-fold greater final yield than the strong promoter P\textsubscript{neo}. This superior expression mode was further validated using two secreted proteins, camelid antibody fragment (VHH) and endoxylanase (XynA). Furthermore, utilizing CRISPRi technology, ethanol utilization blocking strains were created, and P\textsubscript{A256} was shown to be impaired in the phosphotransacetylase (PTA) knockout strains, indicating that ethanol metabolism into the tricarboxylic acid cycle is required for P\textsubscript{A256} upregulation. Finally, this platform was applied to produce the “de novo design” protein NEO-2/15, and by introducing the N-propeptide of CspB, NEO-2/15 was effectively secreted with the accumulation 281 mg/L obtained after 24 h of shake-flask fermentation. To the best of our knowledge, this is the first report of NEO-2/15 secretory overexpression.

1. Introduction

The expansion of biologics and the biopharmaceutical industry has fuelled the demand for recombinant proteins in recent years [1,2]. As the major factor influencing product quality and process cost, the selection of biological platforms used as cell factories is particularly important [3,4]. For structurally simple proteins that do not require posttranslational modifications, prokaryotic cells, particularly Escherichia coli, are still the preferred expression systems [5]. However, considering that secreting the desired protein into the supernatant significantly simplifies downstream processing and reduces the total cost [6,7], several gram-positive bacterial expression systems with prominent secretory capacity have recently been developed [8,9]. Among them, Corynebacterium glutamicum, the long-established industrial workhorse for the production of a variety of amino acids [10], possesses enormous potential for the secretory production of recombinant proteins due to its safety, excellent fermentation performance and concise secretory environment (minimal endogenous secretory proteins and rarely detectable protease activity) [11].

For a high-performance expression system, optimal promoter
sequences and efficient regulatory mechanisms are essential prerequisites; thus, many attempts have been made to enhance protein yield through promoter engineering. The methods for obtaining promoters with desirable properties and strength in C. glutamicum fall into three main categories: 1) mutagenesis of the existing strong promoters, e.g., the tac-M promoter with an extended –10 region mutation [12]; 2) a fully synthetic library, e.g., the H36 promoter from a 70-bp random sequence library [13]; 3) screening naturally occurring promoters (NOPs) [14]. Several regulatable NOPs have been identified by screening under specific conditions, e.g., maltose and gluconate-inducible promoters [15], which are beneficial for protein production without impacting cell proliferation. Moreover, there is usually an extended regulatory region upstream of NOPs, which contains multiple TFBSs, and transcriptional engineering through modification of TFBSs has been demonstrated to be an effective strategy to optimize promoter strength and regulation in eukaryotic cells [16,17].

The AOX1 promoter of Pichia pastoris is the typifier of NOP, which was identified from the methanol utilization pathway and widely used for protein production with extraordinary strength and tight regulation [18]. Inspired by this, we noted the possibility of ethanol as a supplementary carbon source and inducer for protein production in C. glutamicum. First, ethanol as a carbon source is consistent with the biofinery concept due to its unlimited availability and environmental friendliness. Second, C. glutamicum is able to use ethanol as the sole carbon source with an appreciable biomass yield [19]. Third, the utilization of ethanol in C. glutamicum has a delayed effect. Biphasic growth behaviour could be observed on mixed carbon source medium, which was caused by the sequential utilization of glucose before ethanol [19]. Finally, unlike the “permeability repression” of inducers such as isopropyl-l-thiogalactopyranoside (IPTG) and l-arabinose [20], ethanol, as an amphipathic molecule, is freely permeable across bacterial membranes.

In this work, we reanalysed the promoter of isocitrate lyase (ICL), the key enzyme triggering the ethanol utilization (EUT) pathway of C. glutamicum, and demonstrated its capability as a powerful NOP when a conservative translation initiation structure is intact. Through transcriptional engineering based on TFBSs and UTR (Untranslated Region) modification, we generated an enhanced ethanol-activated promoter P_{A256}, which maintains low activity in early growth but exhibits extraordinary strength during the ethanol utilization phase. By comparing the activity of P_{A256} in various EUT-blocking strains, we found that the activation of acetyl-CoA by acetyl phosphate is the key premise for the upregulation of P_{A256} on ethanol. Finally, efficient protein production with P_{A256} over the EUT pathway was successfully demonstrated with three recombinant proteins: VHH, XynA and NEO-2/15.

