Identification and Heterologous Expression of the Chaxamycin Biosynthesis Gene Cluster from *Streptomyces leeuwenhoekii*

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*Streptomyces leeuwenhoekii*, isolated from the hyperarid Atacama Desert, produces the new ansamycin-like compounds chaxamycins A to D, which possess potent antibacterial activity and moderate antiproliferative activity. We report the development of genetic tools to manipulate *S. leeuwenhoekii* and the identification and partial characterization of the 80.2-kb chaxamycin biosynthesis gene cluster, which was achieved by both mutational analysis in the natural producer and heterologous expression in *Streptomyces coelicolor* A3(2) strain M1152. Restoration of chaxamycin production in a nonproducing ΔcxmK mutant (cxmK encodes 3-amino-5-hydroxybenzoic acid [AHBA] synthase) was achieved by supplementing the growth medium with AHBA, suggesting that mutasynthesis may be a viable approach for the generation of novel chaxamycin derivatives.

Antibiotic resistance is rapidly becoming a worldwide medical problem that requires urgent attention (1–3). Historically, natural products produced by actinomycetes and fungi have been the major source of clinically used antibiotics (4). However, the repeated rediscovery of known chemical entities from natural sources has led to a marked decline in the discovery of novel microbial antibiotics with clinical utility (5, 6). One potential solution to this problem is to screen microorganisms isolated from previously little-studied ecological niches, and bioprospecting of unexplored environments has indeed already resulted in the discovery of novel microorganisms and new chemical scaffolds (4, 6).

In 2004, a considerable number of new actinomycetes were, surprisingly, isolated from the hyperarid Atacama Desert, the oldest and driest desert on Earth, located in northern Chile (7, 8). Strikingly, some of these strains produce novel chemical structures with potent biological activity (9–11). One of the strains isolated from this environment is *Streptomyces leeuwenhoekii* DSM 42122 (previously known as *Streptomyces* strain C34) isolated from the Chaxa Lagoon (7, 12). It produces the new polyketide antibiotics called the chaxamycins (9) and the chaxalactins (10). Chaxamycin A to D (compounds 1 to 4 in Fig. 1) are four new naphthalenic ansamycin-type polyketides derived from a 3-amino-5-hydroxybenzoic acid (AHBA) starter unit. Chaxamycin D (compound 4) displays, through an unknown mechanism, highly selective antimicrobial activity against *Staphylococcus aureus* ATCC 25923 and against a panel of methicillin-resistant *S. aureus* (MRSA) strains (9), while chaxamycin A (compound 1) inhibits the intrinsic ATPase activity of the human Hsp90 protein, which is involved in cancer proliferation (9). The chaxamycins differ from other naphthalenic ansamycins, such as the rifamycins, naphthomycin, geldanamycin, and the herbimycins, in lacking a methyl group at the olefinic C-16 next to the amide bond present in all of these molecules.

The biosynthesis of ansamycin-type polyketides commences with priming of the polyketide synthase (PKS) with the precursor molecule AHBA. In the case of the rifamycins and other ansamycin-type polyketide gene clusters, a sub-gene cluster, located near the PKS, is largely responsible for the synthesis of AHBA (13). One of the enzymes encoded in the core of this AHBA gene cluster is AHBA synthase, which converts 5-deoxy-5-amino-3-dehydroshikimic acid (amino-DHS) into AHBA, the final step in its biosynthesis (14), and an AHBA synthase gene indeed was identified previously by PCR analysis of genomic DNA of *S. leeuwenhoekii* (9). In addition to AHBA, we predict that chaxamycin is synthesized from 10 additional extender units (three malonyl coenzyme As [malonyl-CoAs] and seven methylmalonyl-CoAs) by a modular type 1 PKS.

Recent improvements in next-generation sequencing technologies and genome mining have markedly accelerated the identification and characterization of gene clusters for specialized metabolites, many potentially encoding novel compounds with antimicrobial activity. In this work, we describe the application of these technologies to identify and partially characterize the chaxamycin biosynthesis gene cluster of *S. leeuwenhoekii*.

**MATERIALS AND METHODS**

Strains, culture conditions, and general methods. The bacterial strains used and generated during this study are listed in Table 1. *Escherichia coli* and *Streptomyces* strains were cultured, maintained, and manipulated genetically following the methods of Sambrook et al. (15) and Kieser et al. (16), respectively. The plasmids and oligonucleotides used or constructed during this work are listed in Table 2 and Table 3, respectively. Mobilization of phage P1-derived artificial chromosome (PAC) clones into *Streptomyces* strains was performed as described previously (17). Molecular biology enzymes, reagents, and kits were used according to the manufac-
turer’s instructions. High-fidelity PCR amplification was performed with Phusion or Q5 DNA polymerase following the manufacturer’s instructions (NEB, Ipswich, MA) with a nucleotide proportion of 15A:15T:35G:35C to improve the amplification efficiency of high-moles-percent G+C Streptomyces DNA. 3-Amino-5-hydroxybenzoic acid (AHBA) was purchased from Sigma (catalog no. PH011754) and dissolved in dimethyl sulfoxide.

**Specific methods for S. leeuwenhoekii.** Spore stocks of *S. leeuwenhoekii* were prepared following standard methods (16) from cultures grown on soy flour-mannitol (SFM) agar medium at 37°C for 5 to 7 days. Mobilization of plasmids into *S. leeuwenhoekii* was performed as established for *Streptomyces coelicolor* A3(2) by conjugation from *E. coli* ET121567 (carrying either pUZ8002 or pR9406) using 10⁸ spores that were heat shocked in 2% YT medium at 50°C for 10 min; the treated spores were allowed to cool to room temperature before mixing them with the *E. coli* cells, and the conjugation mixtures were plated on SFM agar medium supplemented with 10 mM MgCl₂ and 10 mM CaCl₂.

