Internal promoters of the epothilone biosynthetic gene cluster and their activation in Myxococcus xanthus by CRISPR-dCas9

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Research

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Abstract

Background

Multiple genes involving in a complex pathway are often clustered into a giant operon with no transcription terminator before the end, and this leads to frangibility of the transcriptional process and arduous engineering work to control the transcription of operon genes. Internal promoters might occur in operon to coordinate the transcription of individual genes, but their effects on the transcription of operon genes have been less investigated.

Results

Epothilones are a kind of polyketides synthesized by seven multifunctional enzymes, which are encoded by a 56-kb operon of the myxobacterium *Sorangium cellulosum*. In this study, we determined that the epothilone operon contained multiple internal promoters. These promoters were activatable by the CRISPRa technique, and the yields of epothilones were accordingly increased. However, the activation efficiencies of promoters in operon and separate forms were greatly different. Further, we found that the transcriptional levels of the epothilone genes were always increased at a greater extent than the epothilone yields, which suggested that the transcriptional activation of single genes probably had a weak effect on the final epothilone yield, and higher yield required an overall transcriptional increase of the multiple operonic genes. Finally, we combined the activation of the starting promoter $P_{epoA}$ together with internal promoters in different epothilone-producing strains, and obtained the highest 15-fold increase of epothilone yield in *Myxococcus xanthus* ZE5.

Conclusions

This is the first time to report the internal promoters in epothilone gene clusters in *Myxococcus xanthus* and the first time to assay the activation effects of these internal promoters by CRISPR-dCas9. Our results highlight that tuning internal promoter activities is critical to control the transcription of operon genes and the production efficiency of microbial secondary metabolites.

Background

Operons, a well-known feature of prokaryotic genomes, are clusters of co-regulated genes with related functions [1]. Big operons, such as pathways for the biosynthesis of secondary metabolites, may contain multiple genes with no transcription terminator before the end. The transcription of multiple genes in operon is initiated by the starting promoter, forming a single polycistronic mRNA. However, the transcription of big mRNA molecules is easily subject to various influences in cells, leading to frangibility of the transcriptional process and arduous engineering work to control the transcription of operon genes. Internal promoters have been observed in operons for many years. According to experiments and genome prediction, internal promoters are universal in operons in different bacteria [2–7]. For example, Ma et al. demonstrated that there are at least three weak internal promoters $P2$, $P3$ and $P4$ in addition to two strong promoters $P_L$ and $P1$ in the *E. coli rpoBC* operon, which encodes four ribosomal proteins and the $\beta$ and $\beta'$ subunits of RNA polymerase [2]. In the 57-kb jamaicamide operon from the marine cyanobacterium *Lyngbya majuscula*, 17 genes are co-transcribed from the starting promoter, and six internal promoters are present in intergenic regions, which potentially assist management of the toxin production in various environments [3].
Similarly, seven internal promoters have also been observed in the microcystin operon of *Microcystis aeruginosa* [4]. However, the effects of internal promoters on the transcription of operon genes have been less investigated.

Epothilones are a kind of antitumor polyketide compounds with the microtubule-stabilizing activity, which were originally discovered in the extracts of some *Sorangium cellulosum* strains [8]. In 2000, Molnár and Julien reported the sequence and construction of the epothilone gene cluster, which is approximately 56 kb, containing seven open reading frames (ORFs) in the same transcriptional direction [9, 10]. These ORFs encode polyketide synthases and a non-ribosomal peptide synthetase with multiple functioning domains for elongation, modification and release of epothilones in a sequential mode [9]. This gene cluster contains no terminator between ORFs before the end, thus forming a giant operon.