2. Materials and methods

2.1. Bacterial strains and growth conditions

E. coli DH5α was used as a cloning host for plasmid construction and cultivated in Luria–Bertani (LB) medium (per litre 1% tryptone, 0.5% yeast extract and 1% NaCl), which was supplemented with 30 μg/mL chloramphenicol or 50 μg/mL kanamycin when necessary. C. glutamicum GCMCC 1.15647 (GenBank: CP073911.1) was used for the construction and analysis of expression systems and cultivated at 30 °C in LBB medium (LB supplemented with 1% brain heart infusion, 10 μg/mL chloramphenicol and 25 μg/mL kanamycin when necessary). Shake-flask fermentation (10 mL medium in each 100-mL shake flask) and 24-deep-well plate fermentation (1 mL medium in each 5-mL hole) were carried out to assess the protein expression of different constructs. Absolute ethanol was added directly to the LBB medium by volume ratio when needed. Noncolony-type monoclonal culture (NCM) medium supplemented with cell wall weakening agents was used to make competent cells [21]. LBBhis medium (5 g/L tryptone, 2.5 g/L yeast extract, 18.5 g/L brain heart infusion broth, 91 g/L sorbitol and 5 g/L NaCl) was used for transformation of C. glutamicum.

2.2. Protein extraction and analysis

For intracellular protein analysis, the culture of C. glutamicum (adjusted to 1 mL, OD_{600}~10.0) was harvested by centrifugation and washed twice with PBS, then resuspended in 1 mL PBS. Cell lysis was carried out using Ultrasonic breaker Vibra cell VCX500 (SONICS) and lysis supernatant after centrifugation was used for SDS-PAGE analysis. For extracellular protein analysis, after centrifugation, the medium supernatant was collected and directly subjected to denaturation (25 μl supernatant+5 μl protein loading buffer, 98 °C, 8 min), with 15 μl denatured sample loaded on each lane for SDS-PAGE analysis.

The protein band of interest in the ethanol-induced sample was extracted as granules (1 mm), which were then destained in decolorizing buffer and lyophilized at 4 °C. The lyophilized sample was immersed in porcine trypsin solution and incubated within NH_{4}HCO_{3} buffer at 37 °C for 12 h. MALDI-TOF-MS was applied to analyse digested fractions from the SwissProt database to identify the target protein.

Western blotting assays were performed to confirm NEO-2/15 (6× histidine tag at the C-terminal) expression and secretion in C. glutamicum. The proteins of the identical SDS–PAGE gel were electrothoretically transferred onto a Polyvinylenidenfluoride (PVDF) membrane. The membrane was washed with TBST and incubated in blocking buffer (TBST containing 5% nonfat milk powder) for 2 h. Then, the PVDF membrane was incubated in antibody solution (monoclonal horseradish peroxidase-conjugated anti-His6 antibody) for 1 h and washed three times with TBST. Finally, the binding of the antibody was detected using an ECL kit and imaging system.

2.3. Promoter reporter vectors and protein expression vectors

The main strains and plasmids used in the study are listed in Table S1. The important primers and oligonucleotides are listed in Table S2. Tag or KOD-plus-neo polymerases (Toyobo) were used for PCR. The shuttle vector p19-0 [22] was the backbone for all expression vector construction. Genomic DNA of C. glutamicum and Corynebacterium ammoniagenes ATCC 6872 were extracted using a TIANamp Bacteria DNA Kit and used as a template to amplify the promoter fragment. For enhanced green fluorescent protein (eGFP) expression, the promoter fragment and eGFP fragment (additional SD sequence if necessary) were integrated into p19-0 by Gibson assembly [23]. For sfGFP expression, the plasmid p19-RiboJ-sfGFP containing the spacer, RiboJ, RBS and sfGFP sequences was constructed, and the XhoI restriction site between spacer and RiboJ was used for the insertion of promoter variants by Gibson assembly. To generate VHH- and XynA-expressing plasmids, the codon-optimized vhh (GenBank: MZ622246) and xynA (GenBank: MZ622248) for C. glutamicum were PCR-amplified with a C-terminal tetracycline tag and cloned into the p19-0 vector with the promoter-RiboJ-RBS fragment and signal peptide fragment. To generate NEO-2/15-expressing plasmids, the P_{A256}-RiboJ-RBS fragment and codon-optimized neo fragment (6His-tag*, GenBank: MZ545408) were integrated into the p19-0 vector with various N-terminal peptide fragments. All plasmids were confirmed by Sanger sequencing and introduced by electroporation into competent C. glutamicum cells according to protocols described previously [21].