**Production and measurement of chaxamycins.** *S. leeuwenhoekii* strains were grown in modified ISP2 medium (containing, per liter, 4.0 g of yeast extract, 10.0 g of malt extract, and 10.0 g of glycerol) pH 7.2 with NaOH. All strains were cultured in 250-ml Erlenmeyer flasks that had been treated with 5% dimethyldichlorosilane (DMDCS) in toluene (Sylon CT; Supelco, catalog number 33065-U) to deter the mycelium from sticking to the glass; a stainless steel spring was placed at the bottom of each flask to improve aeration. Seed cultures were inoculated with 10⁶ spores and incubated at 30°C with orbital shaking (250 rpm) for 24 to 48 h. Mycelium from the seed cultures was harvested by centrifugation, washed with 0.9% NaCl, resuspended in fresh medium, and used to inoculate production cultures to a final optical density at 600 nm (OD₆₀₀) of 0.2; production cultures were incubated at 30°C with orbital shaking (250 rpm). Samples were harvested at the indicated times: 1 or 2 ml of homogenized culture was pipetted into a clean Eppendorf tube and centrifuged at 13,000 rpm at 4°C for 10 min, the supernatant was moved to a new Eppendorf tube and centrifuged again at 13,000 rpm at 4°C for 10 min before carefully pipetting 0.5 ml of the supernatant into a high-pressure liquid chromatography (HPLC) vial, the mycelial pellet was extracted with 0.5 ml of methanol and centrifuged at 13,000 rpm at 4°C for 10 min, and 0.3 ml of the methanol extract was pipetted into an HPLC vial containing 0.3 ml of water. Chaxamycins were detected in both the supernatants and the mycelial extracts, but larger amounts were always found in the supernatants, and thus only these fractions were analyzed in most experiments. When analyzing total chaxamycin production in a single HPLC run, both fractions were mixed in a 1:1 ratio (thus injecting samples in 50% methanol). Samples were analyzed on a Shimadzu liquid chromatography-mass spectrometry (LC-MS) system equipped with a NexeraX2 liquid chromatograph (LC30AD) fitted with a Prominence photodiode array detector (SPD-M20A) and an LCMSIT-TOF mass spectrometer; samples (typically 5 μl) were injected in a Kinetex XB C₁₈ 2.6-μm, 100-Å, 50-2.10-mm column (part no. 00B-4496-AN; Phenomenex, USA) fitted with a KrudKatcher Ultra HPLC in-line filter (part no. AF0-8497; Phenomenex, USA) and eluted at a flow rate of 0.6 ml per min with a gradient of 0.1% formic acid in water (mobile phase A) and methanol (mobile phase B) as follows (minute, percentage of B reached at that minute): 0 min, 2% B; 1 min, 2% B; 8 min, 100% B; 9.5 min, 2% B; and 11.2 min, 2% B for equilibration. The column was kept at 40°C. Mass spectrometry detection was performed in negative ion mode since all of the chaxamycin species were detected with higher sensitivity than in positive mode. Purified chaxamycin standards for compounds 1, 2, and 3 were kindly provided by Mostafa Rateb and Marcel Jaspars (University of Aberdeen, Scotland). Data acquisition and analysis were performed with LCMSolutions version 3 (Shimadzu); spectrum visualization was also performed with Mass ++ version 2.7.2 (http://www.first-ms3d.jp/english/). The (M – H)⁺ ions detected for the chaxamy-

### Table 1 Bacterial strains used in this study

| Species and strain | Description | Reference or source |
|--------------------|-------------|---------------------|
| **Escherichia coli** | Strain used for routine cloning | 65 |
| DH5α | Strain used for routine cloning | 65 |
| DH10B | Methylation-deficient strain used for conjugation with *Streptomyces*; pUZ8002 provides conjugation machinery | 66 (pUZ8002, J. Wilson and D. Figurski, unpublished) |
| ET121567/pUZ8002 | Strain used for routine cloning carrying conjugation plasmid pR9406 | pR9406, A. Siddique and D. Figurski, unpublished |
| TOP10/pR9406 | Strain used for routine cloning carrying conjugation plasmid pR9406 |  |
| **Streptomyces coelicolor** | M1152 | M145 Δact Δred Δpck Δeda rpoB(C1298T) | 67 |
| M1650 | M1152 carrying pJ12853 | This work |
| **Streptomyces leeuwenhoekii** | M1653 | S. leeuwenhoekii ΔconK::neo | This work |
| M1655 | S. leeuwenhoekii/pGUS | This work |
| M1656 | S. leeuwenhoekii/pJ10740 | This work |

*FIG 1 Structures of chaxamycins A to D (compounds 1 to 4).*
Amplify upstream flanking region of AHBA synthase gene with pIJ12852, pIJ10257 derivative with pIJ12861, pBluescript II KS(−)

pIJ12860 Derivative of pGM1190 with JFC010/JFC009-JFC011/JFC012 fragment; for deletion of

pIJ12858 Derivative of pKC1132 with JFC011/JFC012 fragment; for deletion of

pIJ12857 Derivative of pKC1132 with JFC010/JFC009 fragment; for deletion of S. leeuwenhoekii

assembly of the genome sequence of elsewhere (19). Confirmatory Sanger sequencing was performed using the pIJ12853 Derivative of pESAC13; contains 145 kb of

pIJ12850 Derivative of pGM1190 with JFC010/JFC009-JFC054 TTCCGGAAGATGAACGTGAC

pIJ12849 Derivative of pESAC13 with JFC026/JFC034 fragment; for confirmation of deletion of AHBA synthase gene

TABLE 2 Plasmids used and constructed during this study

| Plasmid          | Description                                                                 | Reference or source |
|------------------|------------------------------------------------------------------------------|---------------------|
| pBluescript II KS(+) | General cloning vector                                                         | 68                  |
| pKC1132          | Cloning vector, conj (oriT from RK2)                                          | 69                  |
| pSET152          | Cloning vector, conj (oriT from RK2), integrative (phiC31 attP)               | 69                  |
| pTGC192-Km       | Source of neo (kanamycin resistance gene)                                     | 70                  |
| pGM1190          | pG5 deriv, tsr, aac(3) IV, oriT, to terminator, P_repA, RBS, fd terminator   | 40 (pGM1190, G. Muth, unpublished) |
| pESAC13          | PAC vector (P1 phage rep) for genomic library construction; conj (oriT from RK2), integrative (phiC31 attP), tsr, neo, P1 rep, sacB | 27 (pESAC13, M. Sosio, unpublished) |
| pGUS             | gusA (β-glucuronidase gene, codon-optimized for streptomycetes) in pSET152    | 38                  |
| pIJ10740         | pGUS deriv with ermE* p driving gusA transcription                            | Morgan Feeney, unpublished |
| pIJ10257         | Expression vector for Streptomyces, with ermE*, hyg, conj (oriT from RK2), integrative (phiC31 attP) | 71                  |
| pIJ12850         | Derivative of pGM1190 with JFC010/JFC009-neo-JFC011/JFC012 fragments; for deletion of cxmK | This work |
| pIJ12851         | Derivative of pKC1132 with JFC010/JFC009-neo-JFC011/JFC012 fragments; for deletion of cxmK | This work |
| pIJ12857         | Derivative of pKC1132 with JFC010/JFC009 fragment; for deletion of cxmK     | This work |
| pIJ12858         | Derivative of pKC1132 with JFC011/JFC012 fragment; for deletion of cxmK     | This work |
| pIJ12859         | Derivative of pKC1132 with JFC010/JFC009-JFC011/JFC012 fragment; for deletion of cxmK | This work |
| pIJ12860         | Derivative of pGM1190 with JFC010/JFC009-JFC011/JFC012 fragment; for deletion of cxmK | This work |
| pIJ12861         | pBluescript II KS(+) deriv with JFC026/JFC034 fragment; for cloning of cxmK | This work |
| pIJ12852         | pIJ10257 deriv with cxmK; for complementation of deletion mutant            | This work |
| pIJ12853         | Derivative of pSAC13; contains 145 kb of S. leeuwenhoekii chromosome, including the chaxamycin biosynthesis gene cluster | This work |

