Complete genome sequencing and antibiotics biosynthesis pathways analysis of *Streptomyces lydicus* 103

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More and more new natural products have been found in *Streptomyces* species, which become the significant resource for antibiotics production. Among them, *Streptomyces lydicus* has been known as its ability of streptolydigin biosynthesis. Herein, we present the genome analysis of *S. lydicus* based on the complete genome sequencing. The circular chromosome of *S. lydicus* 103 comprises 8,201,357 base pairs with average GC content 72.22%. With the aid of KEGG analysis, we found that *S. lydicus* 103 can transfer propanoate to succinate, glutamine or glutamate to 2-oxoglutarate, CO2 and L-glutamate to ammonia, which are conducive to the the supply of amino acids. *S. lydicus* 103 encodes acyl-CoA thioesterase II that takes part in biosynthesis of unsaturated fatty acids, and harbors the complete biosynthesis pathways of lysine, valine, leucine, phenylalanine, tyrosine and isoleucine. Furthermore, a total of 27 putative gene clusters have been predicted to be involved in secondary metabolism, including biosynthesis of streptolydigin, erythromycin, mannopeptimycin, ectoine and desferrioxamine B. Comparative genome analysis of *S. lydicus* 103 will help us deeply understand its metabolic pathways, which is essential for enhancing the antibiotic production through metabolic engineering.

*Streptomyces* species are high-GC Gram-positive bacteria found predominantly in soil1. Through a complex process of morphological and physiological differentiation, *Streptomyces* species could produce many specialized metabolites used for agricultural antibiotics2. Some fungi can degrade difficult decomposition by lipase and cellulase, which play an important role in soil ecology3. Besides, the resistance genes of insecticide and herbicide in *Streptomyces* are widely used in transgenic plants4. These secondary metabolites are not essential for bacterial growth but have important roles in microbe-microbe communication5. As a root-colonizing actinomycete, *Streptomyces lydicus* can produce antibiotics or siderophore for suppressing fungal growth6. The elucidation of the related antimicrobial mechanism will facilitate the finding of novel antibiotics.

With the development of genome sequencing technology, more and more complete genomes of *Streptomyces* species have been announced. *S. lydicus* could produce streptolydigin which acts on catalytic function of RNA polymerase and inhibits RNA synthesis7. Our previous studies have identified its biosynthesis pathways of fatty acids8, type II thioesterase9 and nitrogen metabolism10 which are responsible for streptolydigin biosynthesis. Besides, proteomics and metabolomics approaches have been demonstrated in our previous studies on the responses of *S. lydicus* to pitching ratios during streptolydigin production11,12. However, only one complete genome sequence of *S. lydicus*, i.e., *S. lydicus* A02 (accession number CP007699.1), was available in GenBank. Therefore, we have carried out the complete genome sequencing of *S. lydicus* 103 and constructed its metabolic pathways of antibiotic biosynthesis, including primary metabolism and secondary metabolism. Previous work has shown that heterologously expression of *chit42* gene from *Trichoderma harzianum* P1 in *S. lydicus* A01 could enhance the chitinase activity and natamycin production13. Further functional characterization of the gene cluster

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will advance our understanding of the related pathways of antibiotic biosynthesis, and provide insight into the further analysis of the metabolism and gene targets for strain improvement.

**Results**

**Genomic characteristics of *S. lydicus* 103.** *S. lydicus* 103 has a chromosome of 8.20 Mb with 72.22% GC content, which contains 6,872 annotated protein-coding genes (Fig. S1 and Table S1). Mostly, the chromosomes of *Streptomyces* species are linear\(^{14}\). However, the chromosome of *S. lydicus* 103 in this study is circular, which may lead to more genetic stability. Phylogenetic analysis of *S. lydicus* 103 with other *Streptomyces* species has been carried out using CVTree (Fig. S2). BLASTP searches have been performed based on the whole amino acid sequences of *S. lydicus* 103 against those of other *Streptomyces* genomes listed in Fig. S2 with E-values less than $10^{-5}$. The protein-coding genes with the percent of identity and coverage larger than 80% in all *Streptomyces* genomes listed in Fig. S2 are defined as core genes (Figs 1 and 2). The protein-coding genes that have no hits...
### Table 1. Putative gene clusters coding for secondary metabolites in *S. lydicus* 103. Secondary metabolite types detected by antiSMASH: **T1pks** Type I PKS cluster; **T2pks** Type II PKS cluster; **T3pks** Type III PKS cluster; **NRPs** Nonribosomal peptide synthetase cluster; **Bacteriocin** Bacteriocin or other unspecified ribosomally synthesized and post-translationally modified peptide product (RiPP) cluster; **Lasso** peptidase cluster; **Other** cluster containing a secondary metabolite-related protein that does not fit into any other category. The "similarity" means the percentage of the homologous genes in the query cluster that are present in the hit cluster. According to the definition by the antiSMASH, the homologous genes were selected by BLAST e-value < 1E-05, 30% minimal sequence identity, shortest BLAST alignment covers over >25% of the sequence.