2.4. Quantification of protein expression levels

To quantify the expression levels of eGFP/sfGFP in C. glutamicum, cells cultured for 24/36 h were harvested, washed and resuspended. Cell suspensions were diluted to remain within the linear range of the plate reader (1:40-fold, i.e., Fifty microlitres of cell suspension was added to 1950 μl of ddH_{2}O, 200 μl for fluorometry and 1000 μl for OD measurement. The fluorescence intensity of each sample was measured using
2.4.2 Construction and validation of the all-in-one CRISPRi system

To construct a single plasmid CRISPRi system, the codon-optimized dCas9 gene (GenBank: MZ622250), terminator and sgRNA expression cassette were chemically synthesized as one fragment and cloned into the pEC-X998 vector. The sgRNA expression cassette contained a PtcR sequence without the 5′UTR, a 20 bp sgRNA coding sequence, and a sgRNA scaffold (dCas9 handle and S. pyogenes terminator) sequence. Two BsaI restriction sites separated by spacers (as exemplarily shown in Fig. S5) allowed easy insertion of sgRNA fragments using the Golden Gate method [24]. For sgRNA design, N20NGG motifs were searched in the template strand of the target gene, and those located within the first 150 bp of the open reading frame (ORF) were selected as candidates. Off-target effects were checked in the C. glutamicum CGMCC 1.15647 genome using the SeqMap package [25]. Candidates with a tolerance setting of five mismatches and 40–60% GC content were used for interference experiments, which were obtained by annealing two oligonucleotide chains. Nontargeting sgRNAs were designed by random generation of N20NGG with the proper GC content, and those with more than two mismatches within 12 bp of the PAM region were selected.

To repress the expression of the egfp gene, a sgRNA targeting -32 position of the ORF was designed, and the corresponding CRISPRi plasmid was transformed into P_{ICL}-B constructs. The blank vector without sgRNA was also transformed as a negative control. Fluorescence was characterized as mentioned above, and intracellular expression of egFP was observed through fluorescence microscopy. We further tested the reliability of our system by targeting the CRISPRi machinery to porB, ppg, and pyk (three genes had been knocked down or knocked out in other studies) as a control. The results showed that in knockdown mutants cultivated in ethanol, the corresponding band was missing (Fig. 1a, Lane 4). Meanwhile, knockdown mutants exhibited a defect in ethanol utilization (Fig. 1b), which is consistent with a previous report [19] that ICL is required for the growth of C. glutamicum on ethanol. These results suggested that aceA exhibits the most visible changes in protein abundance upon shifting to ethanol as a carbon source; thus, we refocused our attention on its native promoter P_{ICL}. The gene aceA and the malate synthase gene aceB are transcribed in opposite directions, forming a single operon [29], and the mRNA levels of aceA and aceB were 6-fold and 7-fold higher, respectively, in ethanol-grown cells than in glucose-grown cells [19]. However, in the updated study [30], RNase E/G breakage was detected in the 3′UTR of mature aceA RNA (approximately 3-fold higher mRNA level and 4 times longer half-life of the aceA gene in mrgE knockout mutants), indicating that the practical activity of P_{ICL} is greater. Given that the endogenous 3′UTR is not employed in the expression of foreign genes, we speculated that P_{ICL} would be an appropriate choice for building the recombinant protein manufacturing platform.