DNA sequencing and bioinformatic analysis. The sequencing and assembly of the genome sequence of S. leeuwenhoekii has been reported elsewhere (19). Confirmatory Sanger sequencing was performed using the Eurofins Genomics service (Ebersberg, Germany) and analyzed using the Staden package (20) version 2.0.069 (http://staden.sourceforge.net). BLAST (21) searches were performed at the NCBI server. Local BLAST searching was performed using NCBI package 2.2.26+ (22) and Sequenc-eServer (23) or pfectBLAST (24). Automatic primer design was carried out using the Primer3Plus web server (25). DoBISCUIT (http://www.bio.nite.go.jp/pks/top) (Database of Biosynthesis clusters Curated and Integrated) (26) was used to obtain curated and annotated versions of ansamycin-related gene clusters. The sequence of phase I-derived artificial chromosome (PAC) clone 2-11L (named pIJ12853), selected from a library of S. leeuwenhoekii genomic DNA made in pESAC13 (E. coli-Streptomyces artificial chromosome [27]; http://www.biost.com/page/PAC.aspx) and containing the chaxamycin biosynthesis gene cluster (see “PAC library construction and PCR screening for the chaxamycin gene cluster” below) was obtained using a Roche 454-Junior sequencer (commissioned from the DNA sequencing facility in the Department of Biochemistry, University of Cambridge, Cambridge, United Kingdom). A high-quality sequence of the complete PAC insert was obtained by mapping the assembled contigs from the 454-Junior sequencing onto the genome sequence (19) using the Burrows Wheeler Aligner BWA-MEM algorithm with default parameters (28); the resulting BAM alignment file was processed with SAMTOOLS (29) and loaded into GAPPS (30) to manually produce a consensus sequence. This consensus was processed with Prodigal (31) to identify protein-coding sequences, automatically annotated using the BAys (32) web server (https://www.basys.ca/), and

TABLE 3 Oligonucleotides used in this studya

| Name | Sequence (5’ → 3’)b | Use |
|------|---------------------|-----|
| JFC009 | TTTCTAGAGGAGCCATCAGCCAGGACT | Amplify upstream flanking region of AHBA synthase gene |
| JFC010 | TTAAGCTTGTAGTCCAGCAGCTCCAGT | Amplify downstream flanking region of AHBA synthase gene |
| JFC011 | TTTCTAGAGCTCGGTCTCGTGCTGACTG | Screening of genomic library for chaxamycin gene cluster |
| JFC012 | TTTGATCCACTGCTCTGCTGCTGACTCG | Screening of genomic library for chaxamycin gene cluster |
| JFC022 | CGATTCGAGAAGTGAGCA | Cloning of cxmK in pIJ10257 |
| JFC023 | GTTTAGTGTTGGCGATTTG | Amplify upstream flanking region of AHBA synthase gene |
| JFC024 | CGCGAGAAGAAGAGCGAGGT | Amplify downstream flanking region of AHBA synthase gene |
| JFC025 | CACACATTGTGACTGAGCAC | Screening of genomic library for chaxamycin gene cluster |
| JFC026 | AGACATATGGAAGGGGACAG | Screening of genomic library for chaxamycin gene cluster |
| JFC034 | AAGCTTATGACCTGTGAGTAG | Cloning of cxmK in pIJ10257 |
| JFC032 | GTCTGTCCTCATTTTCCTC | Confirmation of deletion of AHBA synthase gene |
| JFC033 | GACCTGTTAGCCGCTTGTG | Resolution of a frameshift within one of the PKS genes |
| JFC036 | GAGCAGAGACACACAGGAGGT | Resolution of a frameshift within one of the PKS genes |
| JFC037 | GACAACCGGACACTACGG | Resolution of a frameshift within one of the PKS genes |
| JFC053 | AAGCGAGTTGACGAGATCA | Resolution of a frameshift within one of the PKS genes |
| JFC054 | TTCCGGAAGATGAACTGAGC | Resolution of a frameshift within one of the PKS genes |

a Oligonucleotide pairs are shown in consecutive shaded or unshaded rows, and the information in the “Use” column applies to both relevant oligonucleotides.

b Incorporated restriction sites are underlined.
analyzed and curated with Artemis (33). Analysis of the sequence of the insert of pJJ12853 revealed that it corresponded to nucleotide (nt) positions 1151661 to 1305894 of the published *S. leeuwenhoekii* chromosome sequence (European Nucleotide Archive study accession number PRJEB8583) and contained many of the genes predicted to be required for chaxamycin biosynthesis (located between nt 1211049 and 1289829). GC Frame Plot analysis (34) performed within Artemis revealed a clear frameshift in the 454 sequence within one of the PKS genes. The corresponding region was PCR amplified using primers JFC036 and JFC037 and sequenced with the same primers by Sanger sequencing. Analysis revealed the presence of two close frameshifts (two missing Gs), and the consensus sequence was corrected accordingly (matching that obtained with Illumina MiSeq sequencing of the *S. leeuwenhoekii* genome (19)). GC Frame Plot analysis revealed possible frameshifts between approximate nucleotide positions 1214200 and 1215100. Indeed, BLASTX analyses (21) confirmed that this region had the potential to encode a type II thioesterase (see Discussion) but contained three frameshifts in a putative ancestral coding sequence that begins at nt 1215047 and ends at nt 1214242. To assess the quality of this sequence, the region was PCR amplified using primers JFC053 and JFC054 with pJJ12853 (see below) DNA as the template, and the fidelity of the sequence and the presence of the frameshifts were confirmed by Sanger sequencing using the same primers.

**PAC library construction and PCR screening for the chaxamycin gene cluster.** A genomic library of *S. leeuwenhoekii* DNA was constructed in pESAC13. Briefly, high-molecular-weight DNA was digested partially with BamHI and ligated with pESAC13 digested with the same enzyme, resulting in loss of pUC19 present in the original vector. The genomic library was screened for clones containing the chaxamycin biosynthesis gene cluster by PCR using two primer pairs, JFC022/JFC023 and JFC024/JFC025, designed to amplify the predicted ends of the gene cluster. One PAC clone (2-11L) was identified, named pJJ12853, and sequenced (see “DNA sequencing and bioinformatic analysis” above). Both library construction and screening were carried out by Bio S&T Inc. (Montreal, Canada).

