Intraspecific Variability of the Terminal Inverted Repeats of the Linear Chromosome of *Streptomyces ambofaciens*

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The sequences of the terminal inverted repeats (TIRs) ending the linear chromosomal DNA of two *Streptomyces ambofaciens* strains, ATCC23877 and DSM40697 (198 kb and 213 kb, respectively), were determined from two sets of recombinant cosmids. Among the 215 coding DNA sequences (CDSs) predicted in the TIRs of strain DSM40697, 65 are absent in the TIRs of strain ATCC23877. Reciprocally, 45 of the 194 predicted CDSs are specific to the ATCC23877 strain. The strain-specific CDSs are located mainly at the terminal end of the TIRs. Indeed, although TIRs appear almost identical over 150 kb (99% nucleotide identity), large regions of DNA of 60 kb (DSM40697) and 48 kb (ATCC23877), mostly spanning the ends of the chromosome, are strain specific. These regions are rich in plasmid-associated genes, including genes encoding putative conjugal transfer functions. The strain-specific regions also share a G+C content (68%) lower than that of the rest of the genome (from 71% to 73%), a percentage that is more typical of *Streptomyces* plasmids and mobile elements. These data suggest that exchanges of replicon extremities have occurred, thereby contributing to the terminal variability observed at the intraspecific level. In addition, the terminal regions include many mobile genetic element-related genes, pseudogenes, and genes related to adaptation. The results give insight into the mechanisms of evolution of the TIRs: integration of new information and/or loss of DNA fragments and subsequent homogenization of the two chromosomal extremities.

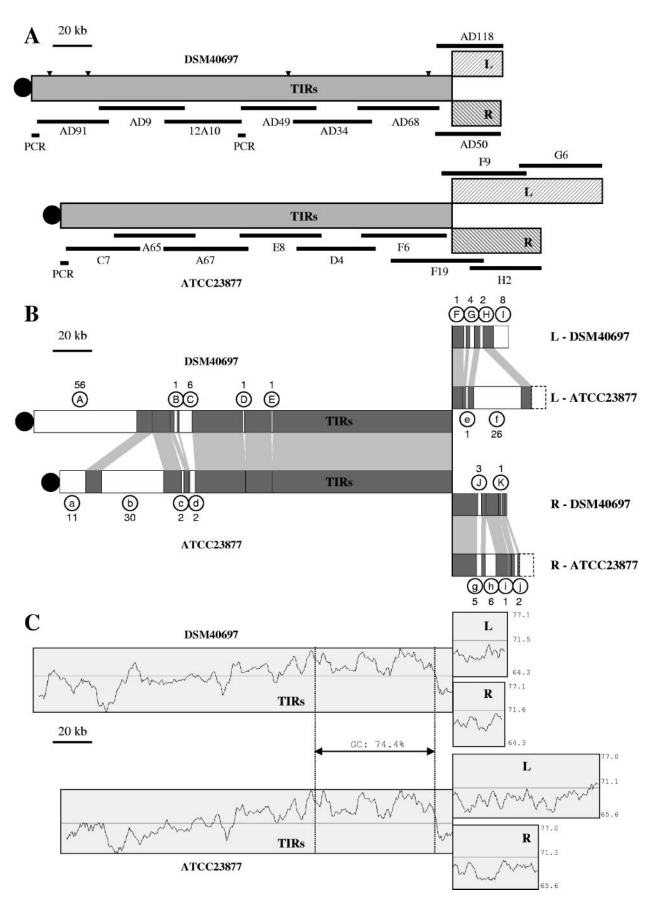
Streptomyces chromosomal DNA is linear and is among the largest described for bacteria, typically 8 to 10 Mb (6, 19). Streptomyces linear replicons (chromosomes and plasmids) share an invertronic structure including the presence of terminal inverted repeat sequences (TIRs) ended by bacterial telomeres covalently linked to terminal proteins (35). The lengths and sequences of the TIRs are extremely variable, and their sizes are not correlated to that of the replicon. The terminal duplications can be as large as several hundreds of kilobases, e.g., in the chromosome of *Streptomyces coelicolor* M600 (~1 Mb) (43), or can be restricted to the telomeric palindromes, e.g., in the chromosome of Streptomyces avermitilis (167 bp) (19). Two Streptomyces annotated genomes have been released so far: that for S. coelicolor A3(2), with 7,825 predicted coding DNA sequences (CDSs) (6), and that for Streptomyces avermitilis MA-4680, with 7,577 putative CDSs (19). Comparison of the two genomes revealed a common general organization with a conserved central region of about 5 Mb and terminal regions (or "arms") carrying mainly nonessential variable genes (6, 19). The terminal regions appear poorly conserved at

the level of gene content and organization, which contrasts with the strong synteny observed for the central region. The fact that the terminal regions are dispensable for vegetative growth in laboratory conditions was revealed earlier by the characterization of instability phenomena (25). In S. ambofaciens, up to 2.5 Mb located at the ends of the chromosome can be lost and is not essential for vegetative growth in laboratory growth conditions. In addition, large DNA rearrangements, such as duplications, deletions, and amplifications, frequently occur in the subtelomeric regions. One of the most spectacular rearrangements affects the size of the TIRs, which can vary from 5 kb to 1.4 Mb in spontaneous mutant strains (46), while the wild-type strain DSM40697 harbors 210-kb TIRs (26). This variation implies nonreciprocal translocations of chromosomal extremities that in some cases result from homologous recombination between duplicated genes (14). Another phenomenon triggering TIR variation implies exchanges of replicon extremities between plasmids and chromosomes. This was demonstrated for a strain of S. coelicolor in which chimeric chromosomes can be generated by crossover of the wild-type chromosome and the linear plasmid SCP1 (47). This phenomenon was also shown for Streptomyces rimosus by interaction between plasmid pZG101 and the chromosome (30) and strongly suggested by the analysis of the terminal structure of the Streptomyces lividans chromosome, which could result from partial integration of the linear plasmid SLP2 (17).

In order to gain further insight into the mechanisms of chromosomal end diversification in *Streptomyces*, we analyzed

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the variability of the TIRs of two isolates belonging to the *S. ambofaciens* species. It was previously shown by cross-hybridization experiments that they could be distinguished by the terminal regions (13). These two independent soil isolates, ATCC23877 (32) and DSM40697 (18), were assigned to the same species according to classical morphological and physiological traits, e.g., antibiotic production. These two isolates indeed share the same antibiotic synthesis profile, both producing the three known antibiotic compounds spiramycin, congocidin, and alpomycin (31, 32). In contrast, the closely related species *S. coelicolor* (1.1% divergence of 16S rDNA sequence from that of *S. ambofaciens* ATCC23877) shows a completely different secondary metabolite profile.

