Characterization of the Self-Resistance Mechanism to Dityromycin in the *Streptomyces* Producer Strain

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ABSTRACT  
Dityromycin is a peptide antibiotic isolated from the culture broth of the soil microorganism *Streptomyces* sp. strain AM-2504. Recent structural studies have shown that dityromycin targets the ribosomal protein S12 in the 30S ribosomal subunit, inhibiting translocation. Herein, by using *in vitro* protein synthesis assays, we identified the resistance mechanism of the producer strain to the secondary metabolite dityromycin. The results show that the self-resistance mechanism of the *Streptomyces* sp. strain AM-2504 is due to a specific modification of the ribosome. In particular, two amino acid substitutions, located in a highly conserved region of the S12 protein corresponding to the binding site of the antibiotic, were found. These mutations cause a substantial loss of affinity of the dityromycin for the 30S ribosomal subunit, protecting the producer strain from the toxic effect of the antibiotic. In addition to providing a detailed description of the first mechanism of self-resistance based on a mutated ribosomal protein, this work demonstrates that the molecular determinants of the dityromycin resistance identified in *Streptomyces* can be transferred to *Escherichia coli* ribosomes, where they can trigger the same antibiotic resistance mechanism found in the producer strain.

IMPORTANCE  
The World Health Organization has identified antimicrobial resistance as a substantial threat to human health. Because of the emergence of pathogenic bacteria resistant to multiple antibiotics worldwide, there is a need to identify the mode of action of antibiotics and to unravel the basic mechanisms responsible for drug resistance. Antibiotic producers’ microorganisms can protect themselves from the toxic effect of the drug using different strategies; one of the most common involves the modification of the antibiotic’s target site. In this work, we report a detailed analysis of the molecular mechanism, based on protein modification, devised by the soil microorganism *Streptomyces* sp. strain AM-2504 to protect itself from the activity of the peptide antibiotic dityromycin. Furthermore, we demonstrate that this mechanism can be reproduced in *E. coli*, thereby eliciting antibiotic resistance in this human commensal bacterium.

KEYWORDS  
self-resistance, antibiotic, ribosomal protein S12, translation

The emergence of multidrug-resistant (MDR) bacteria is a growing problem that represents one of the top three threats to global public health worldwide (1–4). Moreover, it has been estimated that approximately 70% of infection-causing bacteria are already resistant to at least one antibiotic available in routine clinical practice. Therefore, all the efforts to provide a detailed molecular picture of a specific resistance mechanism can contribute positively to improve our understanding of antimicrobial resistance (AMR) and its dynamics.
Dityromycin is a cyclic decapeptide antibiotic (Fig. 1A) discovered as a secondary metabolite produced by *Streptomyces* sp. strain AM-2504 (5, 6). Its structure is almost identical to that of GE82832, a translocation inhibitor produced by *Streptosporangium* spp. (7–9). Both molecules interact with the same ribosomal site and display the same mechanism of action, even if they are produced by different microorganisms. The interaction of dityromycin with the ribosome is peculiar compared to other ribosomal inhibitors, since it interacts exclusively with the ribosomal protein S12 without a direct interaction with the 16S rRNA (Fig. 1B and C) (8, 9). Indeed, the binding of dityromycin to the ribosome is mediated by its interaction with five amino acids at the ribosomal protein S12, namely, arginine 30, valine 32, arginine 55, histidine 76, and valine 78 (Fig. 1C). Dityromycin inhibits protein synthesis by blocking the translocation of tRNA through the ribosome, while it has almost no effect on the accommodation of aminoacyl-tRNA into the P-site or the A-site, in the context of either the 30S ribosomal subunit or the 70S ribosome (7). Recent studies have shown that dityromycin blocks the EF-G-dependent translocation of peptidyl tRNA and mRNA without preventing the ribosomal binding of the elongation factor. In particular, dityromycin hampers the interaction between domain III of EF-G and protein S12, which is located near the decoding center of the ribosome, at the interface of two subunits (9, 10).