**Construction of *S. leeuwenhoekii* M1653 (∆cxmK::neo).** The AHBA synthase gene (*cxmK*) was deleted and replaced with the kanamycin resistance gene *neo* by double-crossover homologous recombination. Two DNA segments flanking *cxmK*, each about 1.7 kb in length, were PCR amplified with primer pairs JFC010/JFC009 (containing 5′ HindIII and XbaI sites, respectively) and JFC011/JFC012 (containing 5′ XbaI and BamHI sites, respectively) and cloned separately in pKC1132 that had been cleaved with HindIII plus XbaI or XbaI plus BamHI to generate pIJ12852. The HindIII-BamHI insert of pIJ12852 was cloned in the self-replicative temperature-sensitive plasmid pGM1190 that had been cleaved with HindIII and BamHI to yield pIJ12860. Finally, *neo* was excised with XbaI from pTC192-km and cloned in the XbaI site of pJJ12860 to yield pIJ12850. This construct was transferred into *S. leeuwenhoekii* by conjugation, and kanamycin-resistant exconjugants were selected at 30°C, restreaked on SFM agar containing kanamycin, and cultivated at 37°C, a temperature at which pGM1190 cannot replicate, therefore forcing chromosomal integration of neo by either single or double homologous recombination. The resulting spores were plated on S. SFM agar without antibiotic selection and incubated at 30°C, and the resulting colonies were replicated consecutively onto S. SFM agar plates containing just kanamycin, just apramycin, or neither antibiotic, which were then incubated at 37°C. Colonies resistant to kanamycin but sensitive to apramycin were subjected to PCR analysis using primers (JFC032/JFC033), which flanked *cxmK*, and the required ΔcxmK::neo deletion mutant (M1653) was identified (see Fig. S7 in the supplemental material). Note that we resorted to using a derivative of the replicative, multicopy pKM1190 for gene replacement after attempts to use pJJ12851 (Table 2), a derivative of the nonreplicative pKC1132 containing the same deletion cassette as pJJ12850, failed to yield single-crossover recombinants, which possibly was a consequence of a relatively low recombination frequency in *S. leeuwenhoekii*. We hoped to compensate for any such deficiency by using the multicopy, replicative pGM1190 derivative.

**Construction of pJJ12852.** cxmK was amplified from *S. leeuwenhoekii* genomic DNA by high-fidelity PCR using primers JFC026 and JFC034 (containing 5′ Ndel and HindIII sites, respectively); the PCR product was blunt-end ligated into the Smal site of pBluescript II KS(+) to yield pJJ12861, and the sequence of the PCR product was verified by sequencing with universal primers corresponding to the vector. cxmK was excised from pJJ12861 with Ndel and HindIII and inserted into pJJ10257 that had been cleaved with the same enzymes to generate pJJ12852 with cxmK transcribed from the constitutive ermE′ promoter.

**Generation of illustrations.** Illustrations of LC-MS/MS data were generated using Microsoft Office Suite. ImageJ version 1.47 (http://imagej.nih.gov/ij/) and Inkscape version 0.91 (https://inkscape.org) were used to create and edit drawings and illustrations.

**RESULTS** Establishment of microbial and genetic manipulation procedures for *S. leeuwenhoekii*. *S. leeuwenhoekii* grew well on SFM, R2, and DNA agar media, with the most copious and rapid sporulation observed after growth for 4 to 6 days at 30°C on SFM agar (see Fig. S1 in the supplemental material). Since *S. leeuwenhoekii* was isolated from the Atacama Desert (Chile), we compared growth at 30°C, 37°C, and 43°C. Growth and sporulation occurred more rapidly with increasing temperature, with sporulation occurring at 37°C and 43°C after just 2 days of cultivation (see Fig. S2 in the supplemental material). Consequently, we used 37°C to prepare spore stocks, a compromise between high growth rate and sporulation and a temperature at which the medium did not dry out too quickly. Chaxamycin production was also assessed at the same three temperatures in modified ISP2 production medium and shown to be very low at 37°C and 43°C compared to 30°C (see Fig. S3 in the supplemental material); consequently, all studies of chaxamycin production were performed at 30°C.

Before establishing techniques for the genetic manipulation of *S. leeuwenhoekii*, we first tested the susceptibility of the strain to a range of antibiotics commonly used with streptomycetes (16). Growth was prevented at concentrations of >1.0 μg/ml kanamycin, >0.5 μg/ml apramycin, >2.0 μg/ml hygromycin B, and >2.5 μg/ml thiostrepton; *S. leeuwenhoekii* was not sensitive to phosphomycin up to 50 μg/ml, while growth was retarded in the presence of nalidixic acid at >25 μg/ml (see Fig. S4 in the supplemental material).

Many plasmids used to manipulate streptomycetes genetically utilize the attachment sites of temperate phages (16). Putative bacterial attachment (*attB*) sites for the phiC31 and phiBT1 actinophages were identified in the chromosome of *S. leeuwenhoekii* (European Nucleotide Archive accession number PRJEB8583 [19]) by BLAST searches using the sequences of the corresponding sites from *S. coelicolor* (35, 36) (see Table S1 in the supplemental material). In both cases, the putative *attB* sites were located in homologues of the genes in which they reside in *S. coelicolor* (see Table S2 in the supplemental material).

**Conditions for transconjugation** DNA into *S. leeuwenhoekii* by conjugation were established using pSET152, which integrates in the phiC31 *attB* and carries the apramycin resistance gene *aac(3)IV*, and pJJ10257, which integrates at the phiBT1 *attB* and carries the
The hygromycin B resistance gene *hyg*; conjugations were plated on SFM agar supplemented with 10 mM MgCl₂ and 10 mM CaCl₂. The most common constitutive promoter used for gene expression in *Streptomyces* is *ermE*<sup>*</sup> <sup>(37)</sup>. To test its usability in *S. leeuwenhoekii*, we introduced pGUS (a pSET152 derivative with the *gusA* reporter gene <sup>[38]</sup>) and pIJ10740 (a pGUS derivative with *ermE*<sup>*</sup> driving the transcription of *gusA* [Morgan Feeney, personal communication]); *gusA* expression (assessed as described previously [39]) was readily observed in the pIJ10740 exconjugants but not in the pGUS exconjugants or in the *S. leeuwenhoekii* wild-type strain, confirming the utility of *ermE*<sup>*</sup> in this species.

*P. GM1190* is a self-replicative, temperature-sensitive plasmid (reported to be nonreplicative at over 34°C) used for gene replacement in *Streptomyces* (40). We readily obtained *S. leeuwenhoekii* exconjugants with pGM1190 and verified plasmid loss after cultivation at 34°C, 37°C, and 43°C (see Fig. S5 in the supplemental material). Since we observed some persistence of pGM1190 at 34°C, 37°C was used to cure the plasmid.