In previous work, we have successfully integrated the epothilone biosynthetic gene cluster from *S. cellulosum* So0157-2 [11] into different sites of the *Myxococcus xanthus* genome, and produced dozens of epothilone producers [12]. We found that the transcriptional levels of operon genes greatly varied in either *M. xanthus* or the original *S. cellulosum* producers. We constructed a CRISPRa (CRISPR-dCas9-mediated transcription activation) system in *M. xanthus* and have successfully improved the production of epothilones by activating the starting promoter of the operon [13]. However, the transcriptional levels of operon genes still varied. In fact, uneven gene expression of operon genes was observed many years ago [14], and genome-wide transcriptomics studies have also revealed varied transcriptional levels of the consecutive genes within operons [15], which suggest that the operon genes are complexly regulated, rather than transcribed from the single starting promoter. Internal promoters are suggested to coordinate the transcription of operon genes, but their effects remain mysterious. In this study, we determined the existence of multiple internal promoters in the epothilone biosynthetic gene cluster. We employed the established CRISPRa technique to promote the transcription of internal promoters, and found that the activation efficiency of promoters in operon was distinct from that of them in separate form. We combined the activation of the starting promoter PepoA together with internal promoters in different epothilone-producing strains, and obtained the highest 15-fold increase of epothilone yield in *Myxococcus xanthus* ZE5. Our results highlight that coordinating transcriptional activities of internal promoters is critical for the transcription of operon genes and the production efficiency of microbial secondary metabolites.

**Results**

1. **The epothilone gene cluster is a big operon containing multiple internal promoters**

The epothilone gene cluster contains seven ORFs (*epoA* to *epoF*) in the same transcriptional direction, which are separated with a short distance (15 ~ 147bp) or shortly overlapped (Fig. 1a). No terminator structure exists in the intergenic regions of these ORFs, which means that the epothilone gene cluster is a huge operon. The continuous transcription of the gene cluster was also proved by co-transcriptional detection of two adjacent ORFs (Figure S1). However, these operon ORFs exhibited significant differences in their transcriptional levels. For example, as determined in *M. xanthus* ZE9, an epothilone-producer constructed with the whole gene cluster of *S. cellulosum* So0157-2 [12], the last ORF *epoF* showed the highest expression level by approximately 2.5 times of that of the first ORF *epoA* (Fig. 1b). We suggested that internal promoters might be present in this operon to tune the transcription of individual ORFs.
We predicted the existence of internal promoters in the epothilone operon derived from *S. cellulosum* So0157-2 using the “Neural Network Promoter Prediction” program (https://www.fruitfly.org/seq_tools/promoter.html), and found ten potential promoters (shown as red arrows in Fig. 1a). Prediction with “BPRoM” (http://www.softberry.com/berry.phtml) revealed similar results but excavating a new promoter upstream of *epoD* (shown as green arrows in Fig. 1a). Detailed information of the predicted promoters is shown in Table S1. These promoters mostly appeared in the junctional regions between ORFs. Up to now, four *S. cellulosum* strains have been reported of their epothilone biosynthetic gene clusters with the GenBank accession numbers of AF210843.1 in SMP44 [9], AF217189.1 in So ce90 [10] and GU063811.1 in KYC3013 [16], in addition to EU414841.1 in So0157-2 [11]. The sequences of these four epothilone gene clusters exhibited more than 98.6% similarity. In the epothilone operons from other three *S. cellulosum* producers, internal promoters were similarly present but with some differences (Figure S2; detailed information is provided in Table S2 ~ S4).

To analyze activities of internal promoters, we selected six regions that locate in the junction between two ORFs in the So0157-2 epothilone operon. These regions were 1000 bp in length, including the 800-bp fragment of an upstream gene and the 200-bp fragment of a downstream gene, which were amplified from *M. xanthus* ZE9 using the corresponding primer pairs (listed in Table S5). Each region, containing at least one predicted internal promoter, was cloned into the pKK232-8 plasmid to control the expression of chloramphenicol acetyltransferase (CAT) (Figure S3). The *aphII* promoter was constructed upstream of the CAT reporter gene as a positive control. According to the CAT activities, assayed with the CAT ELISA kit, all the six regions exhibited transcriptional activities in *E. coli*. The *P_{epoP}* promoter showed the highest activity, which was close to that of the *aphII* promoter, while the *P_{epoB}* and *P_{epoE}* promoters showed weak activities, slightly higher than that of the negative control (without a promoter before the CAT gene) (Fig. 1c).