| Type                      | From (bp) | To (bp) | Most similar known cluster                  | Similarity |
|---------------------------|-----------|---------|---------------------------------------------|------------|
| Lantipeptide              | 140342    | 176012  | Chlorzadine A biosynthesis                   | 11%        |
| Lassopeptide              | 365678    | 367939  | SapB biosynthesis                           | 100%       |
| NRPs                     | 1043276   | 1065896 | —                                            | —          |
| Siderophore               | 4508815   | 4560693 | A-500359 biosynthesis                        | 10%        |
| Ectoine                   | 7791219   | 7850756 | Mannopeptimycin biosynthesis                 | 81%        |
| Bacteriocin               | 2288536   | 2299192 | Ectoine biosynthesis                         | 75%        |
| S. lividans               | 2380917   | 2392976 | Desferrioxamine B biosynthesis               | 80%        |
| Butyrolactone             | 6256663   | 6271630 | —                                            | —          |
| Terpene                   | 3366951   | 3377184 | —                                            | —          |
| Butyrolactone             | 4574162   | 4586126 | —                                            | —          |
| Bacteriocin               | 5991578   | 6003917 | —                                            | —          |
| Siderophore               | 4485349   | 4511671 | Isorenieratene biosynthesis                  | 100%       |
| Lassopeptide              | 5409699   | 5436377 | Hopene biosynthesis                          | 69%        |
| Terpene                   | 5579311   | 5600591 | Kanamycin biosynthesis                       | 46%        |
| Butyrolactone             | 7923608   | 7945845 | Salinomycin_biosynthesis                     | 4%         |
| Other                     | 4723717   | 4734682 | Hygrocin biosynthesis                        | 6%         |
| Butyrolactone             | 6059826   | 6070904 | —                                            | —          |
| T1pks                     | 5055818   | 5097455 | A-503083 biosynthesis                        | 7%         |
| T2pks                     | 632282    | 678623  | —                                            | —          |
| T1pks-NRPs                | 3269415   | 3311930 | Spore pigment biosynthesis                   | 83%        |
| T1pks-Terpene-NRPs        | 679064    | 735066  | Erythromycin biosynthesis                    | 8%         |
| NRPs-T1pks                | 2855210   | 2966492 | Streptolydigin biosynthesis                  | 97%        |
| NRPs-T3pks                | 3761274   | 3830863 | SW-163 biosynthesis                          | 10%        |
| NRPs-Melanin              | 6090695   | 6154188 | Arylomycin biosynthesis                      | 55%        |
| Thiopptide-Lantipeptide   | 4933499   | 4994082 | WS9326 biosynthesis                          | 10%        |
| Lassopeptide-NRPs-Nucleoside | 3985339   | 4037579 | Cyclothiazomycin biosynthesis                | 38%        |
| Lassopeptide              | 5150871   | 5208226 | Toyocamycin biosynthesis                     | 30%        |

The related metabolism of streptolydigin synthesis in *S. lydicus* 103. Primary metabolism significantly influences secondary metabolism and serves as building precursors for antibiotic biosynthesis, including acetyl-CoA, glucose-6-phosphate, glyceraldehyde-3-phosphate, and oxaloacetate. With the aid of KEGG analysis, metabolic network was obtained, including the central carbon metabolism, nitrogen, amino acids and fatty acid metabolism, and stress responses. The regulatory mechanism of TA loci in *S. lydicus* 103 may be necessary to the environmental stress responses and complex secondary metabolisms.
In the central carbon metabolism, \textit{S. lydicus} 103 has the complete glycolysis, citrate cycle and pentose phosphate pathway. Acyl-CoA is the important precursor of acetyl-CoA, malonyl-CoA, methylmalonyl-CoA, and ethylmalonyl-CoA (Fig. 3). Cutting phosphofructokinase would transfer carbon metabolic flux of glycolytic pathway to the pentose phosphate pathway, and acetyl-CoA could be significantly accumulated and further converted to antibiotics and pyruvate\cite{18}. In the carbohydrate metabolism, \textit{S. lydicus} 103 harbors the complete pathway that transfers xylitol to D-ribulose-5P, involving pentose phosphate pathway, and contains endoglucanase and beta-glucosidase, which transfer cellulose to glucose. Furthermore, \textit{S. lydicus} 103 contains the PTS system and sugar-specific component, thus utilizing the extracellular trehalose and maltose. In the propanoate metabolism, \textit{S. lydicus} 103 harbors the complete pathway that transfers propanoate to succinate, involving pyruvate metabolism. In the nitrogen metabolism, we found two cycle pathways to transfer CO2 and L-glutamate to ammonia, respectively.