Using egFP as a reporter, we evaluated P_{ICL} activity under three different strategies. First, the 483 bp cis-regulatory region (called P_{ICL-M}) upstream of the start codon ATG was intercepted and directly followed by the egFP gene; however, only a few fluorescent signals were detected. The RBS Calculator (predict mode, version: v2.1.1) [31] indicated that the translation initiation rate of this architecture was extremely low (only 2.06, au). Hence, P_{ICL-U} was constructed by replacing the endogenous 5′UTR with the 5′UTR from monocistronic Ptac [32] to ensure effective translation initiation. Furthermore, the bicistronic design (BCD) expression cassette, a ‘context preservation’ strategy often employed to repair NOPs [33,34], was used to preserve the original translation initiation structure and improve expression via the translational coupling effect [35]. For this, 62 bp from the N-terminal coding region of ICL and additional SD2 were inserted as fore-cistrons (encoding a 25-amino-acid peptide), and the resulting bicistronic promoter was designated P_{ICL-B} (Fig. 1c). Given the increased biomass from ethanol utilization, the relative fluorescence unit (rfu) of each construct was divided by the corresponding OD_{600} to normalize the expression of fluorescent protein. As the results showed (Fig. 1d), when grown with 1% ethanol added, both P_{ICL-U} and P_{ICL-B} constructs showed obvious fluorescence, with the latter being 3.5-fold higher, in P_{ICL}-B constructs, the egFP expression level after induction was 10 times higher than that in the uninduced state. Overall, after translational structural adjustment, the NOP of isocitrate lyase displayed significant.
activity for heterologous protein overexpression and tight regulation by ethanol.

3.2. Transcriptional engineering of the isocitrate lyase promoter

The transcription start site (TSS) of P_{ICL} has been reported to be located −112 bp upstream of the start codon ATG, where a bidirectional core promoter (separate −35 elements and an overlapping −10 element) and a TG trimer were found nearby [36]. Furthermore, P_{ICL} has been demonstrated to be regulated by the global transcription factors RamA, RamB, and GlxR, similar to the promoters of other central metabolism genes [37]. Six RamA binding sites (Abs) were found in the upstream regulatory sequence of P_{ICL} [38] and a RamB binding site (Bbs) was located between the 4th Abs and the 5th Abs [39]. In addition, a binding site of GlxR (a global TF of the cAMP receptor protein family) [40] closely follows the 3rd Abs (Fig. 2a, Text S1). To investigate the effect of various TFBSs on promoter activity, we created a series of promoters with truncated upstream regulatory sequences of 486 bp, 371 bp, 301 bp, 256 bp, 234 bp, 223 bp, 205 bp, 184 bp, 166 bp, 116 bp, 87 bp, 72 bp, and 59 bp (before the TSS) while keeping the native UTR, which we called P_{A256}UTR. Considering the abovementioned negative impact of the native UTR on translation efficiency and to ensure that all modifications act at the transcription level, the insulator Riboj for generating uniform transcripts for sfGFP expression (Fig. 2b). The strength of these truncated variants was compared to the powerful synthetic promoter H36 (P_{H36}, with Riboj) under ethanol and nonethanol conditions (culture time t = 24 h and t = 36 h, respectively). After induction with 1% ethanol, the specific fluorescence of full-length P_{ICL} (P_{A486}UTR) increased 2.85-fold compared with the uninduced state but was approximately 10% lower than that under the P_{H36} promoter (Fig. 2c). The promoter strength did not change significantly when shortened from 486 to 301 bp, demonstrating that there is no effective cis-acting element in this scope. Surprisingly, after removing the 1st Abs (i.e., P_{A256}UTR), the expression level was 3.62-fold greater than before it was eliminated. Despite the fact that sfGFP expression with P_{A256}UTR increased dramatically in the absence of ethanol, identical induction ratios (2.78-fold) were maintained. When the 2nd Abs (P_{A234}UTR) was deleted, the promoter’s activity was drastically reduced, and subsequent truncated versions maintained a very low expression level. No additional inducing activity was observed when the Bbs (P_{A256}UTR, P_{A234}UTR) were removed. Overall, we investigated the significance of several TFBSs in P_{ICL} transcription by truncating the upstream regulatory area, finding that the most distant RamA binding motif (1st Abs) to the TSS has a significant inhibitory effect, but the 2nd Abs has a positive effect on transcription. The P_{A256}UTR promoter generated by removing a hammerhead ribozyme and downstream hairpin [41] was introduced to generate uniform transcripts for sfGFP expression (Fig. 2b). The strength of these truncated variants was compared to the powerful synthetic promoter H36 (P_{H36}, with Riboj) under ethanol and nonethanol conditions (culture time t = 24 h and t = 36 h, respectively). After induction with 1% ethanol, the specific fluorescence of full-length P_{ICL} (P_{A486}UTR) increased 2.85-fold compared with the uninduced state but was approximately 10% lower than that under the P_{H36} promoter (Fig. 2c). The promoter strength did not change significantly when shortened from 486 to 301 bp, demonstrating that there is no effective cis-acting element in this scope. Surprisingly, after removing the 1st Abs (i.e., P_{A256}UTR), the expression level was 3.62-fold greater than before it was eliminated. Despite the fact that sfGFP expression with P_{A256}UTR increased dramatically in the absence of ethanol, identical induction ratios (2.78-fold) were maintained. When the 2nd Abs (P_{A234}UTR) was deleted, the promoter’s activity was drastically reduced, and subsequent truncated versions maintained a very low expression level. No additional inducing activity was observed when the Bbs (P_{A256}UTR, P_{A234}UTR) were removed. Overall, we investigated the significance of several TFBSs in P_{ICL} transcription by truncating the upstream regulatory area, finding that the most distant RamA binding motif (1st Abs) to the TSS has a significant inhibitory effect, but the 2nd Abs has a positive effect on transcription. The P_{A256}UTR promoter generated by removing...
the 1st Abs displayed extraordinarily high sfGFP expression, which was
2.43-fold greater than that achieved with P\textsubscript{H36} on 1% ethanol.