This phenetic classification was supported by more-recent molecular analyses. First, at the whole-genome scale, pulsed-field gel electrophoresis analysis, which is a highly sensitive approach to distinguishing and identifying bacterial isolates, showed that the two strains differ only slightly, whereas the *S. coelicolor* chromosome diverges much more (26). Further, mapping experiments with large chromosomal DNA fragments and linking clones showed that most of them were common to both strains, leading to the conclusion that the two chromosomes show a colinear organization (26). An additional piece of evidence is the presence of a recent gene duplication affecting a sigma factor-encoding gene (*has* gene). This duplication is common to the two *S. ambofaciens* isolates, while a single copy of the *has* gene is present in the two complete genome sequences of *S. coelicolor* and *S. avermitilis* (34).

Furthermore, the most convincing evidence for their close relationships comes from the analysis of their 16S-23S internal transcribed spacer sequences, whose evolution is the most effective (among the *rm* operon) to infer close phylogenetic relationships. The six internal transcribed spacer regions were isolated from the two strains, and their sequences were compared to those of *S. coelicolor* (45). While the two isolates share identical sequences, these sequences differ from that of *S. coelicolor*, thereby showing the divergence of the strains. Although gene conversion between the *rm* loci could be identified, no crossover has occurred between them, maintaining the colinearity of the two chromosomes (45).

Altogether, these data show that the two isolates are closely related strains and that their assignment to the same species is supported by a multicriterion analysis. In this work, we undertook a detailed sequence analysis of the TIRs of these two *S. ambofaciens* strains to determine the strain-specific gene content and to gain a deeper insight into the mechanisms important for the evolution of chromosomal extremities.

MATERIALS AND METHODS

Sequencing. For each strain, ATCC23877 and DSM40697, a cosmid library was constructed from partially BamHI-digested *S. ambofaciens* genomic DNA cloned into the Supercos1 (Stratagene) vector. As the size of the *S. ambofaciens* TIRs greatly exceeds that of a fragment readily clonable into a cosmid vector, each copy of the TIRs cannot be isolated as a single recombinant molecule. Consequently, the sequences were obtained from a set of ordered recombinant cosmids (Fig. 1A). Note that for cosmids with insert sequences entirely corresponding to the TIRs, e.g., from cosmid C7 to F6 for strain ATCC23877, the chromosomal origin (i.e., from the right or left arm) of the recombinant cosmids cannot be deduced. Therefore, the TIR sequence is likely to consist of a chimera between the left and right repeats.

For sequencing, cosmids were mechanically fragmented and cloned with a BstXI adaptor into either pcDNA2.1 vector (Invitrogen) or pCNS (a derivate of pSU18 [4]). Ligation products were then introduced into Escherichia coli DH10B. It was not possible to obtain a cosmid clone containing the terminal fragment that includes the telomeres, and a PCR strategy was therefore used to amplify and sequence this region in both strains. Primers were designed from the cosmid C7 sequence (5'CACCCAGCGAGCCCAGCA3') for strain ATCC23877 and from the cosmid AD91 sequence (5'AGCTGCAACGGTGCGTTCTATTG GG3') for strain DSM40697, and a third primer was designed according to a consensus of the telomere sequences derived from several Streptomyces species (5'CGGAGCGGTACCACATCGCTG3'). PCR was performed using 50 ng of DNA, with 800 µM deoxynucleoside triphosphates, 2 units of LA Taq polymerase (Takara), 2.5% dimethyl sulfoxide, and 20 pmol of each primer in a 50-µl final volume. After a denaturation step (95°C, 5 min), 30 cycles of denaturation (95°C, 30 s), annealing (58°C, 30 s), and polymerization (68°C, 3 min) were used to amplify the terminal fragment.

The 10 cosmids of strain ATCC23877 were completely sequenced (mean coverage, 10×). For strain DSM40697 (mean coverage, 10×), 17 gaps were remaining after the shotgun sequencing of the eight cosmids. PCR products were obtained for each gap and were cloned into pGEM-T Easy vector (Promega) for production of single-stranded DNA. For each gap, sequencing reactions were performed on all available templates, i.e., PCR product, double-stranded recombinant pGEM-T Easy vector, and the single-stranded DNA. In addition, sequencing reactions were carried out using two different reagents, CEQ Dye terminator (Beckman) and BigDye Terminator (Applied Biosystems). Four gaps remained, all localized in intergenic regions, suggesting they probably resulted from the formation of intrastrand secondary structures (e.g., terminators). Gap 1 (between DSMT0010 and DSMT0011) and gap 2 (between DSMT0032 and DSMT0033) are included in a strain-specific region (cosmid AD91) (Fig. 1A), and they are estimated to be less than 50 bp in length. Gap 3 (between DSMT0146 and DSMT00147) and gap 4 (between DSMT0204 and DSMT0205) belong to regions highly conserved between the two strains, and their sizes can be estimated as less than 10 bp and 200 bp, respectively, by comparison to the sequence of strain ATCC23877.

Annotation. The gene finder Glimmer 2.10 (11) was used for CDS prediction, with a minimum size of 40 codons arbitrarily chosen as the threshold. Results were then refined by RBSfinder (39). The Basic Local Alignment Search Tool (BLAST 2.2.6) was used to find similarities (1), and the Interpro package was used to describe protein domains (48). CDSs were assigned a functional category where their best cluster of orthologous groups (COG) homologue is classified (40). Then, BLASTX translations were realized for each intergenic region in order to detect initially unpredicted CDSs and pseudogenes. While comparing the predicted protein sequences with BLASTP, proteins sharing more than 30% identity over at least 80% of the length of the query sequence were considered to be homologous.

FIG. 1. (A) Schematic representation of the *S. ambofaciens* ATCC23877 and DSM40697 TIRs and adjacent regions (L, left arm; R, right arm) and of the cosmids and PCR products used for the sequencing. Gap positions are represented by triangles (see Materials and Methods). (B) Comparison of the TIR sequences and flanking regions of *S. ambofaciens* ATCC23877 and DSM40697. Strain-specific regions are represented by white rectangles, whereas conserved regions are represented in gray. The strain-specific regions are named as follows: from A to K for strain DSM40697 and from a to j for strain ATCC23877, and the number of genes (including pseudogenes) carried by each specific region is indicated. Since the regions sequenced outside of the TIRs are larger for strain ATCC23877, sequences which cannot be compared because they were not sequenced in the second strain are represented by dashed rectangles. The terminal protein, covalently bound to the telomere, is represented by a black circle. (C) G+C content of TIRs and adjacent regions for the two strains (displayed with a 5,000-bp window size). Minimum, maximum, and mean (represented by a straight line) values of G+C percentages are indicated at the right side. These values are calculated for the whole contigs including the TIRs and the left or right adjacent region.