Simultaneously, we also analyzed promoter activities of the 1 kb sequence upstream of the translation initiation codon (ATG or GTG) of each ORF in *M. xanthus*. The six fragments were constructed respectively in front of the *EGFP* (Enhanced Green Fluorescent Protein) reporter gene in pZJY41, an autonomously-replicating plasmid in *M. xanthus* [17] (Figure S4). The identified starting promoter *P_{epoa}* in front of the epothilone operon [18] was used as the positive control. We introduced these plasmids into *M. xanthus* DZ2 and assayed the green fluorescence values of the mutants after 24 h, 48 h and 72 h of incubation. The results showed that these fragments, even the sequence upstream of *epoF*, where no promoter had been predicted, exhibited promoter activities; *P_{epoa}* always had the highest activity, and the activity of *P_{epoP}* is the lowest (Fig. 1d). Notably, the detected promoter activities, either in *E. coli* or *M. xanthus*, were significantly inconsistent with the transcriptional levels of the genes in the epothilone operon. For instance, we did not find a strong promoter upstream of the *epoF*, which, however, displayed the highest transcriptional level among the ORFs in the epothilone operon (Fig. 1b). These results suggested that these internal promoters might be complexly regulated of their transcriptional activities in operon.

2. Operon and separate internal promoters exhibit different transcriptional activities

CRISPR (clustered regularly interspaced short palindromic repeat) Cas system is an RNA-mediated immune system existing in many bacteria and archaea to protect cells from foreign DNA invasion [19–21]. Peng *et al.* constructed a CRISPRa system in *M. xanthus* and successfully activated the *P_{epoa}* promoter of the epothilone operon, which increased the transcriptional levels of operon genes and improved the yield of epothilones by 230% [13]. The schematic diagram of CRISPRa construction in *M. xanthus* is shown in Fig. 2a.
We performed the CRISPRa-based activation on separate promoters in *E. coli*. To achieve activation, we transferred three plasmids into the HB101 strain: the pSWcuomxdCas9-ω plasmid carrying dCas9 protein and transcription activator Omega (ω) [13], the pZJY41-sgRNA series plasmid carrying sgRNA, and the pkk232 series plasmid carrying promoter $P_{epo}$ and reporter gene CAT (Figure S5). To construct the pZJY41-sgRNA series plasmids, we employed one spacer for each promoter (refers to Table 1), which were designed using the online software “CasOT” [22]. We found that the transcription activities of these separate promoters were all significantly improved: weak promoters were more easily activated by CRISPRa, and the promoters with high transcription activities were also activated but to low extents (Fig. 2b), which is consistent with the previous report [23]. For example, the transcription activity of the weakest promoter $P_{epoB}$ was increased by nearly 33-fold, while the strongest promoter $P_{epoA}$ was activated by approximately 1.6-fold.

| Name | Spacer | Coding strand | Off-target | Distance(bp) | $\Delta G$ (kcal/mol) | Hairpin | GC% |
|------|--------|---------------|------------|--------------|----------------------|---------|-----|
| CuOm-P2 | TCCGGGGGATGATGCTCGAG | + | 8 | -617 | 43.6 | N | 65.0 |
| CuOm-B3 | TGAGGAGCCTGTTGCAGAAG | + | 14 | -376 | 38.2 | N | 55.0 |
| CuOm-C2 | ACCGTACC GGCAACGCTTGTG | + | 4 | -165 | 45.3 | N | 61.9 |
| CuOm-D2 | TGCGGCCGGTATCCTGGACGA | + | 3 | -117 | 47.8 | N | 66.7 |
| CuOm-E1 | TGGATGTATCCCAAGGTGCT | + | 2 | -160 | 38.1 | N | 50.0 |
| CuOm-F2 | AGCTCTTCTTCCGAAATGCCG | + | 2 | -193 | 43.5 | N | 52.4 |

We further performed in-situ activation on the internal promoters within the epothilone operon in *M. xanthus* ZE9 by using the same spacer sequences for the activation of each of the six internal promoters (Table 1). The spacer sequences were constructed into the pZJY41 plasmid, respectively (Figure S6), producing six series plasmids (Table S6), which were separately introduced into the CuOm strain by electro-transformation. The CuOm strain was constructed by introducing the pSWcuomxdCas9-ω plasmid into ZE9 (a diagrammatic sketch for the construction is shown in Figure S7). The results showed that the transcriptional levels of most of the activated genes were significantly increased, except that the expression of *epoD* in CuOm-D2 was increased insignificantly ($t$-test, $p > 0.05$) (Fig. 2c). The best activation in *M. xanthus* was achieved on the *epoE* gene, which was activated by about 5-fold. Notably, the results were markedly different from that of the activation on the separate promoters although using the same spacer sequences. Thus, the transcription activities of internal promoters were influenced in operon.