Another key element determining promoter transcriptional activity is RNA polymerase escape efficiency, which is influenced by both the core promoter and the first transcribed sequence \cite{42}. The core region and the native UTR of P\textsubscript{A256}UTR were shortened for this purpose to study their impact on transcription. A series of forward and reverse truncated variants surrounding the TSS were generated and evaluated for sfGFP expression. As the results showed (Fig. S1), different degrees of 5’UTR retention resulted in significantly varied transcriptional activity, with P\textsubscript{A256}U20 exhibiting the highest expression level, approximately 1.45-fold higher than P\textsubscript{A256}U0 and 2.59-fold higher than P\textsubscript{H36}. Therefore, retaining the 20 bp endogenous UTR is required to enable the optimum activity of P\textsubscript{CIL}. Furthermore, removing the entire native UTR, including the −35 element responsible for aceB transcription start (P\textsubscript{A256}U0 constructs), only resulted in a minor decrease in promoter activity compared to P\textsubscript{A256}UTR, demonstrating that native UTR and bidirectional transcription structure are not essential for P\textsubscript{CIL}-oriented transcription. The TG dimer near the −10 element, on the other hand, has been reported to significantly increase promoter activity in C. glutamicum \cite{43}, but our results showed that the TG trimer of P\textsubscript{CIL} has no significant effect on its transcriptional activity because similar sfGFP expression levels between the P\textsubscript{A256}U0 and P\textsubscript{A256}M20 constructs were observed. As expected, the removal of the −10 element prior to the TG trimer (i.e., P\textsubscript{A256}M20) caused the deactivation of the promoter. We attempted to further minimize the redundant sequence of the regulatory region using internal deletions on the basis of P\textsubscript{A256}U20. However, eliminating the in-between sequence of Gbs and the 3rd Abs or deleting the internal Abs and Bs motifs severely impaired promoter activity (Fig. S1). As a result, in all subsequent tests, the P\textsubscript{A256}U20 structure (hereafter abbreviated as P\textsubscript{A256}) was employed.

3.3. Expression capabilities and limitations of P\textsubscript{A256}

To better understand the protein overexpression controlled by P\textsubscript{A256} on ethanol, the growth and sfGFP expression curves of P\textsubscript{A256}-sfGFP constructs were examined (Fig. S2a). The results showed that, whether ethanol was given or not, P\textsubscript{A256}-sfGFP constructs had significantly lower expression levels than those under the P\textsubscript{H36} promoter in the early growth period (3 h, \(P < 0.01\)) and on plates (Fig. S3). Furthermore, diauxic growth could be observed with 1% ethanol added, and the sfGFP expression level with P\textsubscript{A256} dramatically increased in the second stage (after 6 h). Thus, P\textsubscript{A256} provides a better production mode with both a proliferation phase and a production phase, which is critical for the overexpression of heterologous proteins, particularly toxic proteins. To find the best ethanol concentrations for the induction of P\textsubscript{A256} promoter, several native promoters of corynebacteria were cloned and integrated into the p19-RiboJ-sfGFP vector, and their expression levels were tested under ethanol and nonethanol conditions. As a result, in all subsequent tests, the P\textsubscript{A256}U20 structure was employed.

