Redirection of acyl donor metabolic flux for lipopeptide A40926B0 biosynthesis

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Summary
The metabolic flux of fatty acyl-CoAs determines lipopeptide biosynthesis efficiency, because acyl donor competition often occurs from polyketide biosynthesis and homologous pathways. We used A40926B0 as a model to investigate this mechanism. The lipopeptide A40926B0 with a fatty acyl group is the active precursor of dalbavancin, which is considered as a new lipoglycopeptide antibiotic. The biosynthetic pathway of fatty acyl-CoAs in the A40926B0 producer Nonomuraea gerenzanensis L70 was efficiently engineered using endogenous replicon CRISPR (erCRISPR). A polyketide pathway and straight-chain fatty acid biosynthesis were identified as major competitors in the malonyl-CoA pool. Therefore, we modified both pathways to concentrate acyl donors for the production of the desired compound. Combined with multiple engineering approaches, including blockage of an acetylation side reaction, overexpression of acetyl-CoA carboxylase, duplication of the dbv gene cluster and optimization of the fermentation parameters, the final strain produced 702.4 mg l⁻¹ of A40926B0, a 2.66-fold increase, and the ratio was increased from 36.2% to 81.5%. Additionally, an efficient erCRISPR-Cas9 editing system based on an endogenous replicon was specifically developed for L70, which increased conjugation efficiency by 660% and gene-editing efficiency was up to 90%. Our strategy of redirecting acyl donor metabolic flux can be widely adopted for the metabolic engineering of lipopeptide biosynthesis.

Introduction
Actinomycetes have proven to contribute various clinically applicable antibiotics. However, in the process of switching from primary to secondary metabolism, competition often occurs because different pathways may share the available precursors and energy in the host (Wan et al., 2015; Lu et al., 2016). Therefore, the rational redistribution of key precursors plays an important role in the efficient biosynthesis of the target product.

Lipopeptides are an important class of non-ribosomal peptide family compounds with a lipophilic acyl chain attached to the peptide core (Strieker and Dr, 2010). The fatty acyl side-chain in lipopeptides assists in dimer formation, membrane anchoring and stabilizing interactions with lipid II. These events enhance target site binding, prolong the half-life, and improve biological activity against gram-positive organisms (Zhanel et al., 2010). However, congeners differing in the fatty acyl side-chain have different efficacies, which hampers the industrialization of many microbial drugs (Zerilli et al., 2010; Alt et al., 2019).

Fatty acid biosynthesis is an essential metabolic pathway mediated by fatty acid synthase (FAS). This biosynthesis shares similar building blocks and catalysis mechanisms with polyketide biosynthesis in microorganisms (Hopwood and Sherman, 1990). In addition, a series of acetyl coenzyme A (acyl-CoA) molecules are considered to be the common precursor in both systems. Short-chain acyl-CoA precursors, such as acetyl-CoA and isobutyryl-CoA, are typically used as a starter unit in fatty acid biosynthesis, while malonyl-CoA functions as an extender unit for a growing acyl chain (Galina et al., 2002; Hertweck, 2009). Many studies have focused on increasing the supply flux of acyl-CoAs into the fatty acid pathway to stimulate the production of polyketides (Li et al., 2020; Ng et al., 2021). However, few studies have addressed redirecting the...
flux of acyl donors for lipopeptide biosynthesis (Shuai et al., 2018, Wu et al., 2018).

Dalbavancin is a new generation antibiotic for the treatment of severe infections, including those caused by vancomycin-resistant bacteria (Boucher et al., 2014). Lipopeptide A40926, the precursor of dalbavancin, is a mixture of compounds differing in the fatty acyl side-chain. According to the approval requirements for dalbavancin of the Food and Drug Administration, component B0 with an isododecanol side-chain accounts for >75% of the whole complex (Cavaleri et al., 2005). However, due to the broad substrate specificity of the acyltransferase Dbv8, a variety of A40926B0 analogues are produced during fermentation. Previous studies showed that supplementation with L-valine could increase the production of component B0 by 1-fold from approximately 4 to approximately 8 mg l⁻¹ (Jovetic et al., 2008). However, improving the biosynthetic efficiency of A40926B0 remains a key problem in industrial production.

Rare actinomycetes are generally not very accessible for genetic manipulation. This is a serious obstacle for further improvement of industrial strains. In Nonomuraea gerenzanensis L70, several genetic tools have been used for gene-editing. Among them, PCR targeting is time-consuming and incapable of achieving traceless mutagenesis (Yi et al., 2010). Cre/LoxP recombination is even more tedious as two rounds of conjugation are necessary (Xi et al., 2016). Homologous recombination (HR) with low recombination efficiency results in difficult conjugation transfer and necessary double crossover screening. In addition, the most popular gene-editing tool, CRISPR-Cas9, seemed to be useless in strain L70, since no transformants could be observed through conjugation. Therefore, efficient genetic tools are urgently required for this industrial strain.

In this study, we redirected acyl donor metabolic flux towards A40926B0 biosynthesis based on the developed endogenous replicon CRISPR (erCRISPR) system (Fig. 1). Firstly, a type II polyketide pathway was knocked out to channel the malonyl-CoA flux for lipopeptide biosynthesis. Second, the production of malonyl-CoA from acetyl-CoA was enhanced by overexpression of acetyl-CoA carboxylase (ACC). Thirdly, the branched-chain α-keto acid dehydrogenase (BCDH) complex encoded by bkdA2B2C2 was also overexpressed to further compete for malonyl-CoA between branched-chain fatty acid (BCFA) and straight-chain fatty acid (SCFA) biosynthesis. After eliminating the multilevel competition for malonyl-CoA, we introduced two extra copies of the dbv gene cluster. Finally, we successfully constructed a high-producing strain for A40926B0 named N. gerenzanensis IPB-9.

Results

Mining of the endogenous replicon in an indigenous plasmid-deficient strain

The industrial strain N. gerenzanensis L70 was obtained from Nonomuraea sp. ATCC 39727 by UV-induced mutation. Fermentation analysis showed that the yield of A40926B0 in L70 was 49.0% higher than that in the original strain (Fig. S1). Therefore, L70 was chosen for study to produce a higher yield of A40926B0. In our previous research, traditional genetic tools were shown to work with low efficiency (Fig. S2). After unsuccessfully testing several wide-host-range replicons, we focused on the mining of a suitable replicon in this industrial strain.

