Two Chimeric Chromosomes of *Streptomyces coelicolor* A3(2) Generated by Single Crossover of the Wild-Type Chromosome and Linear Plasmid SCP1

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*Streptomyces* species are gram-positive filamentous soil bacteria with a high G+C composition (70 to 74%) and are well known for the production of a large number of secondary metabolites. Moreover, this genus is unusual among bacteria because its members have a linear chromosome (1, 7, 38, 39, 52) and a linear plasmid(s) (16, 30, 33). *Streptomyces* linear chromosomes and plasmids have the same principal structural features: terminal inverted repeats (TIRs) are present at both ends and a terminal protein is covalently linked to the 5′ ends (1, 55).

*Streptomyces coelicolor* A3(2), the genetically best studied *Streptomyces* species, contains a linear chromosome (45) as well as two sex plasmids, the linear plasmid SCP1 (18, 50) and the circular plasmid SCP2 (3, 4, 47). Recently, genome projects have been completed for all three of these replicons, namely, the *S. coelicolor* A3(2) chromosome (8,668 kb) (2), SCP1 (356 kb) (3), and SCP2*, a derivative of SCP2 (31 kb) (15).

Considerable attention has been given to SCP1 because of its interaction with the host chromosome. Hopwood and colleagues isolated variants carrying a hybrid SCP1-chromosome structure (18, 19, 21). Namely, SCP1 is integrated into the central region or other regions of the *S. coelicolor* A3(2) chromosome. Free SCP1-prime plasmids containing a certain chromosomal DNA stretch were also found. In matings with SCP1-free partners, SCP1-integrated strains showed either a unidirectional or bidirectional gradient of transfer of genetic markers with respect to the SCP1 integration site.

Previous studies reported the integrated structures of SCP1 in the normal fertility (NF) strain 2612 and the NF-like strain A634 (14, 54). These strains show similar types of bidirectional DNA transfer, although the directions of SCP1 integration are opposite and the deletion sizes at each end are totally different. Therefore, the molecular reason for directional DNA transfer is still unknown. On the other hand, genetic studies of the *cysB* donor strain 1984 and the *cysD* donor strain 2106 suggested that they contained an SCP1-prime plasmid, either SCP1′-*cysB* or SCP1′-*cysD* (20). A preliminary physical analysis revealed that SCP1′-*cysB* and SCP1′-*cysD* are giant linear plasmids, of 550 and 1,700 kb, respectively (35). However, their structural details and interaction with the host chromosome have not been clarified.

For this paper, we precisely analyzed the structures of SCP1′-*cysD* and the chromosome of *S. coelicolor* A3(2) strain 2106 and revealed the interaction between SCP1 and the wild-type chromosome in these linear DNA elements. Based on our results, we discuss the universality and function of TIRs of *Streptomyces* linear replicons and the origin and evolution of linear chromosomes.

**MATERIALS AND METHODS**

**Bacterial strains, cosmids, plasmids, and media.** *S. coelicolor* A3(2) strain 1147 is the wild-type strain (17), and strains M145, M138, and 2106 are derivatives of strain 1147. The genotype and plasmid status of each strain are as follows (29): 1147, wild type/SCP1′ SCP2′; M145, prototroph/SCP1′ SCP2′; M138, argA1 proA1 cysD/SCP1′ SCP2′; and 2106, cysD donor pheA1/SCP1′SCP2′. *Escherichia coli* SURE2 was used as a host for a cosmids library, and *E. coli* XL1-Blue was used for cloning and nucleotide sequencing. A cosmids library of strain 2106 was constructed by a previously described method (44, 45) using the vector Supercoos1 (10) and a Gigapack III Gold packaging kit (Stratagene, La Jolla, Calif.). pSCP201 contains the 3.9-kb SpeI end fragment of SCP1 (34), and pSU22.21 contains the 1.3-kb BamHI end fragment of the M145 chromosome (22). YEME medium (29) was used for liquid cultures of *S. coelicolor* A3(2) strains. MYM agar, which was used for methylenomycin sensitivity assays, contains 0.4% maltose, 0.4% yeast extract, 1.0% malt extract, and 2.0% agar.

**DNA preparation and pulsed-field gel electrophoresis.** Total DNAs of *S. coelicolor* A3(2) strains were prepared as described by Kieser et al. (29). DNA gel samples were prepared by the mycelium method (31, 38), and SCP1′-*cysD* and SCP1 were separated by contour-clamped homogeneous electric fields (CHEF) (9). CHEF assays were performed with 0.5× Tris-borate-EDTA buffer and 1.0% Tris-borate-EDTA buffer.
agaro gels at 14°C. E. coli strains were cultured in Luria-Bertani medium, and cosmids and plasmid DNAs were isolated as described by Sambrook et al. (46).