Fig. 2. Transcriptional engineering of the isocitrate lyase promoter. (a) The cis-acting elements characterized in the upstream regulatory sequence of P\textsubscript{CIL}. The truncated position is indicated by a black arrow. The TSS is denoted as +1, and the native UTR is highlighted in yellow. The start codon is marked in red. (b) Schematic diagram of expressing sfGFP with RiboJ. All truncated promoter and control promoter sequences were inserted upstream of RiboJ. Under any promoters inputting, ribozyme RiboJ cleaves internally, to remove the 5’UTR sequence from the promoter, thus forming the same mRNA for sfGFP translation. (c) The promoter strength of truncated P\textsubscript{CIL} promoters under ethanol and nonethanol conditions. The sfGFP expression was conducted in shake flasks for 24 h (0% ethanol) and 36 h (1% ethanol). Values and error bars reflect the mean ± s.d. of three biological replicates (\(n = 3\)). \(* P < 0.05\), \(* * P < 0.001\), Student’s two-tailed t-test.
3.4. Activity of \(P_{A256}\) in the EUT blocking strains

After passing through the cell membrane, ethanol is oxidized via acetaldehyde to acetate, which then enters the TCA cycle after being activated to acetyl coenzyme A (Fig. 3a). To identify the key steps determining the upregulation of \(P_{A256}\), we attempted to disrupt each step by interfering with key genes in the EUT pathway. We created a single plasmid CRISPRi system (Fig. S5) on the pECXK99E vector (pGA1 replicon) for this purpose, which is compatible with the A256 expression system established on the pBL1 replicon-derived vector. To control dCas9 expression, the LacI\(^{-}\) repressor system was employed, and the sgRNA expression cassette referred to bacterial CRISPRi system design guidelines [44]. To investigate the efficacy of CRISPRi for gene repression in \(C.\) glutamicum, four genes were chosen as targets (egfp in the \(P_{ICL-8}\) construct and the endogenous genes \(porB, pgi,\) and \(pyk\)) and were all effectively knocked down (Fig. S5).

Based on the foregoing results, we established CRISPRi system dependability and used it to block the EUT pathway in \(P_{A256}\)-controlled sfGFP-expressing strains. We designed one sgRNA for each of the six key genes involved in the EUT pathway, and three nontargeting sgRNAs were utilized as negative controls (Fig. 3b). Each sgRNA was integrated into the CRISPRi plasmid and cotransformed into wild-type strains with the \(P_{A256}\)-sfGFP plasmid. The growth and sfGFP expression of these constructs were evaluated in ethanol and nonethanol environments (Fig. 3c). The results indicated that most knockdown strains grew normally under nonethanol circumstances, with the exception of the ALDH knockdown strains, which had 18.1% less biomass than the control strains. Under ethanol conditions, the knockdown strains exhibited varying degrees of growth impairment (the final biomass was lowered to 44.2–90.6% of the control strains). Among them, the ALDH knockdown strains showed similar growth levels as when no ethanol was given; i.e., the biomass growth due to ethanol addition was reduced from 110.9% to 44.2\(^{-}\). An interesting finding is that the suppression of ALDH caused the strains to lose their capacity to use ethanol almost entirely. Except for the PTA knockdown strains, which had 18.1% less biomass than the control strains under ethanol conditions (a loss of 96% of expression relative to the control strains), although it operates effectively when ethanol is not present. It should be noted that the genes encoding AK and PTA (\(ack\) and \(pta\), respectively) form a bicistronic operon, with \(pta\) upstream of \(ack\), suggesting that there might be a forward polar effect, i.e., targeting the CRISPRi complex to the ORF of \(pta\) is likely to inhibit \(ack\) transcription. As a consequence, \(P_{A256}\) inactivation might be an independent impact of PTA knockdown, or it could be the result of simultaneous suppression of PTA and AK. At the very least, the current data suggest that the elevated activity of \(P_{A256}\) is the consequence of ethanol metabolism, particularly when ethanol is metabolized into the TCA cycle.