Two independent and stable indigenous plasmids (P1 and P2) were detected with sizes of 48321 bp and 7713 bp respectively. Genome annotation revealed 12 putative genes in P2 and 55 in P1 (Fig. 2A and Tables S1, S2). Among the 12 annotated genes in P2, orf0012 encodes the partition A protein (ParA). ParA has ATPase activity that provides energy for plasmid partitioning (Ebersbach and Gerdes, 2005). We then investigated the endogenous replicon in the P2 deletion strain. Many methods have been developed for plasmid elimination, including high temperature, the addition of chemical reagents and the use of transposons. The deletion efficiency usually appears to be extremely low, and injuries may occur in the host (Imre et al., 2006; Leavitt et al., 2010; Zaman et al., 2011). Therefore, a self-targeting CRISPR-Cas9 system was designed, in which the key gene parA was used as the target for disruption (Fig. 2B).

We firstly engineered the CRISPR-Cas9 system based on the suicide plasmid pSET153. The parA gene was ligated to the vector, ensuring its survival only through integration into P2 via HR. The single guide RNA (sgRNA) expression cassette targeting parA and codon-optimized Cas9 from Streptococcus pyogenes was added to generate pSC01. The expression of Cas9 was controlled by a double-induced system comprising the thiostrepton-inducible promoter tipAp and theophylline-inducible riboswitch, which synergistically reduced Cas9 activity at the transcription and translation levels, respectively, during conjugation (Wang et al., 2019). The constructed plasmid was then transformed into L70. Very few transformants (1.5 colonies per plate) integrated into P2 naturally. Induction was later performed in which the Cas9 protein cut the DNA, leading to two cleavages, as shown in Fig. 2B. Replica plating revealed that 60% of 30 single colonies were sensitive to apramycin (Fig. S3). This involved the loss of P2 as subsequently confirmed by diagnostic PCR (Fig. 2C). To our knowledge, this is the first study to use the self-targeting CRISPR-Cas9 strategy for indigenous plasmid elimination, which is
highly efficient and host-friendly. No significant difference was observed between the P2 deletion strain IPB-1 and L70 in biomass and A40926B0 production, indicating that P2 might have no critical effect (Fig. 2D).

Fig. 1. Schematic diagram of redirection of acyl donor metabolic flux to A40926B0 biosynthesis based on the erCRISPR system in Nonomuraea gerenzanensis L70. The engineering strategies include blockage of the competing pathway (shown in blue), overexpression of critical enzyme (shown in green) and enhancement of specific precursor (shown in red). Abbreviations are: ACC, acetyl-CoA carboxylase; ACP, acyl carrier protein; BCDH complex, branched-chain α-keto acid dehydrogenase complex; CLF, chain length factor; KS, ketosynthase; LipAB, lipoyl synthase & lipoyl transferase; NRPs, non-ribosomal peptides; PKs, polyketides.

Fig. 2. Mining of the endogenous replicon NGR70 in Nonomuraea gerenzanensis. A. Plasmid maps of P1 and P2 from N. gerenzanensis L70. Circle 1 displays the predicted ORFs. Circle 2 displays the GC skew (±). B. Overview of the self-targeting CRISPR-Cas9 strategy for P2 deletion in N. gerenzanensis L70. C. Identification of ΔP2 mutant by diagnostic PCR. Lane 1 and 4 show PCR products amplified by genome-F/genome-R primer pairs. Lane 2 and 5 show PCR products amplified by P1-F/P1-R primer pairs. Lane 3 and 6 show PCR products amplified by P2-F/P2-R primer pairs. D. Effect of P2 deletion on cell growth and A40926B0 production. Dry weight of mycelia was measured in the wild-type strain (L70) and the P2 deletion strain (IPB-1) at different incubation times as shown. A40926B0 production of L70 and IPB-1 at different incubation times as shown. Standard deviation (SD) bars are shown. E. Identification of the replication region of P2. A linear map of P2 is shown at the top. The bars below the map representing P2 regions covered in the respective plasmids.
pSR02, pSR03 and pSR04, whereas no transformants grew out using the other vectors (Fig. 2E). The essential region for P2 self-replication consisted of a 5788 bp fragment (named NGR70) with eight open reading frames (ORFs). Sequence alignment indicated that NGR70 is a new replicon in actinomycetes (Fig. S4). Furthermore, NGR70 is the first replicon available in *N. gerenzanensis*, with the potential for development into a powerful genetic tool.

**Construction of a sequential gene targeting erCRISPR-Cas9 system**

To broaden the application of the self-targeting CRISPR-Cas9 system, we combined it with NGR70 for genetic manipulation. The *tipAp* promoter and riboswitch were introduced into vector pSR04 to control the expression of Cas9. Two sgRNA expression cassettes were controlled by two independent inducible promoters, *PnitA* and Potr*, which respectively targeted the specific gene and the *parA* gene of the vector (Herai et al., 2004; Wang et al., 2016). Finally, one pair of homologous flanking regions of the specific gene with 3.0 kb was introduced into the vector (Fig. 3B), which functioned as the template for repairing the double-strand break mediated by Cas9.

Based on the above steps, we constructed the all-in-one vector pSRK04 for the traceless deletion of the ketosynthase (KS) gene named vpkAB (Fig. 3A). Through conjugation based on erCRISPR-Cas9, approximately 24 positive transformants were obtained on each plate. The efficiency was 2.87-fold higher than that mediated by the suicide vector pSK03 with the same homologous arms (Fig. 3E). Then, we transferred three transformants to the YEME liquid culture for 16 h at 28°C. The sgRNA cassette and Cas9 protein were induced for 48 h by supplementing with 5 μM thioestrepton, 8 mM theophylline and 15 mM ε-caprolactam to delete the vpkAB gene. Gene-editing efficiency was evaluated using PCR analysis and sequencing (Fig. 3C and 3D). Nearly 90% (86/96) of the colonies were identified as ΔvpkAB mutants, which was nearly 20-fold higher than that of the traditional HR strategy (4.2%, 4/96) based on pSK03 (Fig. 3E).