Southern hybridization experiments. DNA fragments were separated by CHEF or conventional agarose gel electrophoresis and then transferred to nylon membrane filters as described by Kieser et al. (29). For colony hybridization, cosmids clones were inoculated directly onto nylon membrane filters, cultured on Luria-Bertani ampicillin plates, and treated as described by Sambrook et al. (46).

For the preparation of large hybridization probes, such as the 146-kb AseI fragment of SCP1 and the AseI-B fragment of the M145 chromosome (1,450 kb), a CHEF assay was performed with low-melting-point agarose, and excised bands were melted at 70°C and extracted with phenol and then phenol:chloroform (1:1). DNA fragments were labeled with digoxigenin-11-dUTP (Roche Diagnostics, Mannheim, Germany), and hybridization and detection were carried out as described by the supplier.

Telomere cloning and nucleotide sequencing. The telomere fragments of SCP1-cysD were separated first by CHEF gel electrophoresis and then by conventional agarose gel electrophoresis and were cloned into pUC19 by force cloning. For force cloning (12, 34, 37), the telomere fragment, which should still retain a small peptide covalently bound to the 5’ end after a protease treatment, was directly ligated to the vector and subjected to transformation. Nucleotide sequence analyses were performed by the dideoxy termination method with a Dye Terminator cycle sequencing kit (Amersham Pharmacia Biotech, Uppsala, Sweden) and an ABI-373S sequencer (PE Biosystems, Foster City, Calif.). The Genetyx-Mac 10.1 program (Software Development, Tokyo, Japan) was used for analyses of sequence data.

PCR. Primers for PCR amplification of the junction fragment of SCP1-cysD were designed based on the junction sequence of the 7.2-Mb chromosome, and their sequences were as follows: primer A, 5’-GGTGGGGTG-3’ (complementary to nt 176767 to 176744 of the M145 chromosome sequence [accession no. AL590463]); and primer B, 5’-AAGAGGCGGAAATCCC GTGTTGGGTG-3’ (complementary to nt 176767 to 176744 of the M145 chromosome sequence [accession no. AL023861]). Lowercase letters indicate the bases that were introduced to create recognition sites for HindIII and EcoRI, respectively. PCR products were done with the total DNA of strain 2106 and with the wild-type chromosome DNA as template (data not shown), while both the 146-kb SCP1 fragment and the M145 AseI B fragment hybridized to this fragment (Fig. 1A, panels 2 and 3). These results indicated that the 1,560-kb AseI B fragment was a hybrid of the 146-kb fragment of SCP1 and the AseI B fragment of the wild-type chromosome. Based on the total sizes of the AseI fragments of strain 2106 together with genome project data for the M145 chromosome (2), the chromosomal size of strain 2106 was calculated to be 7.2 Mb. Thus, it was suggested that in strain 2106, SCP1-cysD and the 7.2-Mb chromosome were generated by a single crossover between the 8,668-kb wild-type chromosome and the 356-kb linear plasmid SCP1 (Fig. 1B; the direction of SCP1 is shown opposite to that reported in reference 32).

Nucleotide sequences of SCP1-chromosome junctions. To confirm that there was a single crossover between SCP1 and the wild-type chromosome in strain 2106, we tried to clone the SCP1-chromosome junctions in SCP1-cysD and the 7.2-Mb chromosome. First, a cosmid library was constructed for the total DNA of strain 2106 by use of a partial digest with Sau3AI. Of 2,000 cosmids obtained, 83 clones were selected by positive hybridization to SCP1 DNA. An additional round of hybridization using the chromosomal AseI B fragment identified four candidate cosmids (6B3, 7G1, 8E8, and 11B2). Hybridization with SCP1-cysD and the 7.2-Mb chromosome revealed that these cosmids were not derived from SCP1-cysD but from the 7.2-Mb chromosome.

The four junction clones hybridized to both SCP1 cosmids 32 (44) and the chromosomal cosmid 3C8 (45), with the latter being located in the AseI B fragment. Restriction and hybridization analysis with these cosmids located the junction of the 7.2-Mb chromosome in a 1.1-kb BamHI fragment (pJB9-1) (Fig. 2A). This fragment was subcloned from cosmid 8E8 and then sequenced (Fig. 2B). The results, together with genome project data for SCP1 and the M145 chromosome, determined the SCP1-chromosome junction site in the 7.2-Mb chromosome and also defined another junction in SCP1-cysD. The deduced 1.0-kb DNA fragment containing the latter junction was amplified by a PCR with the two primers indicated in Fig. 2A (arrowheads) (see Materials and Methods for primer sequences), cloned into pUC19, and sequenced.