**Bioinformatic identification of the chaxamycin biosynthesis gene cluster.** Bioinformatic analysis of the genome sequence of *S. leeuwenhoekii* identified a contiguous stretch of DNA that contained many of the genes predicted to be required for chaxamycin biosynthesis, including those needed for AHBA biosynthesis (i.e., the previously sequenced AHBA synthase gene [GenBank accession number FR839674.1]) and a complete set of appropriate PKS modules, including a loading module with similarity to those of other ansamycin-type PKSs such as ansamitocin (DoBISCUIT accession code Ansam_00270), geldanamycin (Gelda2_00080), herbimycin A (Herb_00170), rifamycin (Rifam_00210), and rubradiarin (Rubra_00070).

The putative limits of the gene cluster were identified by comparison with gene clusters encoding other ansamycin-like polyketides, including rifamycin (the *rif* cluster) from *A. mediterranei* S699 (AF040570.3), naphthomycin from *Streptomyces* sp. strain CS (GQ452266.1), and saliniketal from *Salinispora arenicola* CNS-205 (CP000850.1). The right end of the chaxamycin gene cluster (as shown in Fig. 2 and in Fig. S6 in the supplemental material) was defined by the presence of a gene, *cxmJ*, homologous to *rifJ*, encoding a putative aminodehydroquinate (amino-DHQ) dehydratase involved in the biosynthesis of AHBA (catalyzing the
TABLE 4 Proposed chaxamycin biosynthesis gene cluster and gene functions

| sle no. | con name | No. of amino acids | Rifamycin homologue | Proposed function in chaxamycin biosynthesis |
|---------|-----------|--------------------|---------------------|---------------------------------------------|
| sle10390 | conA | 138 | rifA | Small hypothetical protein |
| sle10380 | conS | 327 | rifS | NADH-dependent oxidoreductase |
| sle10370 | conT | 323 | rifT | NADH-dependent dehydrogenase |
| sle10360 | conV | 397 | orfB | Cytochrome P450 |
| sle10350 | conA | 5616 | rifA | Polyketide synthase (module 0 [M0], ASBBK-RDKR-ACP; M1, KS-ATmmal-dh-KR-ACP; M2, KS-ATmmal-dh-ACP; M3, KS-ATmmal-KR-ACP) |
| sle10340 | conB | 5363 | rifB | Polyketide synthase (M4, KS-ATmmal-dh-KR-ACP; M5, KS-ATmmal-dh-KR-ACP; M6, KS-ATmmal-dh-KR-ACP) |
| sle10330 | conC | 1820 | rifC | Polyketide synthase (M7, KS-ATmmal-dh-KR-ACP) |
| sle10320 | conD | 1773 | rifD | Polyketide synthase (M8, KS-ATmmal-dh-KR-ACP) |
| sle10300 | conE | 3488 | rifE | Polyketide synthase (M9, KS-ATmmal-dh-KR-ACP; M10, KS-ATmmal-dh-KR-ACP) |
| sle10290 | conF | 269 | rifF | Polyketide release and ansa-ring formation: amide synthase |
| sle10280 | conG | 368 | rifG | AHBA synthesis, aminodehydroquinic acid synthase |
| sle10270 | conH | 406 | rifH | AHBA synthesis, amino-DHAP synthase |
| sle10260 | conI | 268 | rifI | AHBA synthesis, aminodehydroquinic acid synthase |
| sle10250 | conK | 386 | rifK | AHBA synthesis, AHBA synthase |
| sle10240 | conL | 358 | rifL | AHBA synthesis, oxidoreductase |
| sle10230 | conM | 232 | rifM | AHBA synthesis, AHBA synthase |
| sle10220 | conN | 307 | rifN | AHBA synthesis, AHBA synthase |
| sle10210 | con18 | 295 | orf11 (+) | Flavin-dependent oxidoreductase |
| sle10200 | con19 | 533 | orf19 (−) | Naphthalene ring formation, FDA-dependent-monooxygenase and 3-(3-hydroxyphenyl) propionate hydroxylase |
| sle10190 | con20 | 402 | orf20 | Tailoring, O-acetyltransferase (chaxamycin D) |
| sle10180 | con21 | 63 | | Ferredoxin |
| sle10170 | con22 | 393 | orf4 (+) | Cytochrome P450 monooxygenase |
| sle10160 | con23 | 418 | orf5 (W'), orf13 (+) | Tailoring, cytochrome P450 monooxygenase (hydroxyfuran of chaxamycin D) |
| sle10150 | con24 | 335 | | Tailoring: methyltransferase, S-adenosylmethionine dependent (not homologue of orf4) |
| sle10120 | conY | 433 | | Transcriptional regulator (C-terminal DNA-binding domain found in the NarL/FixJ response regulator family) |
| sle10110 | conZ | 246 | | Transcriptional regulator, atypical response regulator of OmpR/PhoB family |
| sle10100 | conM | 168 | rifM | AHBA synthesis, aminodehydroquinic acid dehydratase |

a Gene names have been assigned as far as possible according to predicted functional homology to the rifamycin gene cluster (see also Table S3 in the supplemental material).

b Rifamycin gene cluster from A. mediterranei S699 (GenBank accession no. AF040570.3). Mutant phenotypes: +, similar production to parent strains; −, production abolished; W, loss of rifamycin B production and accumulation of rifamycin W.

c Based on Pfam motif search and homology to rifamycin biosynthesis proteins. PKS domains (found with antiSMASH [47] and NCBI-CDD [72]): A, adenylation; ACP, acyl carrier protein; AT, acyltransferase; DH, dehydratase (lowercase indicates that it should be inactive); KR, ketoreductase; KS, ketosynthase. The specificity predicted for each AT domain is shown as a subscript: AHBA, 3-amino-5-hydroxybenzoic acid; mal, malonyl-CoA; mmal, methylmalonyl-CoA.

d Genes required for AHBA biosynthesis (43).

e According to reference 56.

f Mutant accumulates linear polyketide intermediates (59).

conversion of 5-deoxy-5-aminodehydroquinic acid to 5-deoxy-5-aminodehydroshikimic acid [amino-DHS] [41]); surprisingly, conM is located some distance from the other genes required for AHBA biosynthesis, as also noted for the conM homologues in the other three gene clusters (13).