The second round of induction was performed to eliminate the plasmid itself in these mutants. 5 μM thioestrepton, 8 mM theophylline and 6 μM oxytetracycline were added to switch on the transcription of the *parA*-sgRNA cassette along with the Cas9 protein when the mycelia were cultured in YEME for 16 h. After a 48-h incubation, the transformants were transferred to a YM1 solid plate for successive subculture. Diagnostic PCR revealed that all 30 transformants lost the plasmid because of the continuous cutting by Cas9 and the absence of a corresponding repair template (Fig. S5). The mode of traceless deletion ensures multiple usages of the erCRISPR-Cas9 system. Moreover, the time for gene-editing was greatly shortened from 52 d to 32 d, as the traditional editing method relying on HR requires multiple subcultures to obtain mutant colonies (Table 1).

**Blocking polyketide synthesis to redistribute the metabolic flux of malonyl-CoA**

*N. gerenzanensis* L70 is known for producing A40926B0, the precursor of dalbavancin. To improve the yield of A40926B0, we first constructed the IPB-2 strain by deleting the *dbv23* gene, which encodes a pathway-specific acetyltransferase responsible for acetylating the mannose moiety of A40926B0 (Sosio et al., 2010). Liquid chromatography-mass spectrometry (LC-MS) analysis showed that the A40926B0 yield was up to 251.6 mg l⁻¹ and eight components differing in the fatty acyl chain were detected in fermentation broths (Fig. 4A, B, S6 and S7). These moieties consisted of a straight-chain or branched-chain fatty acyl group with 11 to 14 carbon atoms. The A40926 complex composition has been closely related to the cell fatty acid pattern (Jovetic et al., 2008). Thus, we speculated that the composition of A40926 could be affected by changing the content of related fatty acids in the strain.

So far, the non-ribosomal peptide A40926, encoded by 37 ORFs, is the only secondary metabolite reported in *N. gerenzanensis* (Sosio et al., 2003). RNA-seq analysis also showed that many other gene clusters were in a cryptic state or low-level expression, except for a type II PKS-encoding cluster (cluster 23), designated here as the vpk-gene cluster (Table S4). The expression of the core synthase gene vpkAB in the vpk-gene cluster (Fig. 5A, 5B and Table S3) was detected at a relatively high level. Since malonyl-CoA is used as the common extender unit in the biosynthesis of type II polyketide and fatty acids, we sought to redistribute the metabolic flux of malonyl-CoA into A40926 biosynthesis by blocking the polyketide pathway (Fig. 4C).

The vector pSRK04 for in-frame deletion of the vpkAB gene was transformed into IPB-2, mediated by the developed erCRISPR-Cas9 system to generate IPB-3. Compared with the parent strain, the pigment production of IPB-3 was significantly reduced in shaker fermentation (Fig. 5C). High-performance liquid chromatography (HPLC) analysis also demonstrated that IPB-3 possessed cleaner metabolite profiles than IPB-2, indicating the elimination of the pigment compounds produced by the vpk-gene cluster (Fig. 5D). The metabolites related to primary metabolic pathways, including the tricarboxylic acid cycle, pentose phosphate pathway, Embden-Meyerhof-Parnas pathway, amino acid metabolism and
lipid metabolism (LPM), were identified and compared. The levels of metabolites in LPM showed a remarkable increase in IPB-3, while no significant change was detected in other pathways (Fig. 5E and S8). The yield of the A40926 components was subsequently tested by analysing the peak area from the HPLC chromatogram. No significant difference was observed in most components, including A40926B0 (3). In contrast, A40926B1 (4), with a straight-chain decanoyl group, was significantly increased in IPB-3 by 92.2% (Fig. 5F). These data indicated that deletion of the vpkAB gene transferred the malonyl-CoA from the PKS pathway to the FAS pathway associated with lipopeptide synthesis, favouring the biosynthesis of SCFAs instead of BCFAs. In addition, the intracellular ATP and NADH/NAD⁺ levels were increased by nearly 2-fold in IPB-3. This might provide more cellular energy and reducing power for the improved productivity of A40926B0 in the future studies (Fig. 5G).

Enhancing metabolic flux in target fatty acid pathway

Overexpression of ACC improves the supply of malonyl-CoA. Since blockage of the vpk-gene cluster biosynthetic pathway resulted in a redistribution of malonyl-CoA flux into fatty acid biosynthesis, the next
and IPB-3, while no significant changes were observed in other components (Fig. S9).

Identifying the pivotal bkd cluster to enhance the precursors in BCFA synthesis. In the biosynthetic pathway of SCFAs, the main initial unit of acetyl-CoA is relatively abundant, as it plays an essential role in primary metabolism. To further enhance the production of A40926B0 with a branched-chain isododecanoyl group, the key bottleneck was to improve the biosynthesis of BCFAs, which may compete for malonyl-CoA with SCFAs. In contrast, isobutyryl-CoA and isovaleryl-CoA, which function as specific precursors of BCFAs, were reported to be only derived from branched-chain amino acid (BCAA) degradation.

In BCFA catabolism, a bkd gene cluster containing bkdA/B/C genes encodes the subunits E1,a, E1,b and E2 of the branched-chain α-keto acid dehydrogenase complex (BCDH complex). The BCDH complex catalyses the oxidative decarboxylation of α-keto acids derived from BCAAs to produce the corresponding acyl-CoA derivatives, which then act as precursors for BCFA biosynthesis (Stirrett et al., 2009).