In addition, we found that the epothilone production abilities in these CRISPRa-promoted mutants were significantly increased, even including CuOm-D2, in which the expression of *epoD* was not increased significantly by the CRISPRa technique (Fig. 2d). We found that the transcriptional levels of the epothilone genes were always increased at a greater extent than the epothilone yields. For example, the highest gene expression was found in
CuOm-E1 (increased by 5-fold), but the epothilone yield was only increased by 1.7-fold. We thus checked the transcription of each ORF in the epothilone operon with the activation of single internal promoters. The results showed that the activation of the front promoters $P_{epoA}$, $P_{epoB}$ or $P_{epoC}$ normally increased the transcription of the front genes $epoA$, $epoP$, $epoB$ and $epoC$, but did not change or even decreased the transcription of the hind genes $epoD$, $epoE$ and $epoF$, whether the genes were specifically activated or not (Figure S8). Similarly, the activation of the hind promoter $P_{epoD}$, $P_{epoE}$ or $P_{epoF}$ increased the transcription of the hind genes, and sometimes the upstream $epoB$ and $epoC$ genes, but had no effect on $epoA$ or $epoP$. The results suggested that the transcriptional activation of single genes probably had a weak effect on the final epothilone yield, and higher yield required an overall transcriptional increase of the multiple operonic genes.

3. Tuning the activity of multiple promoters to increase the epothilone yield

To improve the activation effects to increase the yield of epothilones, we combined the activation of the starting promoter $P_{epoA}$ together with an internal promoter $P_{epoP}$, $P_{epoB}$ or $P_{epoD}$, thus forming the AP, AB or AD combination. We also combined the activation of three promoters, i.e., APB, APD and ABD. The combined sgRNA sequences were cloned into the pZJY41 plasmids, which were introduced into the CuOm strain, respectively (Figure S9). After the CRISPRa-based activation, the production of epothilones was increased in each combination, and the highest yields were obtained in the mutants CuOm-AP (11.17 mg/L) and CuOm-ABD (11.69 mg/L), both of which increased approximately 2.4-fold, compared with the 4.95 mg/L in the initial strain ZE9 (Fig. 3a). However, the combined activation of two or three promoters did not lead to an accumulation of the activation effects of single promoters.

We analyzed the transcription of each operon ORF under different activation combinations (Figs. 3b and 3c). The results showed that the transcriptional levels of the front operon genes $epoA$, $epoP$ were mostly significantly increased by the combined activation, even when $P_{epoP}$ was not specifically activated (in CuOm-AB, CuOm-AD or CuOm-ABD). However, the transcription of $epoD$, as well as the genes behind, was not increased in CuOm-AD, CuOm-ABD or CuOm-APD, in which $P_{epoD}$ was even specifically activated. Seemingly, the genes close to the starting promoters $P_{epoA}$ were more easily activated than the hind operon genes.

Based on the abovementioned results, we combined CRISPRa-mediated activation on promoters in $M. xanthus$ strains with different transcription levels of the epothilone genes. We previously constructed dozens of epothilone-producing $M. xanthus$ strains, in which the same epothilone biosynthetic gene cluster was inserted in different sites of the DZ2 genome, resulting in varied production abilities of epothilones [12]. We chose four strains ZE9, ZE5, ZE10 and ZE14 to assay the transcriptional efficiency using the CRISPRa technique. Among these four strains, ZE9 had the highest production ability of epothilones, followed by ZE10, then ZE14, and ZE5 exhibited the lowest epothilone yields (Fig. 4a). Consistently, the transcriptional levels of the seven operon ORFs in ZE9 were higher than that in the other three strains (Fig. 4b). In ZE14, the front operon genes were transcribed at higher levels, but $epoD$, $epoE$ and $epoF$ were significantly lower than that in ZE10. Similarly, the transcriptional levels of $epoD$, $epoE$ and $epoF$ were extremely low in ZE5, causing the strain to produce the lowest yield of epothilones among the four strains.