Strain	Size of TIR (bp)	G+C content (%)	No. of predicted CDSs (no. of pseudogenes)	No. of proteins with assigned function (%)	No. conserved with proteins of unknown function (%)	No. of orphans (%)
ATCC23877	197,936	71.8	194 (3)	125 (65)	49 (25)	20 (10)
DSM40697	212,655	71.9	215 (10)	135 (63)	54 (25)	26 (12)

TABLE 1. General features of the TIR sequences of the S. ambofaciens strains ATCC23877 and DSM40697

For strain ATCC23877, duplicated genes in the TIRs were named SAM<u>T</u>nnnn, whereas those specific only to either the left or right arm were annotated as SAM<u>L</u>nnnn or SAM<u>R</u>nnnn, respectively (underlining indicates the type of specificity). A similar principle was adopted for strain DSM40697, with the prefix "DSM" used instead of "SAM." Sequences of left and right contigs for the two strains and their corresponding annotations are available through the SAMDB web server at http://www.weblgm.scbiol.ambofaciens.uhp-nancy.fr/.

Nucleotide sequence accession numbers. The sequences were deposited in EMBL under the following accession numbers: AJ937740 (ATCC23877 left TIR), AJ937741 (ATCC23877 right TIR), AM279694 (DSM40697 left TIR), and AM279695 (DSM40697 right TIR).

RESULTS

Intraspecific variability at the chromosomal ends. The large TIR sequences of two *S. ambofaciens* strains were determined using sets of recombinant cosmids spanning the terminal regions of the chromosome (Fig. 1A). Chimeric sequences of 197,936 bp and 212,655 bp (see Materials and Methods) were produced for strains ATCC23877 and DSM40697, respectively (see Table 1 for general features).

Considering a single genome, the 100% nucleotide identity of the two TIR copies was supported by two lines of evidence. First, hybridization of DNA probes corresponding to the TIRs onto genomic DNA did not reveal any polymorphism (26). Second, no mismatch was found within the overlaps of sequenced cosmids, despite the fact that they can originate from either of the chromosomal arms.

The ends of the TIRs will be defined as the first nucleotide of divergence (noted as "internal boundary of the TIRs" in Fig. 2A). Consequently, the terminal duplication does not end at exactly the same nucleotide position in the two strains. However, this situation results from point mutations and small insertions/deletions causing minor differences between the arms in both strains in the flanking small region (over about 5 kb). These regions show near identity and contain two CDSs, a probable cytochrome P450 (DSMT0215, SAML/R0195) and a homologue of afsA (DSML/R0216, SAML/R0196), separated by a large intergenic region that includes stretches of short repeated C/A-rich motifs (Fig. 2A). Another stretch of C/A-rich motifs is located in the 3' part of afsA, and the variation in the number of motifs results in differences between the four afsA copies (two copies in each strain). Consequently, the afsA coding sequences have different sizes (three different alleles for four copies). In addition, these open reading frames (ORFs) might be pseudogenes, since only the 5' end (about 200 codons) shows similarity (between 38% and 46%) with the afsA-like gene of Streptomyces rochei (28).

The two arm sequences diverge totally at the same nucleotide proximal to *afsA* in the two strains. Thus, the two strains share the same ancestral boundaries of TIRs (Fig. 2A).

Thus, 194 CDSs (from SAMT0001 to SAMT0194) belong to the TIRs of strain ATCC23877. This last ORF is similar to that

encoding a truncated transposase. In strain DSM40697, the end of the sequences assumed to be identical occurs within the long intergenic region separating DSMT0215 (putative cytochrome P450) and DSML/R0216 (*afsA* homologues) (Fig. 2A).

When the TIRs of the two strains are compared, two regions can be distinguished: a terminal strain-specific region and a conserved region with a more internal location. In total, the syntenic regions extend over about 150 kb and include 149 genes sharing an average 99% nucleotide identity (Fig. 1B).

Within the whole TIRs, seven strain-specific segments resulting from DNA rearrangements such as insertions, deletions, and/or gene replacements involving from 1 to 56 CDSs can be delimited (Fig. 1B). For strain DSM40697, five specific regions (A to E) represent 62 kb and contain 65 CDSs (including probable pseudogenes). Region A on its own includes 56 of the 65 strain-specific CDSs and represents a quarter of the TIR size. For strain ATCC23877, the specific loci are scattered into four regions (a to d) spanning 49 kb and including 45 strain-specific CDSs. Again, 41 of these 45 specific genes are clustered into two loci, one of which constitutes the chromosomal end (region a, 14 kb).

Although the chromosomal ends are strain specific, highly similar telomere sequences were found in the two strains (96.1% nucleotide identity over 180 bp) (Fig. 3). The seven typical palindromes predicted to form the secondary structures found in other known *Streptomyces* telomeres are present in both strains (7, 17) (Fig. 3).

Outside the TIRs, intraspecific variability can also be noticed. Within the areas sequenced, at least four and five DNA rearrangements have occurred in the left and right arms, respectively (Fig. 1B; also see below).

Altogether, these results suggest that, despite their variability in gene content, the current TIRs of the two *S. ambofaciens* strains appear to derive from a single ancestral event of terminal duplication, since they share the same ancestral internal boundary.

Low G+C content. Despite the differences in gene content observed between the two strains, profiles of G+C percentages are quite similar (Fig. 1C). For each strain, a decrease in G+C content characterized the strain-specific extremities as well as the sequences outside of the TIRs. Thus, the terminal 50 kb of the ATCC23877 and DSM40697 chromosomes show G+C contents of 68.8% and 69.2%, respectively. Similar values are observed for the regions sequenced outside of the TIRs (69.1% in strain ATCC23877 and 68.9% in strain DSM40697). In contrast, G+C contents of *Streptomyces* chromosomes are 72.1% for *S. coelicolor* and 70.7% for *S. avermitilis*. In fact, the TIRs of *S. ambofaciens* seem to be present in regions of even lower G+C content, but the presence of a 61-kb cluster (which includes the alpomycin cluster [31]) showing a G+C content of