A strict requirement to ensure that the antibiotic producer cell protects itself when the active metabolites are produced, accumulated, and released is to possess a self-resistance mechanism(s) (11). Microorganisms can adopt several strategies to develop self-resistance, such as (i) chemical modification of the target, as in the case of rRNA methylation or mutation (12, 13); (ii) activation of efflux pumps (14, 15); (iii) sequestration of the natural product by proteins keeping the active compound in a bound and inactive state (16, 17); (iv) production of the antibiotic in a prodrug form, activated through selective cleavage by peptidases found in the periplasmic space, and subsequently released outside the cell (18, 19); and (v) oxidative inactivation of the matured prodrug following export (20).

Biochemical and genetic similarities demonstrate that such resistance mechanisms have prefigured those found subsequently in antibiotic-resistant pathogens, which have acquired specific genes or gene clusters originally developed by antibiotic producers (21).}

In this work, we have investigated the self-resistance mechanism developed by *Streptomyces* sp. strain AM-2504 toward the antibiotic activity of its natural product dityromycin. To prevent self-toxicity, this microorganism has evolved a strategy based on the production of a modified and self-resistant ribosomal protein variant. Our

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**FIG 1** Interaction of dityromycin with the 30S ribosomal subunit. (A) Chemical structure of the antibiotic dityromycin. (B) Three-dimensional (3D) structure of the 30S ribosomal subunit (gray) and localization of dityromycin (red) in contact with the ribosomal protein S12 (green). A, P, and E tRNA binding sites are indicated in blue, yellow, and violet, respectively. (C) Close-up of dityromycin (red) bound to S12 (green) missing the C-terminal domain. The amino acid residues of S12 involved in the interaction with the antibiotic are also indicated. (Modified from reference 9.)
The approach consisted in analyzing three aspects, namely, identification and biochemical characterization of the self-resistance mechanism, characterization of the amino acid residues of the ribosomal protein S12 involved in resistance by the heterologous expression of site-directed mutants in *Escherichia coli*, and a phylogenetic analysis of the dityromycin binding pocket in the ribosomal protein S12.

### RESULTS

**Identification of the mechanism conferring self-resistance.** To determine the self-resistance mechanism of *Streptomyces* sp. strain AM-2504 to dityromycin, we set up a simple experiment evaluating whether the resistance mechanism relied on possible modifications/secretion of the drug (prodrug or efflux pumps) or was based on mutations/methylations/modifications of the target.

A translational test programmed with poly(U) was used to compare cell extracts prepared from either *E. coli* (S30E.coli) or *Streptomyces* sp. strain AM-2504 (S30Str) in the presence or absence of the antibiotic. Dityromycin, an antibiotic targeting the translation elongation phase, completely inhibited protein synthesis with S30E.coli at concentrations ranging between 1 and 10 μM, whereas it did not show any effect on the poly(U) translation carried out with S30Str (Fig. 2A). The lack of inhibition with the S30Str extract clearly excluded prodrug or efflux pumps as a possible self-resistance mechanism.

To test whether this self-resistance mechanism was caused by protective mutations in the ribosomes or by an enzymatic activity able to modify/deactivate the antibiotic, 70S ribosomes were separated by ultracentrifugation from the post-ribosomal supernatant (S100) and the fractions were tested by crossed *in vitro* poly(U) translation assays. In particular, the effect of dityromycin on *Streptomyces* 70S ribosomes (70SStr) was assayed in the presence of *E. coli* S100 (S100E.coli). Vice versa, *Streptomyces* S100 extract (S100Str) was tested with *E. coli* 70S ribosomes (70SE.coli). The results of this experiment (Fig. 2B) show that the reaction mixtures consisting of S100Str plus 70S_E.coli and S100_E.coli plus 70S_Str were totally inhibited by dityromycin, whereas those formed by S100_E.coli plus 70S_Str and S100Str plus 70S_E.coli were not. These results strongly indicate that the modifications causing the self-resistance of the producer strain lie on the ribosome.

Since the target of dityromycin is the ribosomal protein S12 (9), 30S ribosomal subunits from both *E. coli* and *Streptomyces* sp. strain AM-2504 were purified. Increasing concentrations of these isolated particles were used in a pulldown assay conducted in the presence of a fixed concentration of the antibiotic. After ultracentrifugation at 300,000 × *g*, the supernatants were tested in a translational system based on *E. coli* cell extracts programmed with poly(U) to detect the residual presence of unbound dityro-
mycin (see Materials and Methods). As reported in Fig. 2C, the supernatant resulting from the pulldown of E. coli 30S subunits incubated with dityromycin did not interfere with poly(Phe) synthesis. Conversely, the supernatant obtained after centrifugation of Streptomyces 30S subunits incubated with dityromycin inhibited the reaction. This result indicates that dityromycin is not bound by the Streptomyces 30S subunits and is not pulled down during ultracentrifugation, even at high concentrations of ribosomes. In contrast, dityromycin stably interacts with the E. coli 30S subunits and is thus removed from the supernatant during the centrifugation step (Fig. 2C). In conclusion, the present results point to a self-resistance mechanism based on a mutation/modification residing in the small ribosomal subunit.