We combined the activations either on the front promoters ($P_{epoA}$, $P_{epoP}$ and $P_{epoB}$) or the hind promoters ($P_{epoD}$, $P_{epoE}$ and $P_{epoF}$) in these epothilone producing strains. As expected, the epothilone yields were all increased in these strains, and the highest 15-fold increase of epothilone yield was obtained in ZE5 with the DEF promoter.
activation (Fig. 4a). Consistent with the yields of epothilones, the highest activation efficiency also occurred in the ZE5 strain. In ZE5-DEF, the transcriptional levels of the three activated genes \( \text{epoD, epoE} \) and \( \text{epoF} \) were increased by 9.6, 3.1 and 51.7 times, respectively, and \( \text{epoP} \) and \( \text{epoB} \) were also increased slightly (Fig. 4c). However, the transcriptional changes of operon genes suggested that the interferences between operon promoters were very complex. For example, in the ZE5-APB strain, the transcriptional levels of the three activated genes \( \text{epoA, epoP} \) and \( \text{epoB} \) were all increased, but to different extents; the transcriptional levels of the four hind genes (\( \text{epoC} \sim \text{epoF} \)), which were not specially activated, were also mostly increased.

**Discussion**

Transcription regulation is always a topic of concern. Operons are clusters of co-regulated genes with related functions. Bacteria have established multiple mechanisms to ensure the relative expressional levels of individual genes in operon to meet the requirements of cell and environment [24, 25]. Transcriptional interference between tandem promoters is recognized as a potentially widespread mechanism to regulate gene expression [26, 27]. There are many studies on the regulation of single internal promoters on the expression of operon genes, and a few studies have been performed on transcriptional interferences between multiple operon promoters with no clear conclusions. For example, the 14-kb \( \text{CAP1} \) gene cluster in \( \text{Staphylococcus aureus} \) is transcriptionally controlled by a strong upstream promoter and five weak internal promoters, and the internal promoters showed significant activity only after removing the primary promoter [28]. In the cyanobacterium \( \text{Anabaena} \) sp. strain PCC 7120, a zinc-responsive operon contains 4 distinct promoters, which were induced by metal depletion, and they were constitutively derepressed in a zur mutant, despite the two downstream promoters not being direct targets for this regulator [29]. In this study, we demonstrated that the big epothilone gene operon contained multiple internal promoters, and the transcriptional processes of these internal promoters may intricately interfere with each other.

Interference between tandem promoters may be generated by dislodgement of slow-to-assemble pre-initiation complexes and transcription factors, or prolonged occlusion by paused RNA polymerases (RNAPs) [29, 30, 31]. The direct and in cis suppression of one transcriptional process by another transcriptional process is that RNAP transcribing from one promoter may have effects on the supercoiling state of neighbor promoters [26,32.33]. A study in the human and mouse genomes concluded that RNAP collisions were the primary mechanism of interaction between transcripts, based upon decreasing transcript abundance with increasing overlap length, to almost zero when the overlap exceeded 2,000 nucleotides [34]. Mathematical modeling also shows that the probability that an RNAP can avoid colliding with an RNAP from a convergent promoter decreases exponentially with the firing rate of the interfering promoter and with the inter-promoter distance [35]. However, the transcriptional interference between tandem promoters is often not satisfactorily explained by RNAP collisions or occlusion by elongating RNAP [27], and there are other unknown regulatory patterns [29].

**Conclusions**

The inconsistency in the transcriptional levels of operon genes often limits the yield of secondary metabolites. The inconsistent expression levels of genes in operon observed in different bacterial species are not only challenging the concept of operons [36,37], but also impeding our engineering work to control the transcription of operon genes. Our results present in this study indicated that, multiple internal promoters are present in epothilone gene cluster. Although little is known of the involving mechanism, regulation of operon internal promoters should be crucial for the biosynthetic pathways of secondary metabolites encoded by a big operon. Tuning the
transcriptional activities of operon promoters, such as using CRISPRa technique, can efficiently improve the metabolite yields.

Materials And Methods

Strains and culture conditions

Strains used in this study are listed in Table S7.

*Escherichia coli* DH5α and HB101 were used for routine transformations and sub-cloning. The *E. coli* strains were grown routinely in Luria Broth (LB) medium (10 g/L peptone, 5 g/L yeast extract, and 5 g/L NaCl, pH 7.2). *Myxococcus xanthus* strains were grown in CYE medium [10 g/L casitone, 5 g/L yeast extract, 10 mM 3-(N-morpholino)propanesulfonic acid (MOPS; pH 7.6) and 4 mM MgSO₄] or CMO medium [10 g/L casitone, 5 g/L yeast extract, 10 mM MOPS, 4 mM MgSO₄, and 7 mL/L methyl oleate, pH 7.6]. When appropriate, kanamycin (40 μg/mL), tetracycline (10 μg/mL) and apramycin (25 μg/mL) were added. The growth temperatures were 37 °C for *E. coli* and 30 °C for *M. xanthus*.