In the fatty acids biosynthesis, \textit{S. lydicus} 103 lacks the 3-hydroxyacyl-[acyl-carrier-protein] dehydratase, which is responsible for the dehydration step of the dissociated (type II) fatty-acid biosynthesis system\cite{19}. Moderate control of lipids biosynthesis may distribute more coenzyme A to the streptolydigin biosynthesis. In the fatty acids degradation, \textit{S. lydicus} 103 harbors the complete pathways that transfer propanoate to succinate, involving pyruvate metabolism. In the nitrogen metabolism, we found two cycle pathways to transfer CO2 and L-glutamate to ammonia, respectively.

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Among the amino acids, glutamic acid was the most favorable as the nitrogen source to form streptolydigin\cite{20}. In the glutamine and glutamate metabolism, \textit{S. lydicus} 103 contains the complete pathway to transfer glutamine or glutamate to 2-oxoglutarate, supplying the citrate cycle (Fig. 3). In the cysteine and methionine metabolism, \textit{S. lydicus} 103 harbors the complete pathways to transformation among the L-cysteine, pyruvate, L-homocysteine and L-methionine. L-methionine was not the direct precursor for streptolydigin biosynthesis, but it provided N-methyl of streptolydigin through S-adenosylmethionine, which was catalyzed by S-adanosylmethionine synthase. In the lysine degradation, \textit{S. lydicus} 103 lacks lots of related genes, thus restricting the supplement of acetyl-CoA. In the valine, leucine and isoleucine degradation, \textit{S. lydicus} 103 lacks the 2-oxoisovalerate dehydrogenase E1 component alpha subunit [EC:1.2.4.4] and 2-oxoisovalerate dehydrogenase E2 component (dihydrolipoyl transacylase) [EC:2.3.1.168], thus influencing the biosynthesis of branched chain fatty acid and terpenoid backbone. \textit{S. lydicus} 103 harbors the complete pathways of lysine, valine, leucine, phenylalanine, tyrosine and isoleucine biosynthesis. In addition to the acyl-CoA, L-valine contributes to the biosynthesis for methylmalonyl-CoA and ethylmalonyl-CoA, and L-methionine contributes to the biosynthesis for chloroethylmalonyl-CoA. Proteomics and metabolomics analyses showed that the pitching ratio influenced the activity of glutamate and proline pathways (both precursors of streptolydigin), and exogenously addition can increase the yield of streptolydigin production\cite{21}. We found that \textit{S. lydicus} 103 harbors the complete pathways to transform among the arginine, ornithine, glutamate and proline.

Streptolydigin was a polyketide compound synthesized by type I polyketide pathway, which shares same or similar precursors with the type II polyketide pathways\cite{22}. Complete biosynthetic pathway of streptolydigin has been identified, so that the combined and metabolic processes could be further interpreted\cite{23}. \textit{S. lydicus} 103 harbors 67 ORFs covering a region of 111.2 kb, which are putatively assigned as streptolydigin biosynthesis genes encoding amino-acid permease, isocitrate dehydrogenase, lysophospholipase, erythronolide synthase, phenolphthiocerol synthesis polyketide synthase, cadicidin biosynthesis thioesterase, squalene cyclase, cytochrome P450, methylmalonyl-CoA mutase, glucose-1-phosphate thymidylyltransferase, lipopolysaccharide, biosynthesis protein and electron transfer flavoprotein etc.
The analysis of secondary metabolite pathways in *S. lydicus* 103. *S. lydicus* can produce a lot of important secondary metabolites, and a total of 27 gene clusters were predicted to be involved in secondary metabolism by antiSMASH. They are mainly focused on polyketide (PKS), nonribosomal peptide (NRP)S and terpene, and most of them have the really low similarity with the known clusters (Table 1).