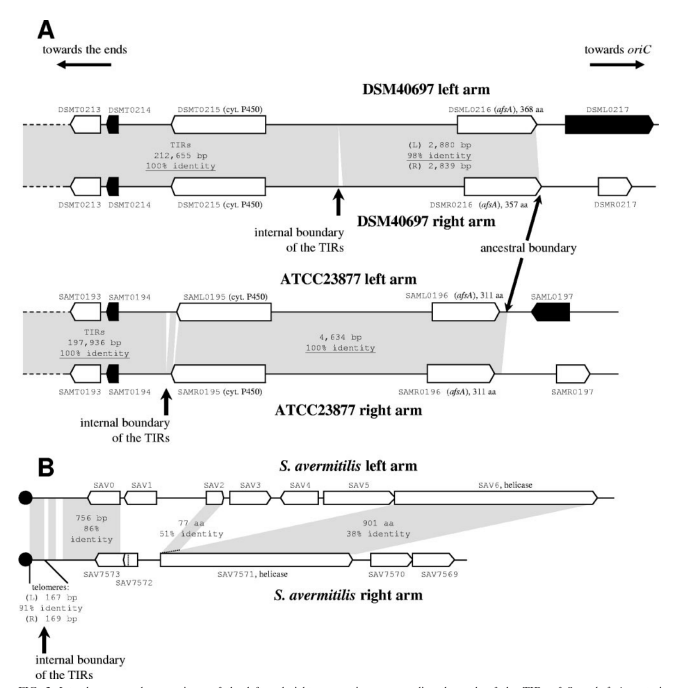


FIG. 2. Intrachromosomal comparisons of the left and right arm regions surrounding the ends of the TIRs of *S. ambofaciens* strains ATCC23877 and DSM40697 (A) and *S. avermitilis* (B). Duplicated regions between both arms are shaded in gray, and their sizes and percentages of identity are indicated. Black arrows represent ORFs similar to genes encoding transposases (or truncated transposases), and the terminal proteins are represented by the black circles. The SAV0 putative CDS was not predicted in reference 19 and was added in this work. cyt., cytochrome. aa, amino acid.

74.4% skews the data by causing a local increase in G+C content (Fig. 1C).

The low G+C content of the chromosomal extremities is reminiscent of that of *Streptomyces* plasmids (SLP2, 68.4% [17]; pSV2, 69.7% [38]; SCP1, 69.0% [5]; and SAP1, 69.2% [19] [see "Plasmid-associated genes" below]). The lower G+C content observed in the regions outside of the TIRs is related to a remarkable abundance of insertion sequences

(ISs) and related genes in the strain-specific regions (found in regions F, G, H, and K in strain DSM40697 and in regions f, h, and j in strain ATCC23877) (Fig. 1B; also see "Mobile genetic element-related genes" below). The lower G+C content of mobile genetic elements in bacteria has been discussed previously (33), and their presence, together with an observed low G+C content, suggests acquisition by horizontal gene transfer.

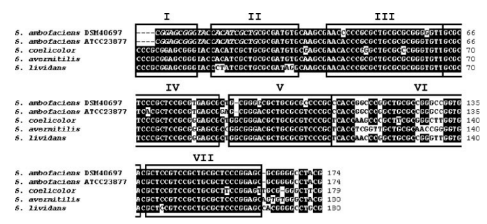


FIG. 3. Sequence alignment of the telomeres of the two *S. ambofaciens* strains with the chromosomal telomeres of *S. coelicolor* A3(2), *S. avermitilis* MA-4680, and *S. lividans* ZX7. The palindromes are labeled, and the primer sequence used for the amplification of the telomeres (see Materials and Methods) is represented in italic. This PCR strategy explains the missing nucleotides at the end of the *S. ambofaciens* telomeres. The boxes labeled I to VII indicate the positions of palindromes starting from the end of the chromosomal DNA. Numbers to the right of the sequences indicate the cumulative length for each of the aligned sequences.

Plasmid-associated genes. A large proportion of the strainspecific CDSs included in the TIRs may have a plasmid origin (see Table 2). This situation is particularly striking for strain DSM40697, for which 16 of the 56 CDSs located in the strainspecific region A show best similarity with plasmid-associated genes. For half of them, the level of identity is particularly high, i.e., more than 80% amino acid identity. Five gene products share best similarity with proteins encoded by linear plasmid SAP1 from S. avermitilis (19) and eight with linear plasmid SCP1 from S. coelicolor (5). Interestingly, one gene fragment (DSMT0048) is similar to tpg from S. lividans linear plasmid SLP2, which encodes the terminal protein involved in the replication of the telomeres (17). Some of the best hits were also with plasmids from other Actinomycetales spp., i.e., pNF1 (circular) from Nocardia farcinica (20) and pREL1 (linear) from Rhodococcus erythropolis (37). Furthermore, syntenic clusters between these strain-specific regions and linear plasmids (Table 2) were found, which strongly supports the hypothesis of integration of plasmid DNA into the terminal regions (9, 42). For example, the DSMT0042-45 cluster is conserved with the cluster SCP1.199-202 from linear plasmid SCP1.

The same conclusion can be inferred from the analysis of region a from strain ATCC23877. First, the probable DNA helicase TtrA (SAMT0002) shows the best BLASTP hit (84% amino acid identity) with that of plasmid SLP2 of S. lividans. This helicase is implicated in the conjugal transfer of the SLP2 plasmid (8). The ttrA gene is present at the extremities of the chromosome of each S. ambofaciens strain, although it is located in the strain-specific regions. It should be pointed out that a ttrA homologue is present close to the telomeres in almost all Streptomyces replicons. In fact, TtrA proteins of the two S. ambofaciens strains, which share 45% identity, may have a different origin: TtrA from strain ATCC23877 shares 84% identity with TtrA from plasmid SLP2, whereas that of strain DSM40697, which is truncated, shares 83% identity with TtrA from the chromosome of S. avermitilis. In addition, the SAMT0010 protein is similar to the KilB protein from different Streptomyces plasmids, pRL2 (linear) and pIJ101 (circular), in

which it is implicated in conjugal transfer and intramycelial spread (36).

Mobile genetic element-related genes. Genomic islands often carry genes implicated in mobility, such as those encoding integrases, recombinases, and transposases (15). Thus, the presence of such genes in variable regions supports the idea of acquisition by horizontal gene transfer. In the conserved part of the TIRs, 149 pairs of orthologues between the two S. ambofaciens strains were predicted. Among them, three gene products showing similarity with transposases (DSMT0057/ SAMT0012, DSMT0058/SAMT0013, and DSMT0214/SAMT0194) and one with an integrase/recombinase (DSMT0060/SAMT0015) were annotated. For three of them, the best similarity is found with transposase (or integrase) from Frankia species (Actinomycetales). Two transposase ORFs are located in a conserved region of the TIRs, just at the borders of the strain-specific regions A and a. The third one is located at the internal boundary of the TIRs as described above. However, none of the transposase-encoding genes seem to constitute a functional IS, the transposase being either inactivated by frameshift mutations or truncated, with no detectable flanking inverted repeats. The terminal strain-specific region A (DSM40697) contains a truncated IS (DSMT0046) similar to an IS from Frankia sp. strain Cci3 and a phage integrase (DSMT0003), close to the telomeres, sharing no homology to Streptomyces but with homology to *Nocardioides* species (see Table 2 for more details). Reciprocally, in the strain ATCC23877, one truncated IS (SAMT0032) is present in strain-specific region b.