Detection of gene co-transcription of epothilone gene cluster

Total RNA of samples was extracted using BIOZOL kits (Total RNA Extraction Regent, BioFast, China) and then transcribed reversely into cDNA with PrimeScript™ Reagent Kit with DNAase (Takara, Japan). Primers were designed at the junction between two genes. Total RNA without inverse transcription and the plasmid containing the epothilone gene cluster were used as templates in the negative control group and positive control group, respectively. In experimental groups, the cDNAs were used as templates.

Prediction of internal promoters in the epothilone gene cluster

We used the promoter prediction software “Neural Network Promoter Prediction” (https://www.fruitfly.org/seq_tools/ promoter.html) to predict internal promoters in the epothilone gene clusters derived from *Sorangium cellulosum* strains So ce90, SMP44, So0157-2 and KYC3013. The threshold was set as 0.8. At the same time, we used another online promoter prediction software “BPROM” (http://www.softberry.com/berry.phtml? Topic=bprom & group=programs & subgroup=gfindb) to correct predictive results. Bold -35 and -10 binding regions were predicted in comparison to the σ70 consensus -35 (TTGACA) and -10 (TATAAT) promoter regions of *E. coli*. Then we selected the prediction results with a threshold greater than 0.8.

Construction of plasmids

The plasmids and primers used in this study are provided in Tables S5 and S6.

pkk-232-P_{epoP} ~ pkk-232-P_{epoF} were used as a promoter activity reporter vector in *E. coli*. P_{epoP} ~ P_{epoF} were obtained by PCR with primers P_{epoP}-F/R ~ P_{epoF}-F/R from epothilones gene cluster in So0157-2. pkk-232-P_{epoP} ~ pkk-232-P_{epoF} were constructed by inserting P_{epoP} ~ P_{epoF} into the HindIII/BamHI sites of pkk-232-8. pKK-232-aph were constructed by inserting *aph* into the HindIII/BamHI sites of pkk-232-8.

pZJY41-Ap-EGFP ~ pZJY41-Fp-EGFP were used as a promoter activity reporter vector in *Myxococcus xanthus*. P_{epoA} was obtained by PCR with primers Ap-F/R from So0157-2 genome, while report gene *EGFP* was amplified
with primers EGFP-F/R, and then overlap PCR was used to obtain P\textsubscript{epoA}-EGFP with primers P\textsubscript{epoA}-F and EGFP-R. Whereafter, P\textsubscript{epoA}-EGFP was inserted into the BamH\textsubscript{i}/Kpn\textsubscript{i} sites of pZJY41 to construct the reporter vector pZJY41-Ap-EGFP. The other six promoter activity reporter vectors were constructed in the same way.

Plasmid pZJY41-sgRNA was used to express sgRNA. Plasmids pZJY41-sgRNA-P2 ~ pZJY41-sgRNA-F2 were constructed by using p41sg [13] as a template to amplify DNA fragment with primers sgRNA-P2-F ~ sgRNA-F2-F and sgRNA-R, and then linking with T4 DNA ligase to form circular plasmids.

pZJY41-sgRNA-AP was constructed by inserting the sgRNA scaffold with spacer A4 and sgRNA scaffold with spacer P3 into the Nde\textsubscript{i} site and Kpn\textsubscript{i}/BamH\textsubscript{i} sites of pZJY41, successively. pZJY41-sgRNA-AB and pZJY41-sgRNA-AD were constructed in the same way. With sgRNA scaffold with spacerA4 inserted into the Nde\textsubscript{i} site, sgRNA scaffold with spacerP3 inserted into the Kpn\textsubscript{i}/BamH\textsubscript{i} sites, sgRNA scaffold with spacerB3 inserted into the EcoR\textsubscript{i} site of pZJY41, pZJY41-sgRNA-APB was constructed. So were the pZJY41-sgRNA-APD and pZJY41-sgRNA-ABD plasmids.