As shown in Fig. 2B, a sequence comparison of the two junctions and their corresponding regions in SCP1 (cosmid 32; accession no. AL590463) and the wild-type chromosome (cosmid 3C8; accession no. AL023861) revealed no homology between the two recombination regions. Thus, the two junctions were generated by nonhomologous recombination between SCP1 and the wild-type chromosome. During this event, a 55-bp DNA from SCP1 and a 15-bp DNA from the wild-type chromosome, shown with dashed lines in the figure, were deleted, and one T residue was inserted. The cross-over points were located on a putative helicase gene
(SCP1.136) in SCP1 and on an unknown gene (SCO6388) in the chromosome. Two fusion genes were generated by recombination: they were a mutated helicase gene in SCP1/H11032-cysD coding for an elongated N-terminal region and a hypothetical gene in the 7.2-Mb chromosome coding for a small protein of only 24 amino acids, which therefore may not function.

Nucleotide sequences of the telomeres of SCP1/H11032-cysD. Most Streptomyces linear chromosomes and plasmids that have been analyzed thus far have TIRs at both ends. In this respect, SCP1'-cysD and the 7.2-Mb chromosome were quite interesting, because they seemed to not have TIRs; rather, the telomere sequences of SCP1 (34) and the M145 chromosome (22) are totally different. When it was probed by the SCP1 end clone pSCP201 (34) and the chromosomal end clone pSUL221 (22), an NcoI digest of SCP1'-cysD showed hybridizing signals at the same positions as SCP1 (3.2 kb) and the M145 chromosome (1.25 kb) (Fig. 3; we used NcoI because BamHI gave terminal fragments of similar sizes [1.25 and 1.33 kb] for both SCP1 and the chromosome). In addition, the 7.2-Mb chromosome also showed the same two signals (Fig. 3).

To clone both ends of SCP1'-cysD, we digested it with SpeI and separated the fragments by CHEF electrophoresis. The 3.9- and 50-kb fragments, which contain the SCP1 (left) and chromosome (right) ends, respectively, were isolated. The

FIG. 1. Comparison of AseI fragments of the linear DNA elements in *S. coelicolor* A3(2) strains by CHEF assay and Southern hybridization (A) and generation model of SCP1'-cysD and the 7.2-Mb chromosome in strain 2106 (B). (A) (1) CHEF electrophoresis. CHEF electrophoresis was done at 150 V with 90-s pulses for 36 h. Fragment sizes were based on data from genome projects. (2 and 3) Southern hybridization. The 146-kb AseI fragment of SCP1 and the AseI B fragment of the M145 chromosome were used as probes for panels 2 and 3, respectively. (B) Generation model. Black and white bars indicate the SCP1 and chromosomal regions, respectively, with the former enlarged to twice its size. AseI recognition sites, fragment names, and TIRs are also included. The two recombination points are connected by a dashed line.
The former was cloned into pUC19 which had been digested with BamHI and HincII. The 50-kb SpeI fragment was further digested with BamHI, and the resultant 1.33-kb end fragment was cloned. The nucleotide sequences of the two right-end clones were identical to that of the 

\[ \ldots \text{S. coelicolor A3(2) chromosome telomere: } \ldots \text{TGTGGTACCCGCTCCGCGGG-3'} \]

On the other hand, the three left-end clones showed minor heterogeneity. Specifically, two clones contained four G residues at the 5’ end and one clone contained three G residues \[ \ldots \text{G(G)GGGCGGAGAGGCCTAACGGC. . .} \]. A similar heterogeneity (four to six G residues) was previously reported for the 5’ end of SCP1 \[ \text{SCP1-H11032} \]. It is not known whether these differences represent real heterogeneity in SCP1 \[ \text{SCP1-H11032} \] or if they were generated during the cloning process. In either case, the terminal sequences of the chimeric molecule are different and therefore are not repeats of each other.