As the typical PKS I, the biosynthetic pathway of erythromycin has been illuminated, including 6-deoxyerythronolide B (6-deB) biosynthesis and glycosylation modification24. The 6-deB was condensed by a molecule propionyl CoA and 6 molecules methyl malonyl CoA. The PKS gene cluster of erythromycin contains eryA I, eryA II and eryA III and encodes acyl wansferase, acyl carrier protein, ketosynthase, ketoreductase, dehydratase and enoyl reductase. The improvements of the erythromycin yield by metabolic engineering has been reported25. The product of 6-deB and erythromycin A was reported in titers of 10 mg·L\(^{-1}\)26. As the typical NRPs, mannopeptimycin was first found in industrial bacterium *Streptomyces hygroscopicus*27. Mannopeptimycin comprises two distinct stereoisomers of amino acids, thus conforming glycosylated cyclic hexapeptide. Besides, with the different R groups, it can form diverse secondary metabolites. We identified a biosynthetic cluster showing 81% similarity with known mannopeptimycin biosynthetic cluster (BGCO00388_c1), which consists of poly-preynl mannose synthase MppG, poly-prenyl phospho-mannosyltransferase MppHIII, mannopeptimycin peptide synthetase MppAB, alpha/beta hydrolase MppK, ABC transporter MppL, isovaleryltransferase MppMN, PLP-dependent aminotransferase MppQ, putative transcriptional regulator MppS, hypothetical protein MppT, two component response regulator MppU, two component sensor kinase MppV, hypothetical lipoprotein MppW, ABC transporter MppX, conserved hypothetical protein MppYZ in *S. lydicus* 103.

Besides, *S. lydicus* 103 harbors the ectione biosynthetic pathway that shows 47% similarity with *Streptomyces albulus* PD-1. As one kind of compatible solute, ectione can be used for protecting enzymes, membranes and whole cells against stresses28. The formation of hydroxyectoine in the ectoine producer *Halomonas elongatwast* was improved by the heterologous expression of the ectione hydroxylase gene from *Streptomyces chrysoomallus*29. We identify two ectione dioxygenases (EctD), L-ectoine synthase (EctC), diaminobutyrate-pyruvate aminotransferase (EctB) and L-2,4-diaminobutyric acid acetyltransferase (EctA) in *S. lydicus* 103, which shows 75% similarity with known ectione biosynthetic cluster (BGCO003853_c1). As the family of siderophores, desferrioxamines can form strong hexadentate complexes with ferric iron. Desferrioxamine B has been used for the treatment of iron overload in human30. *S. lydicus* 103 harbors the desferrioxamine B biosynthetic pathway that shows 80% similarity with known desferrioxamine B biosynthetic cluster (BGCO000941_c1). Previous research has unambiguously identified desferrioxamine E as the major desferrioxamine siderophore produced by *S. coelicolor* M145 and has identified a cluster of four genes (desA-D) that directs desferrioxamine biosynthesis in this model actinomycete31. We also identify tetracopeptide (TPR) protein, DesD-A, HTD domain of SpoOII/ParA/ParB/repB family, 4-nitrophenylphosphatase, desferrioxime E transporter and ABC-type Fe\(^{3+}\)-siderophore transport system in *S. lydicus* 103.

Discussion

Although streptolydigin produced by *S. lydicus* has the activities mentioned above, the yield from the original strain is not very high yet. To achieve higher antibiotic streptolydigin productivity through metabolic regulation, propionate was fed during the fermentation of *S. lydicus*32. The streptolydigin yield, and the carbon fluxes of pentose phosphate pathway and the anaplerotic reaction were significantly increased after propionate feeding. However, it is very difficult to sharply improve the antibiotic production only by the traditional fermentation optimization and mutagenesis treatment. So it is urgent for us to make clear the metabolic network for antibiotic biosynthesis pathways to further improve the production. For example, the cluster slgE1-slgE2-slgE3 is involved in 3-methylaspartate (the precursor of the tetramic acid) supply. SlgE3, a ferredoxin-dependent glutamate synthase, is responsible for the biosynthesis of glutamate from glutamine and 2-oxoglutarate. The expression of slgE3 is increased up to 9-fold at the onset of streptolydigin biosynthesis33. The asparaginyl-tRNA synthetase-like SlgZ and methyltransferase SlgM enzymes are involved in the biosynthesis of the tetramic acid in *S. lydicus*. Over-expression of slgZ and slgM in *S. lydicus* led to a considerable increase in streptolydigin production34. SlmM gene overexpression with different promoters can improve the natamycin production in *S. lydicus* A0235. The biosynthetic genes or regulatory elements of a metabolite must be characterized prior to metabolic engineering36. Furthermore, modifications to the structures of secondary metabolites can often change the biological activity of the compound37. In this study, we presented the complete genome sequence of *S. lydicus* 103 and identified the pathways related to streptomyces biosynthesis from primary metabolism to secondary metabolite, which would provide more accurate analysis of the metabolic network and a more rational adjustment of metabolic regulation38.