Sequencing of the S12 gene reveals three amino acid substitutions in the dityromycin binding site. As shown in the crystallographic structure of Fig. 1, dityromycin binds the 30S subunit by interacting exclusively with five critical positions of the ribosomal protein S12, namely, Arg30, Val32, Arg55, His76, and Val78 (Fig. 1C). To clarify the role played by S12 in reducing the binding affinity of dityromycin to the 30S\textsubscript{str} subunit (Fig. 2), we sequenced the \textit{rpsL} gene, encoding the S12 protein of Streptomyces sp. strain AM-2504. In addition, we sequenced the entire genome of the producer strain, thus obtaining its first annotated draft genome sequence (22). The comparison of \textit{Streptomyces} sp. strain AM-2504 genome with the sequences available in the NCBI genome database allowed us to verify that the strain named AM-2504 described in this study exhibits 98.6% nucleotide identity with \textit{Streptomyces kasugaensis}. Therefore, it is conceivable that \textit{Streptomyces} sp. strain AM-2504 can be classified as \textit{S. kasugaensis}.

DNA sequence data demonstrate that three out of five amino acids located in the dityromycin binding site of S12, namely, Val32Thr, Arg55Lys, and Val78Ile, are different from those found in the model organism \textit{E. coli} (Fig. 3). These substitutions are likely the molecular determinants of the self-resistance mechanism to the antibiotic, preventing dityromycin binding to the ribosome (Fig. 2C).

Expression of the S12 mutant proteins in \textit{E. coli} confers resistance to dityromycin. To verify whether these mutations are responsible for the self-resistance of the producer strain, single substitutions at Val32, Arg55, and Val78 were introduced into the \textit{E. coli rpsL} gene, and the mutant genes were provided in trans to the \textit{E. coli} BL21 strain grown in the presence of increasing concentrations of dityromycin. The experimental system was based on gene expression from the high-copy-number plasmid pET11a, which contained the \textit{rpsL} gene isolated from \textit{E. coli} MC4100 carrying the original mutation Lys42Arg, which confers resistance to the antibiotic streptomycin. As

FIG 3 Sequence of the \textit{rpsL} gene of \textit{Streptomyces} sp. strain AM-2504. Nucleotide and amino acid sequences of ribosomal protein S12 of \textit{Streptomyces} sp. strain AM-2504. The amino acids involved in the interaction with the antibiotic in dityromycin-sensitive ribosomes are circled. Amino acids involved in the self-resistance mechanism are circled in red.
shown in Fig. 4A, *E. coli* cells expressing a wild-type copy of S12, in the absence of the plasmid, are sensitive to streptomycin, but they become resistant once transformed. Moreover, the presence of either Val32Thr, Arg55Lys or Val78Ile substitutions did not affect the streptomycin-resistant phenotype. This demonstrates that the ribosomes effectively incorporate the S12 protein encoded by the plasmid and respond to the mutant genetic background supplied in *trans*. Furthermore, the results shown in Fig. 4B demonstrate that the ribosomal protein S12 carrying the Val32Thr or Arg55Lys substitution confer dityromycin resistance to *E. coli*, whereas the cells expressing the point mutation Val78Ile proved to be susceptible to this antibiotic.

**DISCUSSION**

The growing number of bacterial pathogens uniquely adapted to survive in the presence of one or more antibiotics is an extremely serious threat to global health. The term “antibiotic resistome,” a large repertoire of genes that has been originally developed by bacteria and is currently remodeled by the selective pressure imposed by drug use/abuse, has been coined to describe the genetic determinants involved in antibiotic resistance (23, 24).