3.5. High-level production of secretory proteins with \(P_{A256}\)

To demonstrate the adaptability of \(P_{A256}\) for proteins with a larger range of molecular weights, we applied it to the production of two secretory proteins, VHH (15 kDa) and XynA (47 kDa), both of which have achieved high yield production with the \(P_{H36}\) promoter [45]. To monitor and compare the yield of the target protein in the culture supernatant, we applied the FlAsH-tetracysteine system to our experiments. To this end, an optimized biosensorically binding tetracysteine motif (\(-FLNCPCGCCMEEP-\)) [46,47] was genetically fused to the C-terminus of the target proteins (Texts S2 and S3), which was specifically recognized by the fluorescein derivative with two As(III) substituents, FlAsH-EDT\(_2\) [48], and the fluorescence produced corresponded with the content of labelled proteins (Fig. 4a). The reliability of this approach was demonstrated by inducible \(P_{A256}\)-controlled VHH secretory expression (Fig. S6). For the secretory expression of VHH and XynA, the endogenous signal peptide of CspB [49] (spCspB) was used, and several RBSs with different predicted translation initiation rates were tested to ensure translation efficiency with spCspB as the N-terminal sequence (Fig. S7). With the optimal RBS, we constructed spCspB-guided VHH- and XynA-expressing strains with \(P_{A256}\) and \(P_{H36}\), and the secretory expression under the control of these two promoters was compared on 1% ethanol. In the case of VHH, the target protein band in \(P_{H36}\)-VHH constructs appeared at 6 h but was invisible in the supernatant of \(P_{A256}\)-VHH constructs at the same time (Fig. 4b). The fluorescence intensity from the

![Fig. 3.](image-url)
FlAsH-tetracysteine reaction confirmed this result. This result indicated that the secretory protein expression controlled by \( P_{A256} \) was low during the early fermentation stage, which was compatible with the results of the intracellular sfGFP expression assay. The yield of VHH with \( P_{A256} \) surpassed that with \( P_{H36} \) after 12 h and remained higher until the end of fermentation. In XynA production, a lower preaccumulation and greater final yield expression pattern were also observed. (Fig. 4c). These results demonstrated that the A256 expression system provides a superior model for recombinant protein production in \( C. glutamicum \).

3.6. Secretory production of NEO-2/15 using the A256 expression system

We also applied the A256 expression system to overexpress the new drug protein Neo-2/15 (Neo). As a “de novo design” protein, Neo mimics the effects of the central immune cytokines interleukin-2 and interleukin-15 [50]; to date, there has been no attempt to overexpress it in a secretory form. Initially, we attempted to secrete Neo using the Sec-type secretory signal peptide of CspB and Tat-type secretory signal peptide of CgR0949 [51], but only intracellular soluble NEO was detected in these two constructs (Fig. 5b). This indicated that NEO cannot be secreted through the Sec or Tat pathway only by the guidance of signal peptides. According to recent research, preprotein mature domains not only operate as passive passengers but also contain translocone targeting signals that are essential for secretion [52,53]. Therefore, we devised a novel approach to induce NEO secretory expression, i.e., fusion with a portion of the endogenous secretory protein CspB (Fig. 5a). For this purpose, NEO was genetically fused with the N-terminal 50 amino acid residues of CspB (namely CspB50, Text S4). As the results showed, despite the intracellular retention (target protein with the uncleaved signal peptide), the fused protein CspB50-NEO (C50-NEO, ~19 kDa) could be detected in the culture supernatant (Fig. 5b). Then, the secretory production of C50-NEO controlled by \( P_{A256} \) was evaluated by shaking-flask fermentation under the concentration of 1–2% ethanol (Fig. S8), and the greatest accumulation of C50-NEO (281 mg/L) was achieved after 24 h cultivation with 1% ethanol added (Fig. 5c). To the best of our knowledge, this is the first report of recombinant protein secretion by fusion of the CspB protein in \( C. glutamicum \).