The left end of the chaxamycin gene cluster (as shown in Fig. 2) could not be readily predicted, although from comparisons with the other ansamycin-like gene clusters, we initially speculated that conS encoded a putative NADH-dependent dehydrogenase homologous to the rifS product, which would be the likely limit of the cluster. At 8.7 kb upstream of conS (to the left of conS in Fig. 2), we found a conserved region of 11 genes (sle10460 to sle10560) including a putative hopanoid biosynthesis gene cluster that is also present in S. scabies 87.22 (scab_12881 to scab_13001, excluding scab_12921 [41]) and S. coelicolor (scbo6759 to -6771, except scbo6761 [42]), as well as in many other streptomycetes. Between conS and the hopanoid biosynthesis gene cluster there are seven genes, four of which encode highly conserved proteins with homologues in S. coelicolor: Sle10450, Sle10430, Sle10420, and Sle10400 are homologous to Sco6772, Sco6773, Sco6774, and Sco6776, respectively. Two of the remaining genes (sle10440 and sle10410) encode highly conserved proteins that have no close homologues in S. coelicolor. The gene immediately upstream of conS, sle10390, encodes a hypothetical protein without clear homologues in the databases; since the 3′ end of the coding sequence of sle10390 overlaps with conS and there is a 720-bp intergenic region between sle10390 and the gene immediately upstream, sle10400, it is conceivable that sle10390 is part of the chaxamycin biosynthesis gene cluster, and thus this gene was assigned as the left limit of the cluster and named conM (Fig. 2 and Table 4).

The chaxamycin biosynthesis gene cluster of S. leeuwenhoekii, as defined above, spans a region of 80.2 kb, from nt 1210347 to 1290550 of the published genome sequence (19), and contains 27 genes. The core PKS and AHBA biosynthesis genes show very
similar organization, and the encoded proteins high sequence identity to the rifamycin biosynthesis gene cluster of *A. mediterranei* S699 (GenBank accession no.AF040570.3) (see Fig. S6 in the supplemental material). Consequently, gene names and predicted functions were assigned based on those defined in the rifamycin gene cluster (Fig. 2 and Table 4).

**Heterologous expression of the chaxamycin biosynthesis gene cluster.** One clone, pIJ12853, from a PAC library of *S. leuwenhoekii* genomic DNA was identified by PCR and confirmed by sequencing to contain the proposed chaxamycin biosynthesis gene cluster (see Materials and Methods). pIJ12853 was mobilized into *S. coelicolor* M1152, resulting in integration of the gene cluster at the chromosomal phiC31 attB site, yielding strain M1650.

M1650 was cultivated in R3 liquid medium, and culture supernatants and extracts of mycelial samples harvested at different times were analyzed by LC-MS, in parallel with chaxamycin standards and samples from *S. leuwenhoekii* grown in modified ISP2 liquid medium. The expected molecular ions for chaxamycins A to D (compounds 1 to 4) were readily found in samples from M1650 and with retention times (6.1, 5.9, 5.7, and 6.4 min, respectively) similar to those for the purified standards and the samples from *S. leuwenhoekii* (Fig. 3). In addition, the MS/MS fragmentation patterns of the ions detected for compounds 1, 3, and 4 in the M1650 samples matched those detected from the natural producer, while those for compounds 1 and 3 also matched the fragmentation patterns obtained from the standards; production of compound 2 in M1650 was very low and did not provide clear fragmentation data (Fig. 3B). These results demonstrated that the proposed gene cluster does indeed encode chaxamycin biosynthesis.

**Deletion of the AHBA synthase gene (cxmK) and chemical complementation.** AHBA is the starter unit for the synthesis of the polyketide backbone of chaxamycin. *cxmK* (Table 4) encodes a putative AHBA synthase that catalyzes both the transamination of UDP-3-keto-d-glucose at an early stage of the pathway and the final aromatization of amino-DHS to AHBA (13). Only one copy of this gene was found in the genome sequence of *S. leuwenhoekii*, directly downstream of the PKS genes. *cxmK* was deleted and replaced with the kanamycin resistance gene neo by homologous recombination (see Materials and Methods and Fig. S7 in the sup-

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**FIG 3** Heterologous production of chaxamycins A to D in *S. coelicolor* M1152. Extracted ion chromatogram (EIC) and MS/MS fragmentation patterns are shown for each chaxamycin species detected. (A) *S. coelicolor* M1650 (M1152 containing pIJ12853); (B) *S. coelicolor* M1152 (negative control); (C) chaxamycin A, B, and C standards; (D) *S. leuwenhoekii*. Chaxamycin A (compound 1), m/z (M – H)− 638.29; chaxamycin B (compound 2), m/z (M – H)− 622.29; chaxamycin C (compound 3), m/z (M – H)− 654.29; chaxamycin D (compound 4), m/z (M – H)− 682.29.
FIG 4 Chemical complementation of S. leeuwenhoekii M1653 (ΔcxmK::neo) with 3-amino-5-hydroxybenzoic acid (AHBA). Extracted ion chromatogram (EIC) and MS/MS fragmentation patterns are shown for each chaxamycin species detected. (A) S. leeuwenhoekii M1653; (B) S. leeuwenhoekii M1653 supplemented with 0.36 mM AHBA. Chaxamycin A (compound 1), m/z (M − H)⁻ 638.29; chaxamycin B (compound 2), m/z (M − H)⁻ 622.29; chaxamycin C (compound 3), m/z (M − H)⁻ 654.29; chaxamycin D (compound 4), m/z (M − H)⁻ 682.29.

pllemental material) to generate S. leeuwenhoekii M1653 (ΔcxmK::neo).

M1653 was cultivated in parallel with wild-type S. leeuwenhoe- kii in chaxamycin production medium; LC-MS analysis of samples of culture supernatants harvested at different times after inoculation showed that biosynthesis of all of the chaxamycin species had been abolished in the ΔcxmK mutant M1653 (Fig. 4A). To confirm that the lack of production was due solely to the deletion of cxmK, we attempted to complement the mutant by expression in trans of a wild-type copy of cxmK cloned in the expression vector pJJ10257. The resulting plasmid, pJJ12852, was introduced into M1653 but failed to restore chaxamycin production, presumably reflecting a polar effect of the insertion of neo on the expression of other downstream AHBA biosynthesis genes. We then succeeded in complementing the mutant chemically, restoring chaxamycin production (Fig. 4B) by supplementing the culture medium with commercially available AHBA; after testing several concentrations and times of addition, optimal results were obtained when adding AHBA to 0.36 mM at 24 h after inoculation, presumably coinciding with production of the chaxamycin biosynthetic pathway. When adding AHBA to 0.36 mM at 24 h after inoculation, optimal results were observed when adding AHBA to 0.36 mM at 24 h after inoculation, presumably coinciding with production of the chaxamycin biosynthetic pathway. When adding AHBA to 0.36 mM at 24 h after inoculation, optimal results were obtained when adding AHBA to 0.36 mM at 24 h after inoculation, presumably coinciding with production of the chaxamycin biosynthesis enzymes and minimizing AHBA degradation and/or catabolism. These feeding experiments confirmed a role for cxmK in AHBA synthesis and provided further evidence that the identified gene cluster does indeed encode chaxamycin biosynthesis. It also suggests that mutasynthesis, and the feeding of AHBA analogues, may be a viable approach for the generation of unnatural chaxa- mycin variants.