Resistance to antibiotics is tightly linked to their production, since microorganisms producing these molecules must protect themselves against their action. In many cases, mechanisms of self-resistance have been shown to be mechanistically similar to the ones identified in clinical pathogens (25). Resistance genes are often carried by the same gene cluster of the antibiotic biosynthetic enzymes, and their expression is temporally coupled. Indeed, antibiotic producer cells need to express resistance determinants prior to, or concomitantly with, synthesis of the antibiotic molecule (26, 27). In recent years, the analysis of several genomes suggested that microorganisms produce only a fraction of the secondary metabolites that they encode, and new approaches have been used to identify biosynthetic gene clusters whose expression can be induced. The colocalization of resistance genes and biosynthetic genes in the genomes...
of underexploited groups of secondary metabolite producers, for example, has been used in pursuit of new biosynthetic pathways. This genome mining approach, guided by the search for self-resistant determinants, has been successful for topoisomerase inhibitors whose biosynthetic gene cluster has been identified by looking for putative pentapeptide repeat proteins, which are known to confer self-resistance to these molecules (28).

More than half of the antibiotics currently used inhibit protein synthesis by binding to the functional centers of the ribosome. Recently, important insights into the mechanism of these antibiotics have been obtained by X-ray crystallography and cryo-electron microscopy (Cryo-EM) analysis, opening the possibility of developing new molecules by structure-based drug design (29–36).

In this work, we characterized the mechanism of self-resistance of the dityromycin producer Streptomyces sp. strain AM-2504. Using E. coli and Streptomyces-based in vitro translation assays in combination with an ultracentrifugation pulldown assay, we demonstrated that the resistance determinants are localized in the Streptomyces 30S ribosomal subunit. Since the target of dityromycin is the ribosomal protein S12, we sequenced the gene coding for Streptomyces S12 and compared it with its E. coli homolog. Three amino acids in the Streptomyces protein (out of the five involved in the interaction with the antibiotic) differed from the E. coli protein sequence. Therefore, we suggest that the molecular basis underlying the self-resistance mechanism could be the reduced affinity of the antibiotic toward its target due to the amino acid substitutions on the ribosomal protein S12. To confirm this hypothesis, we introduced the observed amino acid substitutions into a plasmid-borne copy of E. coli rpsL gene harboring another mutation conferring resistance to the antibiotic streptomycin. The observed streptomycin resistance phenotype provides explicit, albeit indirect, evidence that the plasmid gene encoding ribosomal protein S12 is effectively incorporated into the 30S subunits. Even though the setup of the assay did not allow strict quantification of the number of copies of mutant protein incorporated, it showed unequivocally that this incorporation took place and that plasmid-encoded S12 bearing two specific mutations in the dityromycin binding site conferred resistance to the antibiotic. The results shown in Fig. 4B clearly indicate the effects of amino acid replacements at the three key sites of ribosomal protein S12 involved in the binding of the antibiotic. In particular, at position 32, the substitution of the nonpolar aliphatic valine with the polar amino acid threonine interferes with the hydrophobic interaction established with the dihydroxyproline group of dityromycin according to the crystal structure of the dityromycin-30S complex. It has been pointed out by crystallographic analysis that the arginine at position 55 packs against the N,N-dimethyl valine of the antibiotic (9). Our results strongly indicate that the replacement of a positively charged side chain, as in the case of lysine, is not sufficient to maintain the correct structure of the binding site, which presumably requires the chemical properties of the guanidinium group of arginine. This finding is consistent with the high specificity demonstrated by the arginine side chain in the interaction of proteins with their substrates (37, 38). On the other hand, we demonstrate that the substitution of Val78 with isoleucine does not interfere with the interaction of dityromycin with S12.

To identify the degree of conservation of the amino acid residues responsible for dityromycin resistance, we aligned the S12 sequences to 72 Streptomyces spp. randomly selected from the NCBI database. The alignment shows that, of the three residues considered, only Ile78 (Val in E. coli) is conserved in all Streptomyces sequences analyzed. The other two key residues are quite rare, with Thr32 (Val in E. coli) reported only in 7 out of 72 genomes (Streptomyces antioxidans, S. autoliticus, S. gilvosporeus, S. iranensis, S. malaysiensis, S. natalensis, and S. violaceusniger) and Lys55 (Arg in E. coli) in only two strains, S. gilvosporeus and S. natalensis. Therefore, the results of this comparative analysis show that only 2 of 72 Streptomyces strains share the same S12 residues conferring resistance to dityromycin as Streptomyces sp. strain AM-2504 (see Fig. S1 in the supplemental material). When the comparative analysis was extended to 30 bacterial strains randomly covering all relevant
branches of the phylogenetic tree, it was found that all selected bacteria display a
valine at position 32, embedded in a stretch of four amino acids (RGVC), which is
highly conserved and it is part of the “core” of a pseudo beta-barrel of S12 (Fig. S2A
and S2B). The other two residues are less conserved, since Lys55 and Ile78 are found
in 4 strains and 11 bacterial strains, respectively.