**Activity detection of internal promoters in *E. coli***

We constructed plasmids to detect the activity of internal promoters from the epothilone operon in *E. coli*. The promoter activity was characterized by detecting the activity of the report gene chloramphenicol acetyltransferase (CAT). The promoter sequences and reporter gene sequence were cloned into plasmid pKK232-8 by digestion with BamH\textsubscript{i}/Hind\textsubscript{i} and then ligation with T4 DNA ligase. The activity of reporter gene CAT was detected by CAT ELISA Kit. The promoter *aphII* was used as a positive control. The CAT ELISA Kit was purchased from Roche and operated according to the instructions provided (https:// www. sigmaaldrich. com/ catalog/ product/ roche/11363727001? Region = CN).

**Activity detection of internal promoters in *Myxococcus xanthus***

We constructed plasmids to detect the activity of internal promoters from the epothilone operon in *M. xanthus*. The promoter activity was characterized by detecting the fluorescence intensity of the green fluorescence reporter gene EGFP. The promoter sequences and the EGFP gene sequence were seamlessly connected by fusion PCR, and finally cloned into the plasmid pZJY41 by digestion with BamH\textsubscript{i}/Kpn\textsubscript{i} and then ligation with T4 DNA ligase. Related primer information was shown in Table S5. The fluorescence intensity of the green reporter gene was detected at 485 nm/528 nm, and three different incubation times (24h, 36h and 72h) were selected for detection.

**Construction of CRISPRa-dCas9 system in *E. coli***

We transfected three plasmids into *E. coli* competent cells HB101 at the same time: pSWcuomxdCas9-\omega plasmid [13] carrying mxdCas9 protein and transcription activator, Omega(\omega); pZJY41-sgRNA series plasmid carrying sgRNA with different spacer sequences; pKK232 series plasmid carrying internal promoter and reporter gene CAT. Related sequences and plasmid information were shown in Tables S5 and S6.

**Construction of CRISPRa-dCas9 system in *Myxococcus xanthus***

We transfected two plasmids into *Myxococcus xanthus*: pSWcuomxdCas9-\omega plasmid [13] carrying mxdCas9 protein and transcription activator Omega(\omega); pZJY41-sgRNA series plasmid carrying sgRNA. Related sequences and plasmid information are shown in Tables S5 and S6.
**Extraction and detection of epothilones**

ZE9 and mutants were grown overnight in 50 mL of CYE medium supplemented with Apra (30 μg/mL). The cultures were inoculated at a ratio of 2:100 into 50 mL of CMO medium containing 2 % of the XAD-16 resin. The resin was harvested with a strainer after 6 days and extracted with 3 mL of methanol by shaking at room temperature overnight. The supernatant was centrifuged for 10 min at 12,000 rpm and filtered with 0.22 μm filter to remove the impurities. 20 μl of the sample was injected into High Performance Liquid Chromatography (HPLC, SHIMADZU, Japan) and analyzed on a Shim-pack MRCODS RP C18 column (4.6 mm × 250 mm, 4.60 μm; Shimadzu, Japan) and monitored at 250 nm, with a mobile phase of 60% of methanol (HPLC grade) and 40% of H2O at a flow rate of 1.0 mL/min. The yield of epothilone was quantified from the peak area in the UV chromatogram, by reference against a calibration standard.

**Transcriptional analysis of epothilone genes with RT-qPCR**

We collected samples continuously from the fermentation culture at 48 h of incubation. Then, total RNA of samples was extracted using BIOZOL kits (Total RNA Extraction Regent, Bio Fast, China) and then transcribed reversely into cDNA with PrimeScript™ Reagent Kit with DNAase (Takara, Japan). The gapA gene (glyceraldehyde-3-phosphate dehydrogenase gene, MXAN_2815) was chosen as the reference gene for normalization. The transcriptional level of epothilone gene cluster was analyzed by RT-qPCR on LightCycler®480 (Switzerland) with SYBR® Premix Ex Taq™ GC Dye (Takara, Japan). All the primers used in RT-qPCR were listed in TableS5.

**Declarations**

**Author contributions**

YW, XJY and YZL designed researches; YW, XJY, SFY, YH and WFH performed researches; YW, XJY and YZL analyzed the data and wrote the paper.

**Acknowledgments**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

**Availability of data and materials**

All data generated or analyzed during this study are included in this published article and in its Additional files.