DISCUSSION

Functions of the genes present in the chaxamycin biosynthesis gene cluster and a proposed biosynthetic pathway. We have unequivocally identified, cloned, genetically manipulated, and heterologously expressed the chaxamycin biosynthesis gene cluster from S. leeuwenhoekii. The chemical structures of the chaxamycins and the rifamycins share many similarities, and therefore it is not surprising that the biosynthesis gene clusters are also very similar in organization. Based on knowledge of rifamycin biosynthesis, we can now propose a biosynthetic pathway for chaxamycin.

The chaxamycin biosynthesis gene cluster (Fig. 2 and Table 4) contains homologues (cxmG to -N and cxmI) of eight of the genes involved in AHBA biosynthesis in the rifamycin gene cluster of A. mediterranei S699 (rifG to -N and rifI; like rifJ, cxmI is not clustered with the other genes involved in AHBA synthesis). They show the same organization as the rif genes, and their products show high levels of amino acid sequence identity (68 to 86%). Therefore, we propose that biosynthesis of the chaxamycin PKS starting unit (AHBA) occurs as in rifamycin biosynthesis in A. mediterranei S699 (see Fig. S8 in the supplemental material) (13, 14, 43).

In addition to CxmG to -N and CxmJ, a transketolase is also required for AHBA biosynthesis to catalyze the conversion of 3-amino-3-deoxy-D-fructose-6-phosphate to imino-erythrose-4-phosphate (14). Indeed, in Salinispora arenicola CNS-205, deletion of sare_1272 and sare_1273, which encode a two-subunit transketolase, from the shared rifamycin-saliniketal biosynthesis gene cluster abolished the production of both compounds (44). While the proposed chaxamycin biosynthesis gene cluster does not appear to possess such a function, there are two genes (sle09700 and sle09710) located 47.4 kb to the right of the right, contained within the insert of pJJ12853, that encode homologues of transketolase subunits from A. mediterranei (encoded by rifI5A and rifI5B, respectively [45]) that might fulfill this role. Other lines of evidence suggest that such a function can be provided by genes outside the biosynthesis gene cluster. In A. mediterranei, deletion of rifI5A and rifI5B (45), homologues of sare_1272 and sare_1273, respectively, led to the accumulation of rifamycins SV and S, indicating that AHBA was still being produced in the mutant strain. In addition, the heterologous production of AHBA in S. coelicolor after cloning the AHBA genes rifG to -N and rifI suggests that the transketolase step can be performed by an alternative enzyme encoded elsewhere in the genome (14, 43).
Cxml is homologous to RifI, an aminooquinate (amino-Q) dehydratase that catalyzes the conversion of amino-DHQ to amino-Q, with no obvious role in AHBA biosynthesis. Deletion of rifI in *A. mediterranei* led to the accumulation of 20 to 25% more AHBA, and a salvage function has been proposed for the enzyme (43, 46).

The type I PKS responsible for the assembly of the chaxamycin polyketide backbone, is encoded by five genes, *cxmA* to -E. Bioinformatic analysis of their protein sequences was performed with antiSMASH (47), which uses two methods to predict the substrate specificity of the loading and AT domains (48, 49). antiSMASH predicted a loading module with specificity for AHBA, followed by a module selective for methylmalonyl-CoA, one for malonyl-CoA, six more for methylmalonyl-CoA, and finally two more for malonyl-CoA, which is entirely consistent and colinear with the reactions required to produce chaxamycin A (compound 1) (Fig. 2 and Table 4).

The loading module of *CxmA* contains an adenylation domain with 79% sequence identity to the adenylation domain of RifA in *A. mediterranei*, which catalyzes the reaction between a benzoate molecule and ATP to produce an activated aryl-AMP species (50, 51). The *CxmA* adenylation domain contains the AMP-binding and ATPase motifs, suggesting that it could function in a similar fashion to its homologue from *A. mediterranei* (see Fig. S9 in the supplemental material).

The chemical structure of compound 1 suggests that the dehydratase (DH) domains from modules 1, 2, 5, 6, 7, and 8 should be inactive. Active DH domains possess two conserved motifs, DAALH and the conserved histidine residue in the motif HXXX GXXXXP (although the DH domain from module 9 has proline replaced by alanine). DH domains from modules 1 and 2, predicted to be nonfunctional, do not have any of the conserved motifs. DH domains from modules 5 and 8, also predicted to be nonfunctional, are shorter than the rest of the DH domains and do not contain the motif D(A/V)/(V)A/(A/L)(Q/H); thus, despite having the conserved H residue in the active site, they lack the catalytic dyad. Surprisingly, the DH domains of modules 6 and 7 contain both active-site motifs, and we presume that there are other mutations that result in their inactivity (see Fig. S10 in the supplemental material).

The KR domain of module 3 lacks the active-site tyrosine residue (54) and the NADPH-binding motifs (55); the KR domain present in the loading module is shorter and lacks both of sequence features mentioned above. Therefore, these KR domains are predicted to be inactive, again in agreement with the structure of chaxamycin. The remaining KR domains have the tyrosine residue and the conserved motifs for NADPH binding (see Fig. S11 in the supplemental material) and are predicted to be active, consistent with the structure of the polyketide (Fig. 2).

The chaxamycin PKS shows a high level of similarity to that involved in rifamycin biosynthesis in *A. mediterranei* S699 but contains two additional nonfunctional domains (an extra KR domain in the loading module and an extra DH domain in module 2) as well as, unusually, an apparently functional additional acyl carrier protein (ACP) domain in module 6. As for the rifamycin PKS, the chaxamycin PKS lacks a type I thioesterase domain at the end of module 10, and a putative N-acetyltransferase/amide synthase is encoded by *cxmF*, the likely orthologue of rifF, and is similarly located just downstream of the last PKS gene (*cxmE*) (Fig. 2). Therefore, we propose that CxmF catalyzes the formation of the internal amide bond between AHBA and the last extender molecule of the polyketide chain to close the macrolactam ring, as in rifamycin biosynthesis (56), thus releasing the PKS product, which we have called prochaxamycin (Fig. 2).