Genomics-based bottom-up approaches have been developed to unveil biosynthetic pathways of new natural products that were undetected under standard fermentation conditions39. Despite being tapped as antibiotic sources for decades, *Streptomyces* spp. could produce up to 100,000 antimicrobial metabolites, while only a small proportion have been identified40. As an example, a terpene synthase from *S. avermitilis* was expressed in *E. coli*, resulting in the synthesis of the novel tricyclic sesquiterpene, avermitilol41. Chu et al.42 used primary sequence from the human microbiome, and thus bioinformatically predicted and chemical synthesized a new antibiotic. Luo et al.43 applied a plug-and-play synthetic biology strategy to activate a cryptic polycyclic tetramate macrolactams (PTMs) biosynthetic gene cluster from *S. griseus* and discovered three new PTMs. Besides, transcriptome and metabolome can identify the potential biosynthetic genes by correlating the expression of the secondary metabolite related gene44. In *S. lydicus* 103, we found many new gene clusters that have really low similarity with known clusters (Table 1). Thus, further studies are desirable for optimization, isolation and identification of the new bio-active molecule. The availability of the genome sequence of *S. lydicus* 103 provides a framework for biotechnological analysis and characterization of new natural products.
Methods

Bacterial culture and genome sequence. S. lydicus 103, an actinomycete, was isolated from soil. One loop of cells was incubated in a 250 mL flask containing 50 mL seed medium for 48 hours at 28 °C with shaking at 220 r·min⁻¹. The seed medium contained (g·L⁻¹): glucose 5, starch 30, yeast extract 2, peptone 4, K₂HPO₄ 1.5, NaCl 0.5, and MgSO₄·7H₂O 0.5. Isolation of genomic DNA was carried out using SDS method. Total DNA obtained was subjected to quality control by agarose gel electrophoresis and quantified by Qubit. The genome was sequenced by Single Molecule, Real-Time (SMRT) technology. Sequencing was performed at the Beijing Novogene Bioinformatics Technology Co., Ltd. SMRT Analysis 2.3.0 was used to filter low quality reads and the filtered reads were assembled to the chromosome without gaps. The circular skeleton of chromosome was identified by the long fragment across the head and tail.

Genome annotation and bioinformatics analysis. Transfer RNA (tRNA) genes, Ribosome RNA (rRNA) genes, small RNA (sRNA) genes were predicted with tRNAscan-SE65, rRNAmer48 and Rfam database56, respectively. Gene prediction was performed with the integrated model by NCBI prokaryotic annotation pipeline66, and gene functional prediction was performed by Blast64 against the databases, KEGG59 (Kyoto Encyclopedia of Genes and Genomes), COG67 (Clusters of Orthologous Groups), Swiss-Prot52, and GO53 (Gene Ontology). The origin of replication (oriC) and putative DNA boxes were identified using Ori-Finder54. GC-Profile was used to compute the GC content variation in genome sequence and predict the genomic islands65. GenoView Server, a comparative genomics tool for circular genomes, was used to obtain a circular graphical representation of chromosome. A whole genome-based, alignment-free composition vector (CV) method was performed for phylogenetic analysis57 and the phylogenetic tree was generated using the MEGA program58. The toxin-antitoxin (TA) systems were predicted by TADB69. Secondary metabolite gene clusters were predicted by antiSMASH60. BioVenn, a web application for the comparison and visualization of biological lists, was used for Venn diagrams drawing61.

GenBank accession number. The sequence of the S. lydicus 103 genome has been deposited at DDBJ/EMBL/GenBank under the GenBank accession number CP017157.

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Author Contributions

F.G., M.Z.D. and Y.Y.J. designed the project and experiments; N.J. performed the experiments; F.G. and Y.Y.J. contributed reagents/materials/analysis tools; N.J., M.Z.D., H.L. and F.G. analyzed the final data and wrote the manuscript. All the authors read and approved the final version of the manuscript.
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