**Consent for publication**

Not applicable.

**Ethics approval and consent to participate**

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Figures

Figure 1

Internal promoters in epothilone operon. (a)Prediction of internal promoters in epothilone operon from So0157-2. The promoter sequence analysis of the epothilone operon was performed using two online promoter prediction softwares. The red arrow is the promoter site predicted by “Neural Network Promoter Prediction”. The green arrow is the promoter site predicted by BPROM (softberry.com). The threshold was set to 0.8. (b) RT-qPCR analysis of expression levels of the seven genes for epothilone operon in ZE9 at 48h of incubation. The expression of the epoA gene was set as 1, and the expressions of the other six genes were shown as the relative expressions to epoA gene. (c) The activity of the separate internal promoters in E. coli. The activity of the reporter gene chloramphenicol acetyltransferase (CAT) was detected by CAT ELISA kit, with the aphII promoter as a positive control, and the original plasmid pKK-232 as a negative control (no promoter upstream of the reporter gene CAT). (d) The activities of the separate internal promoters in Myxococcus xanthus. The positive control is the identified promoter PepoA upstream of the epothilone operon. The fluorescence intensity of the green reporter gene was detected at
485nm/528nm, and three different incubation times (24h, 36h, and 72h) were selected for detection. The error bars represent the standard deviation of three independent experiments.

**Figure 2**

CRISPRa of internal promoters in epothilone operon. (a) The principle of CRISPRa. mxdCas9: codon-optimized inactivated dCas9 protein (H840A, D10A. Activation factors: Transcriptional activation factor, the Omega subunit of RNA polymerase, can play a role in recruiting RNA polymerase. SgRNA: Single guide RNA sequence, a combination of the CRISPR associated RNA (crRNA) and the trans-activation crRNA (tracrRNA). (b) The activation abilities of the internal promoters in E. coli. The activity of the reporter gene CAT was detected by CAT ELISA kit. As the positive control, the aphII promoter was not activated. (c) RT-qPCR analysis of expression levels of the six genes in epothilones operon in different mutants at 48h of incubation. The expressions of the six genes of epothilone operon in M. xanthus ZE9 were each set as 1, and the expressions of the genes in mutant strains were shown as the relative expressions to that of their corresponding genes in ZE9. (d) The yield of epothilones A and B and their summation in different activated mutants by CRISPRa and the ancestral strain M. xanthus ZE9. The error bars represent the standard deviation of three independent experiments. For statistical analysis between the ancestral strain and mutant strains, the signals of ** and * mean p < 0.01 and p < 0.05, respectively.
Figure 3

Combinatorial CRISPRa of epothilone operon. (a) The yields of epothilone A and epothilone B in wild strain M. xanthus ZE9 and mutants activated by CRISPRa. (b) RT-qPCR analysis of expression levels of the seven epothilone genes in double-activated mutants at 48h of incubation. (c) RT-qPCR analysis of expression levels of the seven epothilone genes in triple-activated mutants at 48h of incubation. The expressions of the seven genes of epothilone operon in M. xanthus ZE9 were each set as 1, and the expressions of the genes in mutant strains were shown as the relative expressions to that of their corresponding genes in ZE9. The error bars represent the standard deviation of three independent experiments. For statistical analysis between the ancestral strain and mutant strains, the signals of ** and * mean p < 0.01 and p < 0.05, respectively.
Figure 4

CRISPRa of different epothilone producing strains. (a) The yield of epothilones in wild strains and mutants activated by CRISPRa. (b) RT-qPCR analysis of expression levels of the seven epothilone genes in ZE9, ZE5, ZE10 and ZE14 at 48h of incubation. The expressions of the seven genes of epothilone operon in M. xanthus ZE9 were each set as 1, and the expressions of the genes of other strains were shown as the relative expressions to that of their corresponding genes in ZE9. (c) RT-qPCR analysis of expression levels of the seven epothilone genes in activated mutants at 48h of incubation. The expressions of the seven genes of epothilone operon in wild strains were each set as 1, and the expressions of the genes in mutant strains were shown as the relative expressions to that of their corresponding genes in their wild strains, respectively. The error bars represent the standard deviation of three independent experiments. For statistical analysis between the ancestral strain and mutant strains, the signals of ** and * mean p < 0.01 and p < 0.05, respectively.

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