**Failure of curing of SCP1’-cysD from strain 2106.** It was easily speculated that SCP1’-cysD is indispensable for the survival of strain 2106 because it contains a 1.6-Mb DNA fragment from the right end of the wild-type chromosome. To study this further, we tried to cure SCP1’-cysD from strain 2106 by the following three mutagenic treatments: incubation with acriflavin or ethidium bromide (26) and incubation at a high temperature (42°C). Since SCP1’-cysD carries the methylenomycin biosynthetic and resistance genes (6), mutants that lost SCP1’-cysD would become sensitive to methylenomycin. Thus, 500 colonies isolated after each mutagenic treatment were grown adjacent to strain M138, which carries SCP1 and produces methylenomycin. Eleven, 9, and 3 colonies of the acriflavin or ethidium bromide or high-temperature treated SCP1’-cysD strains were shown to be methylenomycin-sensitive. These sensitive colonies were used as probes for the detection of SCP1’-cysD in strain 2106 by Southern hybridization. The results indicated that SCP1’-cysD was not cured from strain 2106 by any of the treatments used.

*FIG. 2. Gene organization (A) and nucleotide sequences (B) around the junctions of SCP1’-cysD and the 7.2-Mb chromosome in strain 2106. (A) pJB9-1 contains the junction of the 7.2-Mb chromosome, and the two arrowheads indicate the primers for PCR amplification of the SCP1’-cysD junction. The names of open reading frames are according to genome projects for S. coelicolor A3(2) and SCP1. SCP1.136* codes for the mutated helicase in SCP1’-cysD. Af, AfII; Ba, BamHI; Cl, Clal; Sp, SphI; Xh, XhoI. (B) Nucleotide sequences of the corresponding regions of SCP1 and the M145 chromosome. It was revealed that a 55-bp SCP1 DNA and a 15-bp chromosomal DNA were deleted and a T residue was inserted during the recombination process.*

*FIG. 3. Southern hybridization analysis of the ends of SCP1’-cysD and the 7.2-Mb chromosome of strain 2106. SCP1, SCP1’-cysD, the M145 chromosome, and the 7.2-Mb chromosome were digested with NotI, separated by conventional agarose gel electrophoresis, and analyzed by Southern hybridization. pSCP201 (A) and pSUL221 (B) were used as probes for the SCP1 end and the chromosomal end, respectively.*
flavin, ethidium bromide, and high-temperature groups showed some growth inhibition. All of these colonies, plus 15 well-grown colonies randomly selected from each group, were further analyzed by CHEF electrophoresis for the presence of SCP1-\textsuperscript{-}cysD. All of the tested colonies were confirmed to still carry SCP1-\textsuperscript{-}cysD.

**DISCUSSION**

For this study, we analyzed the structures of the two linear DNA elements, SCP1-\textsuperscript{-}cysD and the 7.2-Mb chromosome, in *S. coelicolor* A3(2) strain 2106. Macrorestriction analysis and nucleotide sequencing revealed that both DNA elements were formed by a single crossover between nonhomologous regions of the wild-type chromosome and SCP1, with deletions of a 15-bp chromosomal DNA and a 55-bp SCP1 DNA. Therefore, the sizes of SCP1-\textsuperscript{-}cysD and the chromosome of strain 2106 were calculated to be 1,843 and 7,181 kb, respectively. The right and left telomeres of SCP1-\textsuperscript{-}cysD were also cloned and sequenced, and the results showed that the telomeres of the wild-type chromosome and SCP1 are conserved at each end. This may also be true for the 7.2-Mb chromosome, as its NcoI digest gave hybridizing signals at the same positions as the end fragments of the chromosome (1.25 kb) and SCP1 (3.2 kb). We were not able to cure SCP1-\textsuperscript{-}cysD by various mutagenic treatments, which indicated that SCP1-\textsuperscript{-}cysD is indispensable for survival. Thus, both the 7.2-Mb chromosome and SCP1-\textsuperscript{-}cysD are chimeric chromosomes, and we named them chromosomes I and II, respectively.

Strain 2106 transferred a cysD phenotype (cysD donor) in a mating with an SCP1-negative strain (19). Consequently, it was believed that the cysD gene is located on the giant linear plasmid SCP1-\textsuperscript{-}cysD. However, from our sequence data, the cysD gene was deduced to be present on the Asel B' fragment of the 7.2-Mb chromosome at a spot that is 356 kb from the junction. Therefore, the 1,843-kb linear DNA element should be called chromosome II instead of SCP1-\textsuperscript{-}cysD, and the gene-transferring property of strain 2106 should be analyzed physically based on the present results. Apart from this problem, the chimeric chromosomes of strain 2106 are noteworthy in two aspects, i.e., the universality and function of the TIRs of *Streptomyces* linear replicons and the origin and evolution of linear chromosomes, which are discussed below in detail.