As reported in other streptomycetes (Stirrett et al., 2009; Shuai et al., 2018), two alleles of bkd gene clusters were located in L70 through BLAST alignment (Fig. 6A and S10). To investigate their functions in this industrial strain, the vectors pSRB01 and pSRB02, which include sgRNA targeting the promoter of the corresponding bkd cluster along with the dCas9 protein, were transformed into IPB-4 (Fig. S11). Transcriptional analysis revealed a significant decrease in the expression levels of

![Redirect flux for A40926B0 biosynthesis](1857)

### Table 1. Efficiency comparison between erCRISPR-Cas9 system and the traditional genetic tool based on HR used in Nonomuraea gerenzanensis.

|                          | erCRISPR-Cas9 | HR by suicide vector |
|--------------------------|--------------|----------------------|
|                          | Plasmid construction 9 days | Plasmid construction 7 days |
| Conjugation              | 9 days       | Conjugation 9 days    |
| Subculture               | 4 days       | Subculture 4 days    |
| First induction gene-editing | 5 days     | Double 4 days        |
|                          |              | crossover screening 8 = 32 days |
| Second induction plasmid elimination | 5 days | Total 32 days |
| Conjugation efficiencya  | 24           | Conjugation efficiencya 6.2 |
| Editing efficiency       | 90.9%        | Editing efficiency 4.2% |

a. Conjugation efficiency was calculated as the number of positive transformants obtained on each plate.

![Fig. 4. A40926 complex and its related fatty acyl side-chain.](1857)

A. HPLC analysis of A40926 components in fermentation extracts of Nonomuraea gerenzanensis L70. The numbered peaks represent eight components of A40926 that were confirmed by LC-MS data and ultraviolet (UV) absorption as shown in Fig. S6.

B. Chemical structures of A40926 complex. The acyl group was shown as FA (fatty acyl). Structures of FA moieties in the A40926 complex are shown with the corresponding marked numbers in HPLC analysis.

C. Partial metabolic pathway in N. gerenzanensis L70. Solid arrow denotes the direct reaction and the dotted arrow denotes the indirect reaction.

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and bkdA2B2C2 in the respective resulting transformants IPB-5 and IPB-6 (Fig. S12). The production levels were examined. The yield of A40926B0 was reduced by 7.7% in IPB-5 and 48.4% in IPB-6 when compared with IPB-4. As a similar tendency was observed in the corresponding ratio, we surmised that bkdA2B2C2 played a critical role in BCFA biosynthesis based on the erCRISPR-dCas9 system (Fig. 6B). On the other hand, lipoylation involving LipAB was reported to be insufficient for BCFA production (Bentley et al., 2016). bkdA2B2C2 and lipAB were overexpressed in IPB-4 to obtain IPB-7. Fermentation analysis showed that the A40926B0 titre in IPB-7 was improved to 342.2 mg l⁻¹, which corresponds to an increase of 36.1%, compared with that in IPB-4. Remarkably, the production of A40926 components with a straight-chain fatty acyl group in the side-chain was reduced to various degrees, among which A40926B1 showed the biggest decline of 43.1% in IPB-7 when compared with IPB-4 (Fig. 6C).

To further evaluate the effect of our rational engineering, we examined the composition of cell fatty acids based on GC-MS analysis of their methyl esters (Fig. 6D and S13). As shown in Fig. 6E and F, deletion of the vpkAB gene resulted in an increase in SCFAs, including n-C16:0 and n-C18:0, in IPB-3. The combinatorial strategy of strengthening the BCFA metabolic flux in IPB-7 enhanced both the amount and ratio of i-C16:0, which may act as the donor of the isododencanoyl moiety in A40926B0. These results agreed with the changes in the A40926B0 ratio and production, indicating that the metabolic flux of malonyl-CoA was redirected into the biosynthesis of the target product A40926B0 after precursor redirection. In addition, transcripts of genes

Fig. 5. Blockage of the vpk-gene cluster biosynthetic pathway results in redistribution of malonyl-CoA flux.
A. Analysis of relative expression levels of putative gene clusters in Nonomuraea gerenzanensis L70 based on RNA sequencing data. The housekeeping gene hrdB was chosen as the reference gene.
B. Genetic organization of the vpk-gene cluster in N. gerenzanensis L70.
C. Phenotypic appearance of culture broths of wild-type and mutant strains after cultivation for 168 h in YS medium.
D. Comparison of metabolite profiles analysis based on isosorbance plot between IPB-2 and IPB-3.
E. Relative levels of the identified metabolites in the different metabolic pathways in IPB-2 and IPB-3 from cultures at 36 h in YS medium. Significant statistical differences are shown (***, P < 0.001; **, P < 0.01; *, P < 0.05; NS, not significant). SD bars are also shown.
F. Production analysis of each component corresponding to the number of A40926 complex (Fig. 4B) by HPLC in IPB-3 compared with that in IPB-2.
G. Relative levels of intracellular NADH/NAD⁺ ratio and ATP concentration in IPB-2 and IPB-3 from cultures at 36 h in YS medium. Abbreviations: AAM, amino acid metabolism; AccA, acetyl-CoA carboxyl transferase subunit; AccB, acetyl-CoA carboxylase biotin carboxyl carrier protein; AccC, acetyl-CoA carboxylase biotin carboxylase subunit; CYC, cyclase; EMP, Embden-Meyerhof-Parnas pathway; KSα, ketosynthase alpha subunit; KSβ, ketosynthase beta subunit; LPM, lipid metabolism; PhoP, response regulator; PhoR, sensor kinase; PPP, pentose phosphate pathway; TCA, tricarboxylic acid cycle.

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including fabD, fabF, fabG, fabl, fabK and fabH, which are essential for fatty acid biosynthesis, were upregulated in IPB-7 (Fig. 6G). Similar trends were observed in the metabolome analysis, which revealed that metabolites associated with fatty acid metabolism showed relatively large differences between IPB-7 and IPB-2. Furthermore, the abundance of metabolites in pyruvate metabolism also exhibited otherness, possibly because pyruvate, as a central metabolic hub, is susceptible to the engineering of other metabolic pathways (Fig. 6H). In general, through our redirection research strategy, fatty acid metabolism, especially the branched-chain one, tended to be more active, which could provide sufficient precursor supply for the efficient biosynthesis of A40926B0.