Interestingly, many transposase-encoding genes are found close to the ancestral boundary of the TIRs (four in strain DSM40697 and three in strain ATCC23877), and this is a common feature of *Streptomyces* replicons. In *S. coelicolor* A3(2), an IS constitutes the ends of the chromosomal TIRs (6), while in plasmid SCP1, Tn5714 is located 3 kb outside the left TIR and IS466 is located at the end of the right one (5).

Given the close relationship between *S. ambofaciens* strains, it is even possible to spot recent IS- or transposon-mediated rearrangements. Indeed, outside of the TIRs, a putative com-

TABLE 2. Strain-specific genes predicted in the TIRs of S. ambofaciens strains DSM40697 and ATCC23877

Strain	Specific region	Gene	Product ^a	Identity (%) ^b	Overlap (%) ^c	Gene name	Plasmid	Organism
DSM40697	A	DSMT0001						
		DSMT0002	1 5					
		DSMT0003		35	93	NocaDRAFT_4522		Nocardioides sp. strain JS614
		DSMT0004						
		DSMT0005	Putative serine/threonine protein kinase					
		DSMT0006						
		DSMT0007						
		DSMT0008		83	88	SAV7571		Streptomyces avermitilis
		DSMT0009		64	98	SAP1 90 (lig)	SAP1	Streptomyces avermitilis
			Putative integral membrane transport protein	72	100	SCO6809		Streptomyces coelicolor
		DSMT0011	Putative secreted protein	75	100	SCO6811		Streptomyces coelicolor
			Putative FAD-dependent oxidoreductase	58	98	pREL1_0108	pREL1	Rhodococcus erythropolis
		DSMT0013	Putative transcriptional regulator	95	96	pFQ25.11		Streptomyces sp. strain F2
			Putative phosphinothricin	77	99	pFQ25.10		Streptomyces sp. strain F2
		DSMT0015	N-acetyltransferase Unknown					
		DSMT0016	Conserved hypothetical protein	85	99	pnf1840	pNF1	Nocardia farcinica
			Putative membrane protein	76	100	SCO3280		Streptomyces coelicolor
			Putative glycosyl hydrolase, BNR repeat	36	98	ArthDRAFT_2101		Arthrobacter sp. strain FB24
		DSMT0019	Putative lipoprotein	72	100	SCO4458		Streptomyces coelicolor
		DSMT0020	Putative membrane protein	60	96	SCO4459		Streptomyces coelicolor
			Putative monooxygenase	78	94	SCO6838		Streptomyces coelicolor
			Putative arsenic resistance membrane transport protein	84	99	SCO6837		Streptomyces coelicolor
			Putative transcriptional regulator	82	100	SCO3699		Streptomyces coelicolor
			Putative arsenate reductase	87	95	SCO6835		Streptomyces coelicolor
		DSMT0025		79	99	SCO6834		Streptomyces coelicolor
		DSMT0026		25	75	Tfu_2935		Thermobifida fusca
		DSMT0027						
		DSMT0028		87	100	SCP1.257	SCP1	Streptomyces coelicolor
		DSMT0029 DSMT0030	Putative RNA polymerase sigma	36 37	83 97	SAP1_88 SAP1_87 (sig)	SAP1 SAP1	Streptomyces avermitilis Streptomyces avermitilis
		DSMT0021	factor Conserved hypothetical protein	35	93	SAP1 86	SAP1	Streptomyces avermitilis
			Putative secreted protein	84	100	SCP1.323c	SCP1	Streptomyces coelicolor
			Putative secreted protein	89	100	SCP1.261c	SCP1	Streptomyces coelicolor
		DSMT0033		84	94	SCP1.262	SCP1	Streptomyces coelicolor
		DSMT0034		80	95	SAV7571	5011	Streptomyces avermitilis
			Conserved hypothetical protein	63	95	ShewDRAFT_1466		Shewanella sp. strain PV-4
		DSMT0037	DUF1099					
			Conserved hypothetical protein (pseudogene)	70	99	SCO0085		Streptomyces coelicolor
		DSMT0039	Putative ATP-dependent DNA ligase	59	100	SAP1_90 (lig)	SAP1	Streptomyces avermitilis
			Conserved hypothetical protein	71	100	SCO0048		Streptomyces coelicolor
		DSMT0041	Unknown					•
			Conserved hypothetical protein	29	95	SCP1.202	SCP1	Streptomyces coelicolor
			Conserved hypothetical protein	84	100	SCP1.201	SCP1	Streptomyces coelicolor
		DSMT0044	Putative secreted protein	94	100	SCP1.200c	SCP1	Streptomyces coelicolor
		DSMT0045	Putative secreted esterase	93	100	SCP1.199c	SCP1	Streptomyces coelicolor
		DSMT0046		55	71	Francci3_3385		Frankia sp. strain CcI3
		DSMT0047		55	93	nfa29650		Nocardia farcinica
		DSMT0048	Tpg protein fragment (pseudogene)	75	97	tpgSLP2	SLP2	Streptomyces lividans
		DSMT0049	regulator (pseudogene)	82	97	SCO3804		Streptomyces coelicolor
		DSMT0050 DSMT0051	Conserved hypothetical protein Putative nourseothricin acetyltransferase	73 69	48 100	SCO3803 nat1		Streptomyces coelicolor Streptomyces noursei
		DSMT0052		30	93	Francci3 1866		Frankia sp. strain CcI3
		DSMT0053 DSMT0054	Conserved hypothetical protein Unknown	47	76	Francci3_1863		Frankia sp. strain CcI3
		DSMT0055	Putative peptidase	38	98	SCO3610		Streptomyces coelicolor
		DSMT0056	Putative major facilitator superfamily	29	93	NocaDRAFT 1725		Nocardioides sp. strain JS614
	В	DSMT0083	Putative 3-oxoacyl-ACP synthase III	40	100	ArthDRAFT 2448		Arthrobacter sp. strain FB24
	C	DSMT0085	Putative ABC transport system ATP- binding protein	64	93	SCO0121		Streptomyces coelicolor
		DSMT0086		44	93	SCO0120		Streptomyces coelicolor
		DSMT0087	Conserved hypothetical protein	46	76	Tfu_1509		Thermobifida fusca
		DSMT0088	Conserved hypothetical protein	31	98	SCO0123		Streptomyces coelicolor
		DSMT0089	Putative polyprenyl synthetase	61	98	SCO0568		Streptomyces coelicolor