In rifamycin biosynthesis, a type II thioesterase, RifR, is thought to function in the release of aberrant intermediates from the PKS, thus restoring functionality for a new cycle (57). However, it is not essential, and the PKS itself may be capable of such activity (58). Interestingly, we identified a nonfunctional rif*F* homologue located between *cxmY* and *cxm24* that contained three frameshift mutations (the frameshift-corrected amino acid sequence shows end-to-end similarity and 64% identity to RifR); possible errors in the PacBio-Illumina assembled DNA sequence were ruled out by PCR amplification and Sanger sequencing (see Materials and Methods). Whether an alternative type II thioesterase encoded outside the *cxm* gene cluster plays an editing role for the chaxamycin polyketide synthase remains to be determined.

We identified eight genes encoding putative tailoring activities homologous to those found in the rifamycin gene cluster of *A. mediterranei*. In rifamycin biosynthesis, the naphthalene ring is formed during the extension of the polyketide backbone by the enzyme 3-(3-hydroxyphenyl)propionate hydroxylase encoded by *orf19* (59); a homologous gene, *cxm19*, is also present in the chaxamycin gene cluster, and the encoded protein is predicted to catalyze the formation of the naphthalene ring of chaxamycin.

A putative methyltransferase is encoded by *cxm24*, but it shows little similarity (25% identity over 87 residues of the 355-amino-acid Cxm24) to the C-27 O-methyltransferase (Rif-Orf14) involved in rifamycin biosynthesis. This is the only putative methyltransferase encoded by the chaxamycin gene cluster, and although Pfam predicts that it is an O-methyltransferase, we speculate that it is responsible for C methylation of the C-3 of AHBA in the naphthalene ring.

An O-acetyltransferase encoded by *orf20* catalyzes the O-acetylation of C-25 during rifamycin biosynthesis (60). *Cxm20*, encoding a homologous protein with 63% identity to Rif-Orf20, is thus proposed to catalyze the O-acetylation of the hydroxyl group of C-25 in chaxamycin D (9). In rifamycin biosynthesis by *A. mediterranei* S699, a cytochrome P450 monooxygenase (CYP) encoded by *orf5* is required for the formation of the keto furan moiety present in rifamycins SV and B but not in rifamycin W; a very similar CYP encoded by *orf13* seems to be nonessential for rifamycin B biosynthesis (59). We identified only one gene, *cxm23*, which encodes a putative CYP homologous to rifamycin Orf5 or Orf13 (68% and 72% amino acid sequence identity, respectively). This is similar to the rifamycin gene cluster from *S. arenicola* CNS-205, where only one encoded protein, Sare1259, is homologous to Orf5 and Orf13 from *A. mediterranei* S699 (with higher identity to Orf13 [79%] than to Orf5 [71%]). Sare1259 performs several oxidation reactions at different positions on the molecule (44) and is 77% identical to Cxm23; therefore, we propose that Cxm23 participates in the formation of the hydroxy furan ring present in chaxamycin D but not in chaxamycin A, B, or C (Fig. 2).

Transcriptional regulation. Two genes encode putative tran-
scriptional regulatory proteins, ccmY and ccmZ. Analysis of CxmY with Pfam (61) revealed a carboxy-terminal helix-turn-helix (HTH) DNA-binding domain (PF00196) typical of the NarF/FixJ family of response regulators (RRs) (62); the rest of the amino acid sequence shows no similarity to that protein family, but BLASTp analysis (21) revealed many homologues of putative transcriptional regulators in other actinomycines, including Rif-Orf36 of the rifamycin gene cluster of A. mediterranei S699, with which it shows end-to-end similarity and 43% amino acid sequence identity. Similar analysis of CxmZ revealed a carboxy-terminal DNA-binding domain of the winged HTH family (Trans_reg_C family, PF00486) typical of the Ompr/Fbo family of RRs (62), and an amino-terminal RR receiver domain (Response_reg family, PF00072). CxmZ lacks the first of a pair of asparagines at positions 13 (serine) and 14 (aspartate) that determine the structure of the phosphorylation pocket in typical RRs, and it is thus unlikely to be phosphorylated by a cognate histidine sensory kinase (no such protein is encoded in the ccm gene cluster), supporting its classification as an atypical RR (see Fig. S12 in the supplemental material). CxmZ shows 41% amino acid sequence identity to JxR, an atypical response regulator involved in the regulation of jadomycin biosynthesis in Streptomyces venezuelae (GenBank accession no. AAB36584.2). Although contiguous, ccmY and ccmZ are transcribed in the opposite direction with an intergenic region of over 600 bp that presumably contains the promoter sequences of both genes.

Transport and immunity. Surprisingly we did not find any gene(s) encoding a putative transport system within, or in close proximity to, the ccm gene cluster. The rifamycin biosynthesis gene cluster of A. mediterranei S699 encodes an efflux transporter (encoded by rjP) whose transcription is likely repressed by a TetR-like transcriptional regulator encoded by the gene immediately downstream (rjQ) (46, 63). Homologues of these two genes are absent from the chaxamycin gene cluster and from the rifamycin/saliniketal gene cluster of S. arenicola CNS-205, which also does not seem to encode a transport system (44). However, two RjP homologues are encoded elsewhere in the S. leeuwenhoekii genome: Sle37020 and Sle08130 (62% and 60% amino acid sequence identity to RjP, respectively). Neither of the corresponding genes is present in $\text{P}1\text{I12853}$. S. coelicolor M145, from which strain M1152 was derived, also encodes a RjP homologue, Sco4024 (62% identity to RjP). Sco4024 is 78% identical to Sle37020 and 64% identical to Sle08130. All of these predicted proteins (as well as RjP) contain the major facilitator superfamily (MFS) domain and could be responsible for export of chaxamycins from their respective strains. A rjP homologue (sare_2597) also occurs in the genome of S. arenicola CNS-205, outside the rifamycin/saliniketal gene cluster.

Rifamycin binds to the β subunit (RpoB) of RNA polymerase, and in A. mediterranei a rifamycin-resistant form of the protein is encoded by an rpoB homologue located immediately downstream of rjP-orf36-orf37 (64). In S. leeuwenhoekii, rpoB (nt 3577583 and 3581068) lies 1.29 Mb away from the chaxamycin gene cluster (nt 1211049 to 1289829), and its product does not have the amino acid residues required for resistance to rifamycin at the expected positions, instead possessing the same amino acids as S. coelicolor M145 RpoB, which is sensitive to rifamycin (see Fig. S13 in the supplemental material). Thus, it seems unlikely that chaxamycin exerts its antibiotic activity by binding to RpoB.

This study has provided new insights into chaxamycin biosynthesis. We have succeeded in expressing the gene cluster heterologously, thus providing a convenient means for a more detailed study of the biosynthetic pathway and the enzymes involved. We have also shown that genetic manipulation of the natural producer is feasible, which could aid in the future engineering of the strain and the directed production of particular chaxamycin species.

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