4. Discussion

Due to its numerous ideal protein secretion properties, \( C. glutamicum \) has attracted attention in recent years as a potential cell factory for recombinant protein production. However, protein production with \( C. glutamicum \) is currently based on sugar-containing medium, which is incompatible with the biorefinery concept of converting renewable biomass to multiple high-value products [54]. Hence, the feasibility of using ethanol as a substrate for recombinant protein production is discussed here. To achieve efficient production of recombinant proteins on ethanol, we were inspired by the regulation of the MUT pathway and the AOX1 promoter in \( P. pastoris \) and identified the promoter of isocitrate
lyase as the representative NOP triggering the EUT pathway in *C. glutamicum*. Although the native 5′UTR of P<sub>ICL</sub> resulted in a very low translation initiation efficiency when expressing heterologous proteins, it could be repaired by bicistronic design or the introduction of a ribozyme-based insulator. Since self-cleavage ribozyme can remove the different upstream 5′ UTRs, transcriptional levels could be compared between promoter variants and positive control promoters. Based on this, transcriptional engineering was performed on the natural P<sub>ICL</sub>, including engineering of TFBSs and UTR, and resulted in a particularly strong promoter, P<sub>A256</sub>. Removal of the most distant RamA binding motif conferred a significantly increased transcriptional strength, despite the fact that RamA was identified as the activator of P<sub>ICL</sub> [55], which indicated that the location of TFBS may correlate with its functionality. In addition, the nondeletability of other TFBSs in P<sub>A256</sub> was validated in our work; thus, duplication and rearrangement of existing TFBSs might be a possible strategy to further improve its transcriptional level in the future.

The protein expression controlled by P<sub>A256</sub> went through three stages: 1) a very low expression level on plate and in the early growth period; 2) an increased expression level during the exponential growth phase with or without ethanol addition; and 3) a dramatically elevated expression level in the secondary growth driven by ethanol. Such an expression curve implies a lower burden on cells during proliferation, whereas higher amounts inhibited growth and protein expression. Tolerance to ethanol can be achieved in bacteria by altering the membrane lipid composition, such as increasing the length of fatty acid chains or the amount of nonpolar lipids [56]. Therefore, generating *C. glutamicum* strains with increased ethanol tolerance by genetic modification or adaptive evolution may be a viable option to further harness the benefits of the A256 expression system.

CRISPRi technology has been successfully applied to optimize the production of various metabolic intermediates and analyse gene function in *C. glutamicum* [57,58]. In this work, a single plasmid CRISPRi system was developed to downregulate the genes encoding enzymes involved in ethanol metabolism, and the performance of P<sub>A256</sub> was investigated in these EUT blocking strains. Ethanol metabolism in *C. glutamicum* begins with a two-step oxidation to acetate, catalysed by ADH and an ALDH, and proceeds via acetyl activation by AK and PTA and the glyoxylate cycle with ICL and MS as key enzymes. Among them, knocking down the PTA coding gene dramatically reduced P<sub>A256</sub> activity under ethanol conditions. This suggests that the PTA-catalysed step, i.e., acetyl phosphatase activated to acetyl-CoA and entering the TCA cycle is a necessary prerequisite for the upregulation of P<sub>A256</sub> on ethanol. Whereas P<sub>A256</sub> is regulated by multiple global transcription factors and the effectors of which have yet to be identified, the exact regulatory mechanisms are a mystery that need more investigation.

By introducing the FlAsH-tetracysteine system, the performance of
different promoters in the secretory expression of various proteins can be easily compared. Thus, except for the intracellular fluorescent proteins, the high-efficiency production of two secretory proteins, VHH and XynA, also confirmed the advantages of the A256 expression system. Furthermore, the “de novo design” protein NEO-2/15 was successfully secreted into the medium by introducing a propeptide at the N-terminus, and a yield of 281 mg/L was obtained at the flask level using an A256 expression system. In conclusion, the ethanol-regulated A256 expression system showed significant potential in recombinant protein production, and we anticipated that it could be further developed through scale-up and fermentation process optimization.

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Data availability
The datasets generated and analyzed during the current study are available from the corresponding author on reasonable request.

Author contributions
XY designed and performed most experiments. XL, XG and XL contributed to experimental planning and data analysis. All authors read and approved the final manuscript.

Ethical statement
This article does not contain any study with human participants or animals performed by any of the authors.

Declaration of competing interest
The authors declare that they have no conflict of interest.

Appendix A. Supplementary data
Supplementary data to this article can be found online at https://doi.org/10.1016/j.synbio.2020.03.004.

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