Further improvement in the composition of A40926B0 by supplying sodium isobutyrate. Given that enhancement
of specific precursors could reduce homologues by competing for malonyl-CoA donors, we attempted to further improve the precursor supply through exogenous addition. The effect of sodium isobutyrate on A40926B0 production was subsequently examined by culturing IPB-7 in YS medium (Fig. S14). 10 mM of sodium isobutyrate was found to be the optimal concentration to achieve the highest yield of 448.2 mg l⁻¹. By supplying sodium isobutyrate to IPB-7, A40926B0 was increased remarkably by 31.0%, while A40926B1 was decreased by 52.1%, compared with that in IPB-7 (Fig. S15). Moreover, calculation of the relative utilization levels of malonyl-CoA revealed a 177.5% increase for A40926B0 together with a 63.2% decrease for other analogues (Fig. 6I). The interaction between Dbv8, the key gene for generating congeners, and different fatty acyl-CoA molecules was analysed by affinity calculation through molecular docking (Lyu et al., 2014). Isododencanoyl-CoA exhibited the strongest binding affinity towards the Dbv8 protein, which explains why sodium isobutyrate supplementation increased A40926B0, rather than A40926D0 (7), although they shared the same precursor isobutyryl-CoA (Fig. S16).

**Construction of a higher A40926B0-producing strain by cluster duplication and fermentation optimization**

Duplication of gene clusters has been reported to effectively improve the titre of target compounds (Li et al., 2017). Here, the vector pMSBBAC1 containing a complete dbv gene cluster was screened from the bacterial artificial chromosome library and introduced into IPB-3 along with pSOA02 containing the overexpression cassettes of accABC, bkdA2B2C2 and lipAB. A further increase of 58.5% was observed in A40926B0 production (Fig. 7B). Based on these results, we attempted to increase the copy number of pMSBBAC1. An artificial ΦC31 attB site was introduced into the chromosome of IPB-3 by replacing the synthase gene of a putative terpene gene cluster (cluster 32) via erCRISPR-Cas9 (Fig. 7A). The vectors pMSBBAC1 and pSOA02 were successively transformed into the parental strain to obtain IPB-9, which contains two extra copies of the dbv gene cluster (Fig. S17). Production analysis revealed that IPB-9 produced A40926B0 at a titre of 507.3 mg l⁻¹, which was nearly 48.2% and 164.2% higher than that of IPB-6 and L70 respectively (Fig. 7B). Scale-up fermentation was further performed to verify the production of A40926B0 by IPB-9 in a 2-L flask with 350 ml working volume in YS medium supplemented with 10 mM sodium isobutyrate. After culturing for 168 h, the titre of A40926B0 reached 550.8 mg l⁻¹, with the productivity of 0.43 mg g⁻¹ dry cell weight h⁻¹ (Fig. 7B and C). The time course profile showed that production of A40926B0 increased continuously from 48 h to 96 h and reached a plateau after culturing for 120 h and 48 h earlier than that of the wild-type strain (Fig. 7D). Therefore, sodium isobutyrate was fed at a rate of 10 mM for 24 h

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**Fig. 7.** Further improvement of A40926B0 production. 
A. Diagram of the introduction of dbv gene clusters in the parental strain. 
B. A40926B0 production in the engineering strains. SD bars are shown. 
C. Heat map of the peak area of the A40926 components in different strains. The numbers of components correspond to those in Fig. 4B. 
D. Time-course profile of A40926B0 fermentation. IPB-9 strain in YS medium with continuous feeding of 10 mM sodium isobutyrate (i), IPB-9 strain in YS medium with continuous feeding of 10 mM sodium isobutyrate (ii), L70 strain in YS medium with continuous feeding of 10 mM sodium isobutyrate (iii). Fermentation broths were sampled for analysis every 24 h. SD bars are shown.
Discussion

Malonyl-CoA is an important central metabolite that serves as the elongation unit for the biosynthesis of polyketides and the acyl chain of lipopeptides. Commonly, intracellular malonyl-CoA availability is regarded as a crucial bottleneck limiting target product formation (Milke and Marienhagen, 2020). Several strategies have been adopted to boost polyketide production by improving the supply of malonyl-CoA, such as enhancement of ACC activity and inhibition of fatty acid or by-product polyketide synthesis (Santos et al., 2011; Zabala et al., 2013; Kallscheuer et al., 2016). However, few studies have focused on the flux of malonyl-CoA for lipopeptide biosynthesis. One typical example is the production of surfactin was improved with a higher level of malonyl-ACP through overexpression of ACC and FabD (Wu et al., 2018). Fatty acids play important roles in growth maintenance and secondary metabolite production, which makes the precise control of malonyl-CoA metabolic flux challenging. In this study, the key synthase enzyme of a major polyketide by-product was deleted to channel the malonyl-CoA flux to the lipopeptide A40926 biosynthesis, which represents the first example of channelling the key precursor malonyl-CoA from polyketide biosynthesis into lipopeptide biosynthesis.

In A40926 biosynthesis, a series of congeners mainly derived from the degradation of cellular SCFAs were produced. However, the biosynthesis of SCFAs seems difficult to block because the end product palmitic acid is essential for the growth and is derived from acetyl-CoA. However, the initial substrate of BCFA biosynthesis is typically controlled by the bkd cluster. Considering that A40926B0 has a branched-chain isododecanoyl group, it seems feasible to compete for malonyl-CoA from the SCFA pathway by regulating bkd cluster expression. bkdA2B2C2 overexpression significantly improved the A40926B0 titre, together with a reduction in by-product production. The yield of A40926B0 was further increased by the addition of sodium isobutyrate. As the feedstock precursor, sodium isobutyrate could be obtained through an equal-molar chemical reaction of isobutyric acid and sodium hydroxide, which enabled the efficient utilization of malonyl-CoA from the fatty acyl pathway in A40926B0 production. A remarkable increase in the A40926B0 titre was achieved based on our strategy, suggesting that insufficient supply of acyl precursors is a key obstacle for A40926B0 production. Besides, the replacement of promoter in the above overexpression cassettes and the increase of dbv gene cluster copy number are expected to enhance the A40926B0 yield further. Overall, our work provides an efficient strategy for reducing the metabolic flux in branching pathways by competing for common precursors, which can be applied to improve the amount of target product in the mixture through rational metabolic engineering.