Continued on following page

TABLE 2—Continued

Strain	Specific region	Gene	Product ^a	Identity $(\%)^b$	Overlap (%) ^c	Gene name	Plasmid	Organism
		DSMT0090	Putative geranylgeranyl diphosphate synthase	48	100	ggdps		Streptomyces sp. strain KO-3988
	D E	DSMT0118 DSMT0134	Conserved hypothetical protein	84	79	SCP1.218c	SCP1	Streptomyces coelicolor
ATCC23877	a	SAMT0001	Conserved hypothetical protein	62	92	SAV7573		Streptomyces avermitilis
		SAMT0002 SAMT0003	Putative helicase Unknown	84	100	ttrA	SLP2	Streptomyces lividans
		SAMT0004	Conserved hypothetical protein	42	100	pFRL1.57	pFRL1	Streptomyces sp. strain FR1
		SAMT0005	Unknown	64	59	pFRL1.57	pFRL1	Streptomyces sp. strain FR1
		SAMT0006	Putative NTP pyrophosphohydrolase	39	67	PFL 4894	_	Pseudomonas fluorescens
		SAMT0007	Conserved hypothetical protein	52	95	nfa38470		Nocardia farcinica
		SAMT0008	Putative glyoxalase	64	96	nfa38460		Nocardia farcinica
		SAMT0009	Putative ferredoxin NADPH reductase	55	97	nfa38450		Nocardia farcinica
		SAMT0010	KilB-like protein, role in intramycelial spread	48	95	pRL2.23	pRL2	Streptomyces sp. strain 44414
	b	SAMT0011 SAMT0023		50	100	SAV2967		Streptomyces avermitilis
		SAMT0024	Putative hydrolase	53	90	Adeh 0548		Anaeromyxobacter dehalogenans
		SAMT0025	Conserved hypothetical protein	38	100	SCO7248		Streptomyces coelicolor
		SAMT0026 SAMT0027	Putative phosphotransferase Unknown	54	91	ard		Streptomyces avermitilis
		SAMT0028	Putative secreted protein	47	80	SCO0072		Streptomyces coelicolor
		SAMT0029	Putative secreted protein	61	89	SCO0072		Streptomyces coelicolor
		SAMT0030 SAMT0031	Unknown Unknown	30	67	Francci3_0808		Frankia sp. strain CcI3
		SAMT0032	Putative truncated transposase	71	80	orf118	pSLA2-L	Streptomyces rochei
		SAMT0033	Putative secreted protein	58	97	SCO0072	r	Streptomyces coelicolor
		SAMT0034	Putative secreted protein	86	100	SCO0072		Streptomyces coelicolor
		SAMT0035	Conserved hypothetical protein	60	100	SCO0073		Streptomyces coelicolor
		SAMT0036	Putative SAM-dependent methyltransferase	67	98	SCO2653		Streptomyces coelicolor
		SAMT0037		82	100	SCO0031		Streptomyces coelicolor
			Putative transmembrane restriction endonuclease	87	100	SCO7763		Streptomyces coelicolor
		SAMT0039	Unknown					
		SAMT0040	Putative stress response protein	77	100	SCO3763		Streptomyces coelicolor
		SAMT0041	Unknown	54	48	Francci3 1126		Frankia sp. strain CcI3
		SAMT0042	Putative membrane protein	37	100	SAV3190		Streptomyces avermitilis
		SAMT0043	Unknown					* *
		SAMT0044	Putative regulator	39	81	SAV1103		Streptomyces avermitilis
		SAMT0045	Putative anti-sigma factor antagonist (pseudogene)	42	81	SCO3692		Streptomyces coelicolor
		SAMT0046		71	96	prpC3		Streptomyces avermitilis
		SAMT0047	Putative hydrolase	79	100	SAV923		Streptomyces avermitilis
		SAMT0048	Putative urease beta/gamma subunit	50	94	DR A0319		Deinococcus radiodurans
		SAMT0049	Putative urease alpha subunit	65	98	SCO1234		Streptomyces coelicolor
		SAMT0050	Putative <i>ureF</i> -like urease accessory protein	76	100	ureF		Streptomyces avermitilis
		SAMT0051	Putative <i>ureG</i> -like urease accessory protein	75	98	ureG		Streptomyces avermitilis
		SAMT0052	Putative <i>ureD</i> -like urease accessory protein	54	81	SCO1231		Streptomyces coelicolor
	c	SAMT0065	Putative truncated transposase	41	100	SAV18		Streptomyces avermitilis
	-		Putative haloacid dehalogenase	89	99	SAV737		Streptomyces avermitilis
	d	SAMT0071	Putative transcriptional regulator	41	96			Saccharopolyspora erythraea
	-		Putative esterase	38	100	SCO4392		Streptomyces coelicolor

^a FAD, flavin adenine dinucleotide; BNR, bacterial neuraminidase repeat; ACP, acyl carrier protein; NTP, nucleoside triphosphate.

posite transposon that is absent from the DSM40697 strain was detected in strain ATCC23877 (SAMR0213 to SAMR0218 [complete region h in Fig. 1B]). This transposon (5.1 kb) consists of two almost identical IS elements (99% nucleotide identity) flanking four CDSs, the products of two of which are also related to IS transposases, plus two orphans. Interestingly, the flanking ISs do not share any homology with *Streptomyces* sequences but do share homology with a transposase from *Frankia* sp. strain EAN1pec (67% amino acid identity).

Coding density and pseudogenes. Acquisition of DNA by horizontal gene transfer and loss of useless genes are the main causes of intraspecific variability, and an equilibrium between these two phenomena leads to genomic flux that shapes bacterial genomes (24). The first step of gene loss is the creation of pseudogenes either by point mutations or by truncations.

In *S. ambofaciens*, the TIRs are characterized by a low coding density. When considering pseudogenes as noncoding DNA, the coding densities of the sequenced regions are

^b Results for the best BLASTP hit are summarized.

^c "Overlap" corresponds to the ratio between the length of the BLASTP alignment and the length of the query protein (for cases in which this ratio was >100%, 100% overlap was indicated).