Most of the linear chromosomes and plasmids isolated so far from *Streptomyces* have TIRs at both ends. The sizes of the TIRs of linear plasmids are quite different, ranging from 44 bp for SL2P in *S. lividans* (8) to 95 kb for pPZG101 in *S. rimosus* (13). The sizes of *Streptomyces* linear chromosomes are also different, ranging from 168 bp for *S. avermitilis* (24) to about 6,500 kb for the end-to-end fused chromosome of the *S. ambofaciens* mutant NSA65 (53). On the other hand, Kalkus et al. (27) reported that plasmid pHG201 in *Rhodococcus opacus* has TIRs of only 3 bp, if they can be called TIRs. Pandza et al. (41) showed that recombination between plasmid pPZG101 and the chromosome of *S. rimosus* led to an exchange of their ends. Based on its restriction map, the generated hybrid plasmid pPZG103 seemed not to have TIRs. Recently, the linear plasmid SLP2 was shown to be a composite plasmid comprising the chromosome and a linear plasmid on each side (23). In addition, this study revealed that neither of the two chimeric chromosomes in strain 2106 has TIRs but that they contain an SCP1 telomere and a chromosomal telomere at each end.

*Streptomyces* linear chromosomes are unstable and frequently cause deletions at one or both ends (7, 36, 51). The deleted chromosomes subsequently display dynamic rearrangements such as circularization, arm replacement, and amplification. We proposed the following hypothesis for the function of TIRs in *Streptomyces* linear replicons (48). When one of the two telomeres is deleted inside the TIR region, recombinational DNA repair may function between the intact and deleted TIR sequences on the same chromosome and recover an intact telomere. This event always repairs telomere damage inside the TIR region. However, we cannot observe it because an identical TIR structure is reproduced. Thus, the TIRs may guarantee homologous sequences for recombination, and this is the reason that most of the *Streptomyces* linear replicons have TIRs at both ends.

When a deletion in one telomere extends beyond the TIR region, the intact TIR structure cannot be recovered in this way. However, if nonallelic similar sequences are present on the right and left arms, homologous recombination between them causes chromosomal arm replacement and reproduces a telomere at the end (11, 48). In this case, some chromosomal sequences are lost and new, longer TIRs are formed. This may be the reason that the sizes of TIRs vary greatly in *Streptomyces* strains and that homologous genes or insertion elements are frequently found at the inside ends of TIRs, and this possibility was pointed out previously (34). When both telomeres are deleted, the extreme ends cannot be recovered, and therefore the chromosome will be circularized by nonhomologous recombination to survive (25, 28). Long TIRs, which are formed by chromosomal arm replacement, also suffer telomere deletions. If a second recombination occurs at the deletion ends inside the long TIRs, a circular chromosome with a large palindromic structure is generated (49). Qin and Cohen (43) analyzed similar structural changes of pSLA2 derivatives and discussed the strategies of *Streptomyces* linear replicons after telomere damage.

Without TIRs at both ends, how can *Streptomyces* linear replicons recover an intact telomere after terminal deletion? Qin and Cohen (43) demonstrated that when a linear plasmid which contained a pSLA2 telomere at one end and a damaged chromosomal telomere at another end was introduced into *S. lividans* ZX7, the damaged telomere was repaired by intermolecular recombination between the plasmid and the chromosome. This result, together with the presence of linear replicons without TIRs, such as pHG210, pPZG103, SLP2, and the two chimeric chromosomes in strain 2106, suggests that TIRs are not essential for *Streptomyces* linear replicons under exceptional conditions. Specifically, if identical telomere sequences are present on two different linear replicons in the same cell, they may compensate for each other intermolecularly in recombinational DNA repair.

*Streptomyces* linear chromosomes contain two replication mechanisms, bidirectional replication from a centrally located replication origin (40) and terminal replication primed by a terminal protein (42). The replication origins of *Streptomyces* chromosomes are quite similar to those of typical bacterial circular chromosomes (5, 56), while their telomere sequences can make a secondary fold-back structure like those of adeno-
viruses and paroviruses (22, 42). Thus, it was proposed that Streptomyces linear chromosomes may have been generated in the evolutionary past by a single crossover between a bacteri- um-type circular chromosome and a linear plasmid or phage which contained TIRs at both ends and terminal proteins bound to the 5′ ends (7, 52). In addition, the results obtained in this study suggested another possible evolutionary process, i.e., that duplication and multiplication of a linear chromosome may have occurred in a similar way to the mechanism for the generation of chromosomes I and II in S. coelicolor A3(2) strain 2106.

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