An efficient genetic manipulation system is a prerequisite for remodelling industrial microorganisms. Efforts to enhance conjugation efficiency have included improving the DNA uptake by the recipient strain via protoplast preparation (Marcone et al., 2010), sonication (Sofia et al., 2010), electroporation (Cho et al., 2017) and optimization of the culture conditions, including concentrations of Mg²⁺ and Ca²⁺ (Wang and Jin, 2014), temperature and medium (Netzker et al., 2016). However, these explorations have been time-consuming and labour-intensive. In N. gerenzanensis L70, we optimized the conjugation efficiency from all aspects and found that many transformants could be obtained by integrative vector-mediated conjugation. Nevertheless, these attempts seem to be useless for episomal vectors. We hypothesize that the incompatibility of replicons might be the key factors affecting conjugation efficiency. Replicons in endogenous plasmids have been developed in Propionibacterium, Pseudomonas alcaligenes and other microorganisms and have proven to be efficient for conjugation (Kwong et al., 2000, Kiatpapan and Murooka, 2002). However, no studies have reported the coupling of endogenous replicons with the CRISPR system. Here, by identifying the endogenous replicon NGR70, we developed an inducible erCRISPR-Cas9 system for traceless mutagenesis. The presence of NGR70 ensured the replication of the plasmid, and increased Cas9 and sgRNA expression levels in the host. The efficiency of conjugation and gene-editing was improved by 6.6-fold and 22-fold respectively. Thus, we conclude that replicon adaptability may be the key for genetic manipulation of many industrial strains. Interestingly, NGR70 had no homology with known replicons, and it may further be developed into an element for universal vectors with potential applications in other actinomycetes. In brief, our work provides a novel genetic strategy based on the examination of replicon compatibility with the host, which will be universally applicable to industrial strains.

Overall, a combinatorial approach was used to concentrate the metabolic flux of acyl donors into the biosynthetic route of the fatty acyl side-chain in A40926B0 based on the developed erCRISPR system. The fermentation yield of A40926B0 is up to 702.4 mg l⁻¹ in flask
increased by 2.66-fold. A40926B0 accounted for 81.5% compared to 36.2% in the original strain. Our study deciphered the metabolic flow distribution of acyl donors during drug biosynthesis in actinomycetes and provided a widely applicable strategy for the efficient biosynthesis of lipopeptides for research and industrialization.

**Experimental procedures**

**Bacterial strains and growth conditions**

The plasmids and strains used in this study are listed in Table S5. *Escherichia coli* DH5α was used as the host for plasmid construction. *E. coli* ET12567/pUZ8002 was used as a donor for intergeneric conjugation. *E. coli* was grown at 37°C in Luria–Bertani (LB) medium (1% tryptone, 0.5% yeast extract, 1% NaCl). *N. gerenzanensis* L70 and its mutants were cultured on YMG solid medium (2% soya malt extract, 0.2% yeast extract) at 28°C for growth. For fermentation, a 1 cm² agar piece was incubated in a 100 ml flask with 20 ml seed medium YEME (4% glucose, 0.5% tryptone, 0.3% malt extract, 0.3% yeast extract) at 28°C with shaking at 220 rpm for 7 days. MS solid medium (2% soya flour, 2% mannitol, 2% agar) supplemented with 10 mM MgCl₂ was used for interspecies conjugation between *E. coli* ET12567/pUZ8002 and *N. gerenzanensis* mycelia. If needed, antibiotics were added as follows: 50 μg ml⁻¹ apramycin, 25 μg ml⁻¹ chloramphenicol, 50 μg ml⁻¹ kanamycin and 25 μg ml⁻¹ nalidixic acid.

**Design of sgRNAs from N. gerenzanensis L70 genome and plasmid sequence**

The sgRNAscas9 software package was applied to rapidly search for CRISPR target sites (Xie et al., 2014). All 20 nucleotide-NGG sgRNA sequences of the target genes in the *N. gerenzanensis* L70 genome or endogenous plasmid were selected as candidate binding sites. Programs were used to search for CRISPR target sites with user-defined parameters, and the potential off-target cleavage sites of CRISPR-Cas9 were analysed. By comparing the total number of the potential off-target sites for each CRISPR target sequence, we selected precise sgRNA candidates with high specificity. All the designed sgRNAs are listed in Table S6.

**Plasmid construction**

The primers used in this study are listed in Table S7. The primers were synthesized using GENEray (Shanghai, China). PCR amplification was performed using Phanta Max Super-Fidelity DNA Polymerase (Vazyme, Nanjing, China) and identification was performed using 2×HiFle™ PCR Master Mix (Yeasen, Shanghai, China). A HiPure Gel Pure DNA Mini Kit (Magen, Guangzhou, China) was used to purify the PCR fragments, and the Plasmid Mini-prep Kit (GENEray) was used to isolate plasmid DNA. All enzymes used for DNA manipulation were purchased from Thermo Fisher Scientific (Waltham, MA, USA). Ligation reactions were performed by seamless cloning using the ClonExpress II One Step Cloning Kit (Vazyme, Nanjing, China). The primers parA-F1/R1, cas9-F1/R1, parA-sgRNA-F1/R1 and parA-sgRNA-F2/R2 were used to amplify the parA gene, tipAp fragment and riboswitch associated with the codon-optimized Cas9, ermE* promoter and the gRNA scaffold from vector pKC1139-TRMA, followed by ligation into the pSET153 plasmid, yielding pSC01 (Wang et al., 2019). To construct pSR01–pSR07, seven fragments were amplified from genomic DNA and generated R1–R7, which covered the position 1–7713 bp, 1926–1925 bp, 3851–3850 bp, 3851–1925 bp, 5776–1925 bp, 4813–963 bp respectively. The R1–R7 fragments were ligated into the EcoRⅠ and HindⅢ sites of pSET153 to obtain the vectors. To construct pSRK04, the primers cas9-F2/cas9-R2, vpkAB-sgRNA-F/R, parA-sgRNA-F2/R3, Potr-F/R and Pnlt-F/R were used to amplify the tipAp-ribo-Cas9, vpkAB-sgRNA, parA-sgRNA, Potr* and PnltA promoters from vector pKC1139-TRMA, plW01 and plL99. (Sun et al., 2012; Wang et al., 2016). Finally, the 3.0 kb homologous arms flanking the vpkAB gene were amplified from genomic DNA and ligated into the EcoRⅠ site of the vector. To construct pSR01/pSR02, the dCas9 fragment was amplified from pCRISPR-Cas9 using the primers dCas9-F/R and fused with the ermE* promoter. The obtained fragment was ligated into pSR04 using EcoRⅠ. The gRNA spacer sequences of the bkdA1B1C1 and bkdA2B2C2 promoters were designed using the primers bkd1-sgRNA-F2 and bkd2-sgRNA-F2. The corresponding sgRNA cassette was ligated into the EcoRⅠ site of the vector. To construct pIB02, the bkdA2B2C2 and tipAp genes were amplified from genomic DNA using the primers ermEp-bkd2-F/R, ermEp-lipA-F2/R2 and ermEp-lipB-F/R respectively. The fused fragment was ligated into the Ndel and EcoRⅠ sites of vector plJ8661. pMSBBBC1 was constructed by Eight Star Biotech (Wuhan, China), and the combinatorial expression fragment amplified by ermEp-bkd2-F2/R2 was ligated into pSOA01 to generate pSOA02.