77.1% and 75.7% for strains DSM40697 and ATCC23877, respectively. These values are not biased significantly by the presence of pseudogenes, since the coding densities are 80.1% (DSM40697) and 77.1% (ATCC23877) when pseudogenes are included as CDSs. These values contrast with values of 88.9% and 86.2% predicted for the complete chromosomes of *S. coelicolor* (6) and *S. avermitilis* (19), respectively. This is again a common trait of *Streptomyces* plasmids, such as SAP1 from *S. avermitilis*, for which the density falls to 79% (19).

Another striking feature of the terminal regions is the strong presence of pseudogenes, notably in the TIRs of strain DSM40697 (10, including two truncated transposases, representing 5% of the gene content). In contrast, pseudogenes represent less than 1% of the CDSs predicted in the whole genome of S. coelicolor (6). Eight out of the 10 pseudogenes are carried by the strain-specific region A. In addition to the truncation of the ttrA gene (DSMT0008; 219 amino acid residues out of 835 in S. avermitilis), two additional TtrA fragments are encoded in strain-specific regions A (DSMT0035; 44 residues [Table 2]) and G (DSML0224; 31 residues), outside of the TIRs. The first two fragments are homologous to different parts of the same gene (ttrA from S. avermitilis). However, the latter pseudogene is not related to them, suggesting acquisitions of extra copies by different integrations of parts of linear replicons. This hypothesis is further supported by the finding, in the same region A, of a gene fragment (DSMT0048; 99 bp) showing best identity with tpgC encoding the terminal protein of the linear plasmid SLP2 and of three gene fragments (DSMT0009, DSMT0038, and DSMT0039) similar to the lig gene encoding a ligase from the S. avermitilis linear plasmid SAP1 (SAP1 90).

In the regions sequenced outside of the TIRs, 10 additional probable pseudogenes were predicted for strain DSM40697, of which 6 belong to the strain-specific regions.

The number of pseudogenes is probably underestimated, and the low coding density could be a consequence of a high mutation rate. Altogether, these data support the hypothesis that the TIRs and the terminal regions of the genome constitute a hot spot for horizontal gene transfer events mediated by linear plasmids.

Horizontal transfer of accessory genes. Horizontal transfer mostly involves accessory genes that are able to confer a selective advantage to the recipient cell. Housekeeping genes are more recalcitrant to transfer (23). In *S. ambofaciens*, functions of many genes predicted for the strain-specific regions, such as resistance to toxic compounds, are related to adaptation to the environment. They are more particularly abundant in the terminal specific region A of strain DSM40697 (Fig. 1B). A good example is the five-gene cluster DSMT0021-25, which is highly similar (from 78% to 87% amino acid identity) to the *S. coelicolor* cluster SCO6838-34, which is implicated in the transport of and resistance to arsenate.

Antibiotic resistance genes are also present. For example, the DSMT0051 gene product shows best similarity (69%) with the *Streptomyces noursei* nourseothricin acetyltransferase, Nat1 (22). In the same way, a homologue of the Ard2-encoding gene (SAMT0026; 54% identity) from *Saccharothrix mutabilis* subsp. *capreolus* (*Actinomycetales*), which confers resistance to A201A antibiotic (3), is present in the ATCC23877 strain-specific chromosomal end.

The DSM40697 strain-specific regions B and C also carry functions related to adaptation. Indeed, the seven specific genes identified here (DSMT0083, DSMT0085-90) all have associations with secondary metabolism: DSMT0083 is a probable 3-oxoacyl-ACP synthase III-encoding gene having homologues in different Actinomycetales spp.; DSMT0085/86/88 are conserved, with three membrane and transport protein-encoding genes of S. coelicolor adjacent to the eicosapentaenoic acid cluster (SCO0124-0129); DSMT0087 shares similarity with a CDS of unknown function located in the fredericamycin biosynthesis gene cluster from Streptomyces griseus (44); and DSMT0089/90 are similar to genes encoding a putative polyprenyl synthase and a geranylgeranyl diphosphate synthase, respectively, from Streptomyces sp. strain KO-3988 (21). Interestingly, although these genes all seem to be implicated in secondary metabolism, they do not correspond to a whole conserved cluster but rather show similarity to genes from many different clusters among Actinomycetales. This could therefore be an example of a locus created by the association of genes derived from different pathways. Examples of transfer of secondary metabolism clusters have already been discussed in reference 27. The alp polyketide synthase cluster present in the TIRs of the two S. ambofaciens strains could also constitute an example of a chimeric cluster (31). While its left part is mostly similar to the kinamycin-biosynthetic cluster of S. murayamaensis (accession number AY228175), most of the right part shows high similarity to and the same genetic organization as a locus identified in the S. rochei linear plasmid pSLA2-L (28).

In addition, the variable region C is replaced by region d in strain ATCC23877 (Fig. 1B), in which one of the two specific CDSs identified encodes a probable transcriptional regulator sharing best identity with the OrfD regulator belonging to a cluster involved in the biosynthesis of a pigment in *Saccharopolyspora erythraea* (10).

However, a significant part of the strain-specific genes have no known function and are, in some cases, orphans.

Species specificity of the *S. ambofaciens* TIR conserved genes. The part of the TIRs conserved at the intraspecific level is highly variable at the interspecific level. No synteny can be observed the other *Streptomyces* with genomes. The regions conserved with *S. coelicolor* are limited to eight small syntenic clusters comprising two to seven CDSs (e.g., a urea degradation cluster). In contrast to what is found for *S. avermitilis*, only one cluster of four CDSs is syntenic.

Seventeen percent of the 149 pairs of orthologues between the two strains do not have any homologue in the NR database. In addition, among the 37 proteins showing best similarity with those from organisms other than *Streptomyces*, the majority (19/37) show highest similarity with those from other *Actinomycetales* genera (especially *Frankia*, *Arthrobacter*, *Nocardia*, *Nocardioides*, and *Kineococcus*). Many of these organisms are fellow soil-dwelling bacteria, which plausibly suggests lateral gene transfer events.

DISCUSSION

Acquisition of new functions by exchange of extremities between linear replicons. The *S. ambofaciens* TIRs contain a large proportion of strain- and species-specific genes as well as

sequences potentially involved in genome plasticity. Genetic organization of the TIRs is consistent with the idea that DNA rearrangements of endogenous sets of genes (duplication and translocation), integration of exogenous information, and/or deletions have been fixed during evolution from their common ancestor.

Several lines of evidence corroborate the acquisition of the terminal strain-specific regions by exchange of replicon extremities with linear plasmids: the presence of gene clusters homologous to plasmid-associated ones; the presence of genes implicated in conjugal transfer (*kilB* and *ttrA*); and their low G+C content (68.8% and 69.2%), which is characteristic of many *Streptomyces* plasmids. In addition, many similarities are found with genes carried by the extremities of the linear plasmids (*ttrA*, *tpgC*, *lig*, and *kilB*). Since the last telomere-proximal ORF conserved in the two strains is a truncated IS, it is plausible that exchange of extremities could have happened by homologous recombination involving two IS copies.