**Transformation of N. gerenzanensis L70**

For the preparation of the donor *E. coli* cells, all recombinant plasmids were transferred into *E. coli* ET12567/
pUZ8002. An overnight suspension culture (200 μl) was inoculated in 20 ml of LB medium containing 50 μg ml⁻¹ apramycin, 25 μg ml⁻¹ chloramphenicol and 50 μg ml⁻¹ kanamycin. After 3 – 4 h incubation at 37°C and 250 rpm, cultured E. coli with an optical density of 0.4 – 0.6 at 600 nm (OD600) were collected by centrifugation at 4000 g for 5 min, washed three times with sterile LB medium, and resuspended in 500 μl of LB medium. To prepare N. gerenzanensis mycelia, a 1 cm² agar piece from plate colonies cultured for 6 – 10 days was inoculated into 30 ml of YEME medium and cultured at 28°C and 220 rpm for 48 h. Then 1 ml of the mycelia was harvested by centrifugation at 4000 g for 5 min, washed three times with sterile LB medium, and resuspended in 500 μl of LB medium. Finally, about 10⁸ donor E. coli cells were mixed with about 10⁷ mycelium cells and spread on the plate with MS medium supplemented with 10 mM MgCl₂. After cocultivation at 28°C and 200 rpm, cultured E. coli cells were mixed with about 10⁷ mycelium cells and spread on the plate with MS medium supplemented with 10 mM MgCl₂. After cocultivation at 28°C and 220 rpm for 10 days and transferred into YMG plates with 0.4 g/l apramycin, 25 gm l⁻¹ chloramphenicol and 50 gm l⁻¹ kanamycin. After 3 h incubation at 37°C and a subsequent isocratic stage of 90% B for 10 min at a constant flow rate of 1 ml min⁻¹ was applied. For the analysis of total metabolites, a linear gradient from 10% to 90% (v/v) B over 30 min and a subsequent isocratic stage of 90% B for 10 min at a constant flow rate of 1 ml min⁻¹ was applied.

For the identification of A40926B0 and its derivatives in the sample, liquid chromatography-mass spectrometry (LC-MS) analysis was performed using the same parameters as those used for HPLC analysis. Electrospray ionisation (ESI)-MS was conducted on a Thermo Finnigan LCQ Deca XP MAX system, with the ESI source operating in positive ionization mode. The scan range was m/z 400–2000. Using an A40926B0 standard as a reference, the retention time of A40926A0, A40926A1, A40926B0, A40926B1, A40926C0, A40926C1, A40926D0 and A40926D1 was 11.092, 11.650, 14.510, 20.250, 21.325 and 21.946 min respectively (Fig. 4A, B and S5).

For the identification of metabolites in the different metabolic pathways, 1 ml extract solution (methanol: acetonitrile: water, 2: 2: 1) containing isotopically labelled internal standard mixture was added to the sample. The samples were sonicated for 20 min and centrifuged at 10 000 for 20 min at 4°C. The resulting supernatant was transferred for analysis. LC-tandem MS (MS/MS) analysis were performed using a Vanquish ultra-HPLC system (Vanquish, Thermo Fisher Scientific) with an ultra-performance LC (UPLC) BEH Amide column (2.1 mm × 100 mm, 1.7 μm) coupled to a Q Exactive HFX mass spectrometer (Orbitrap MS, Thermo Fisher Scientific). The mobile phase consisted of 25 mM ammonium acetate and 25 ammonia hydroxide in water (pH = 9.75) (A) and acetonitrile (B). The QE HFX mass spectrometer was used to acquire MS/MS spectra in the information-dependent acquisition mode using Xcalibur acquisition software (Thermo Fisher Scientific). The ESI source was operated in both positive and negative ionization modes.

Metabolites analysis

After 7 days of cultivation in YS medium, the culture was mixed with methanol in equal proportions and sonicated for 1 h to disrupt the cells. The sample was centrifuged at 10 000 g for 10 min. The supernatant was filtered through a 0.22 μm filter (Millipore, Billerica, MA, USA) for analysis. HPLC analysis of A40926 was performed on an Agilent 1260 infinity system (Agilent, San Diego, CA, USA) equipped with a DAD detector at 220 nm. The column was an Agilent ZORBAX Eclipse Plus C18 (5 μm, 4.6 × 250 mm²). Trifluoroacetic acid (0.1% in water) was mobile phase A and acetonitrile containing 0.1% trifluoroacetic acid was mobile phase B. For the analysis of A40926 production and composition, the chromatographic condition performed a linear gradient from 30% to 50% (v/v) B over 30 min and a subsequent isocratic stage of 90% B for 10 min at a constant flow rate of 1 ml min⁻¹. For the analysis of total metabolites, a linear gradient from 10% to 90% (v/v) B over 30 min and a subsequent isocratic stage of 90% B for 10 min at a constant flow rate of 1 ml min⁻¹ was applied.