This comparison made by use of S. ambofaciens describes for the first time such exchanges at the origin of intraspecific variability in natural isolates. Insertion of DNA extremities of linear replicons may be a favored mechanism for gene acquisition in Streptomyces. Indeed, exchange of the terminal parts requires a single crossover event, the success of which would be guaranteed by the presence of the telomeres in the two replicons. In addition, the presence of a helicase-like gene (ttrA) closely associated with the telomere is a trait common to almost all Streptomyces linear replicons. However, it is truncated in strain DSM40697, and no deleterious effect could be assigned in laboratory growth conditions to the mutation of both copies of ttrA in S. lividans (17). The strong conservation of a gene in a variable region appears paradoxical, but it suggests a role for conferring some long-term advantage, e.g., conjugal transfer, as suggested by C.W. Chen in his "end-first" model (8). This model predicts that this probable helicase would be involved in conjugal transfer by acting on the DNA terminus which would correspond to an origin of transfer (7, 8).

Many of the functions predicted for the strain-specific regions are potentially related to adaptation, which consolidates the idea that horizontal transfer is at the origin of the *S. ambofaciens* terminal variability. In general, genes successfully transferred are responsible for adaptation to the environment. The presence in the DSM40697 strain chromosomal extremities of multiple resistance genes may have conferred advantages responsible for their maintenance in the bacterial population.

In addition to exchange of extremities, multiple events of insertions/deletions have occurred recently both inside and outside of the TIRs. For example, specific region b of the strain ATCC23877 chromosome could be the result of an integration event, but the absence of a detectable target site suggests either integration by illegitimate recombination or deletion of this locus in strain DSM40697. The specific regions C (DSM40697) and d (ATCC23877), which are two different variable regions located at the same locus, could be the consequence of a DNA exchange by double crossover or by different deletions at the same locus in the two strains.

All these multiple rearrangements are posterior to the formations of the TIRs that are themselves totally different from those of the very close *S. coelicolor* genome. Thus, genome

plasticity is extremely strong in the terminal regions, which could be privileged targets for environmental adaptation by loss, acquisition, and creation of new functions. They might be considered as a vector for genetic transfer and also as the "Swiss army knife" of the "boy scout" *Streptomyces* (6, 16).

Origin and evolution of the size of the TIRs. Although the presence of two telomeres is known to be essential for the maintenance of chromosome linearity (2), no role has so far been attributed to the TIRs. Indeed, some replicons, such as the *S. avermitilis* chromosome, have TIRs restricted to a part of the telomeres, while others carry very large TIRs (up to 1.4 Mb [46]). In addition, the duplication of the genes located in the TIRs does not appear to be a mechanism for gene regulation. Thus, there is no obvious difference between the transcriptional levels of the genes duplicated in the *S. coelicolor* M600 TIRs (1 Mb) and those for the same set of genes present in single copy in *S. coelicolor* M145 (22-kb TIRs [43]).

The formation of TIRs could be the consequence of terminal recombination induced either by dysfunction of a telomere or by formation of double-strand breaks (DSBs). Chromosomal rescue could then be achieved by recombination with a DNA fragment including a telomere similar to the endogenous one. This fragment may be a broken daughter chromatid (46) or a plasmidic or chromosomal extremity either endogenously present or resulting from horizontal gene transfer (e.g., conjugal transfer). DSB repair might also result from the breakinduced replication as described previously for *Saccharomyces cerevisiae* (29). Finally, circularization can also trigger chromosomal rescue (reviewed in reference 25).

Analysis of the S. ambofaciens TIR boundary reveals a preliminary step in TIR shortening (Fig. 2A). Thus, while the strict border of the TIRs is located at different positions in the two strains, the imperfect duplication includes the same genes. Point mutations and variability in tandem repeated motifs contribute to the decrease in length of the TIRs. In S. avermitilis, the terminal repetitions are restricted to the telomeres (19), which suggests the loss of ancestral TIRs in this species. Indeed, detailed analysis of the subtelomeric sequences reveals putative traces of ancestral TIRs (Fig. 2B). The last predicted ORF, SAV7573, is in fact imperfectly duplicated at the other end of the chromosome (Fig. 2B), and a putative CDS, which we tentatively called SAV0, can be detected between the telomere and SAV1. SAV0 (372 bp) and SAV7573 (411 bp) share 87% nucleotide identity over most of the CDS length (271/311 nucleotides). Furthermore, although only weakly similar (38%) amino acid identity), a duplicated helicase gene is also present at both ends of the chromosome (SAV6/SAV7571). SAV2 also shows similarity with the helicase-like gene (Fig. 2B) and could correspond to the duplication of the 5' end of SAV6 (69% nucleotide identity). These data suggest that once formed, the TIRs may accumulate point or insertional mutations that result in divergence which might progressively reduce and even prevent the frequent homogenization between TIRs and consequently accelerate the rate of divergence. The TIRs would then tend to degenerate by progressive shortening. The divergence of the duplicated genes present in the TIRs may result in pseudogene formation or leave functional coding sequences evolving toward new functions (34). Alternatively, the loss of TIRs in S. avermitilis could result from a single recombination

event between the ancestral chromosome and an exogenous DNA molecule.

The reduction in TIR size may be balanced by recombination events leading to expansion. Enlarged TIRs have been reported for mutants of S. griseus and S. ambofaciens (12, 41). In both cases, recombination events occurring between duplicated genes (each copy located on a different arm and in a divergent orientation) led to a dramatic increase of the TIR size (from 210 kb to 480 kb and 850 kb in S. ambofaciens and from 24 kb to 450 kb in S. griseus). For S. coelicolor, TIR length variation has been shown to be associated with strain lineage (43). The formation of large TIRs could be ascribed to homologous recombination between transposed IS110 copies. TIR shortening from 1.06 Mb to 22 kb was observed (43). It therefore seems possible that TIR formation corresponds to a byproduct of chromosomal rescue mechanisms. However, once formed, terminal duplications would provide an appropriate substrate for homologous recombination and thus for DNA repair of DSBs occurring in the terminal parts of the Streptomyces linear chromosome. The fact that the S. ambofaciens strain-specific regions are duplicated on both arms in a given chromosome strongly suggests a homogenization mechanism. In other words, these data show that DNA rearrangements occurring within a copy of TIRs can be followed by homogenization of both arms, probably by intrachromosomal recombination between duplicated sequences.

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