Quantitative RT-PCR

Mycelia of N. gerenzanensis L70 were washed twice with RNase-free TE buffer and total RNA extraction was performed using the EASYspin Plus bacteria RNA extract kit (Aidlab Biotech, Beijing, China) according to the manufacturer’s guidelines. The RNA was treated with RNase-free DNase I (TaKaRa Bio, Shiga, Japan) to eliminate residual genomic DNA. After PCR validation, the RNA samples were reverse transcribed into cDNA using the PrimeScript™ 1st Strand cDNA Synthesis Kit (TaKaRa Bio) following the manufacturer’s guidelines. Quantitative real-time PCR (qRT-PCR) was performed on a LightCycler 480 (Roche, Mannheim, Germany)
using TB Green™ Premix Ex Taq™ II (TaKaRa Bio) with
the primers listed in Table S7 according to the manufacturer’s guidelines. Fold changes in genes in different
samples were quantified by normalizing to the expres-
sion level of hrdB.

**RNA-seq analysis**

All the RNA samples were prepared as above from
mycelia cultured for 72 h. A total amount of 3 μg RNA
per sample was used as input material for transcriptome
sequencing. The preparations were sequenced on an
Illumina Novaseq platform (Illumina, San Diego, CA,
USA) and 150 bp paired-end reads were generated.
HTSeq v0.6.1 was used to count the reads numbers
mapped to each gene. Fragments per kilobase of tran-
script per million mapped reads (FPKM) were calculated
to estimate gene expression levels.

**Analysis of fatty acids**

Mycelia cultured for 72 h were suspended in 6 ml of
methanol containing 2% HCl and heated at 85°C for 2 h.
Then 1 ml of 14% boron trifluoride in methanol was
added to form fatty acid methyl esters (FAMEs) after
heating at 100°C for 15 min. The methanolsate was
evaporated to dryness and extracted with 1 ml n-
hexane, which was injected onto a 7890B/7000C GC-
mass spectrometer via HP-5MS column. Helium was used as the
carrier gas at a constant flow rate of 1 ml min⁻¹ and a split
ratio of 20:1. The temperature was maintained at 50°C
for 2 min, increased at a rate of 20°C min⁻¹ to 150°C,
10°C min⁻¹ to 200°C for 10 min, 2.5°C min⁻¹ to 215°C
for 8 min and 5°C min⁻¹ to 270°C for 5 min. The mass
spectrometer ion-source temperature was 250°C, and the scan range was 50–550 m/z.

**Docking analysis of Dbv8 with acyl-CoAs**

The crystal structure of acyltransferase in complex with
decanoyl-CoA (Protein Data Bank code 4MFZ) was used as
the template for docking of eight fatty acid-CoAs. The three-dimensional (3D) structure of the ligand was
downloaded from PubChem, with hydrogen atoms and electric
charges added. The energy of the ligand was optimized
using the MMFF94 field. The UCSF Chimera molecular
visualization program was used to prepare the ligands
and receptors. Decanoyl-CoA in the eutectic structure
was used as a reference ligand to define the combined
pocket by removing water molecules and other ions. A
docking grid box was then generated, and UCSF DOCK
6.9 was employed for docking. Each pose was extracted
with the lowest binding energy and analysed using PyMOL.

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**Conflict of interest**

The authors declare no competing interests.

**Author contributions**

T.Y. Xia and Y.Q. Li designed the study. T.Y. Xia con-
ducted the experiments and wrote the manuscript. T.Y.
Xia and X.A. Chen analysed the data. X.A. Chen, Y.Q.
Liu, D H. Scharf, Q.W. Zhao and Y.Q. Li revised the
manuscript. All authors read and approved the final
manuscript.

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Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

**Fig. S1.** Comparison of morphological phenotype and A40926B0 production in Nonomuraea sp. ATCC 39727 and *N. gerenzanensis* L70.

**Fig. S2.** Evaluation of the conjugation efficiency (A) and gene-editing efficiency (B) of plasmid tools based on the traditional genetic strategies.

**Fig. S3.** Growth status of the corresponding mutants on the MS plates.

**Fig. S4.** Nucleotide sequence alignment of NGR70 in NCBI database.

**Fig. S5.** Growth status of the corresponding mutants on the MS plates.

**Fig. S6.** MS and UV spectra of A40926 components.

**Fig. S7.** Production analysis of each component corresponding to the number of A40926 complex (Fig. 4B) by HPLC in IPB-2 compared with that in L70.

**Fig. S8.** Relative levels of the identified metabolites in the different metabolic pathways.

**Fig. S9.** Production analysis of each component corresponding to the number of A40926 complex (Fig. 4B) by HPLC in IPB-4 compared with that in IPB-3.

**Fig. S10.** Amino acid alignments of partial sequence of BCDH complex in *N. gerenzanensis* L70 and other actinomycetes.

**Fig. S11.** Plasmid map of pSRB01 based on the erCRISPR-dCas9 system.

**Fig. S12.** The relative expression of *bkdA1B1C1* and *bkdA2B2C2* in IPB-4 and IPB-5.

**Fig. S13.** MS spectra of FAMEs for *N. gerenzanensis* cell fatty acids.

**Fig. S14.** Comparison of A40926B0 production with supplementation of sodium isobutyrate at different concentrations.

**Fig. S15.** Production analysis of each component corresponding to the number of A40926 complex (Fig. 4B) by HPLC in IPB-7 supplemented with 10 mM sodium isobutyrate compared with that in IPB-7.

**Fig. S16.** Molecular docking of fatty acyl-CoAs with the acyltransferase Dbv8 and evaluation of Dbv8 substrate affinity for fatty acyl-CoAs.

**Fig. S17.** Validation of two extra copies of the *dbv* gene cluster.

**Fig. S18.** HPLC analysis of A40926 components in fermentation broths.

**Table S1.** Proposed functions of ORFs in P1.

**Table S2.** Proposed functions of ORFs in P2.

**Table S3.** Prediction of possible secondary metabolites of gene clusters in *Nonomuraea gerenzanensis* L70 and transcriptional levels of the core synthases.

**Table S4.** Proposed functions of ORFs in the *vpk*-gene cluster.

**Table S5.** Strains and plasmids used in this study.

**Table S6.** List of protospacers and PAM sequences of each target locus in this study.

**Table S7.** Oligonucleotides used in this study.