To our knowledge, the results of this work represent the first example of a self-
resistance mechanism mediated by a ribosomal protein in the producer strain. Resist-
ance mechanisms identified by isolation of single mutations are extremely rare among
ribosomal proteins. Point mutations in rpsL, the gene encoding S12, have been identi-
cified as the key factor conferring streptomycin resistance (39) and at the same time
activate the expression of cryptic antibiotics, like the blue-colored actinorhodin (40, 41).
This intriguing finding, which is the first reported evidence of an effect induced on gene
expression by a point mutation located on a ribosomal protein, is still lacking a detailed
explanation at the molecular level. A few other examples include protein L3, where a
single mutation has been shown to confer resistance to tiamulin, the semisynthetic
derivative of pleuromutiline, reducing its binding affinity for the ribosome and muta-
tions in L16 conferring resistance to the oligosaccharide antimicrobial agents avilamy-
cin (42) and evenemycin (43). In Pasteurella multocida, a bacterium responsible for
zoonotic diseases, a combination of amino acid changes on ribosomal protein S5 (point
mutation at Ser32 and amino acid deletion at Phe33) and a single transversion in helix
34 of 16S rRNA have been identified as the molecular determinants of spectinomycin
resistance (44).

In summary, we can conclude that Streptomyces sp. strain AM-2504 has devel-
oped a successful strategy to protect its own ribosomes from the effect of dityro-
mycin, combining the effects of three amino acid substitutions, which occur as
isolated point mutations in other Actinomycetes strains and are compatible with the
structural function of ribosomal protein S12. Although this resistance mechanism
does not fall into the category of antibiotic resistance transfer and spreading, in a
strict sense, the characterization of this strategy at the molecular level adds a new
piece to the complex puzzle underlying the coevolution of antibiotic production
and self-resistance mechanisms.

MATERIALS AND METHODS

Media. ISP2 medium (sporulation agar) consists of 0.4% yeast extract, 0.4% glucose, 1% malt extract,
and 2% agar. T1 medium contains 1% glucose, 2% starch, 0.5% yeast extract, 0.5% peptone, and 0.4%
CaCO3. JM medium consists of 0.1% yeast extract, 0.1% MgCl2 (6H2O), 0.3% Tryptone Soya, and 10%
sucrose. In all cases, the pH of the medium was adjusted to 7.0 prior to autoclaving. Luria-Bertani medium
was used for the growth of Escherichia coli strains utilized for propagation of plasmids and for the
preparation of ribosomes and cell extracts.

Culture conditions of Streptomyces sp. strain AM-2504. For the production and isolation of spores,
Streptomyces sp. strain AM-2504 was grown in ISP2 medium for 7 to 10 days at 28°C. The white aerial
mycelium, consisting of spores, was stripped from the upper layer of the plate using a spatula and
dispersed into 1 ml of sterile water. The spore suspension was filtered through a wad of absorbent
cotton, and the spores were concentrated by centrifugation at 4,000 rpm for 15 min. The spores were
resuspended in 100 μl of 20% glycerol and stored at –20°C.

Genomic DNA extraction. Streptomyces spores (20 μl) were inoculated in 20 ml of JM medium at
28°C for 18 h with aeration. Cells were collected by centrifugation, and genomic DNA was extracted using
the Invisorb spin plant minikit (Invitek).

DNA amplification and sequence of rpsL from Streptomyces sp. Oligonucleotide primers GSF1 and
GSR1 (Table 1), designed to cover the coding region of the rpsL gene from Streptomyces, were used to
amplify by PCR the target gene using genomic DNA as the template. The DNA sequence of the amplicon,
covering approximately 95% of the coding region, was analyzed to identify the amino acid variants
carried by the antibiotic producer strain.

Site-directed mutagenesis. Mutations in the E. coli rpsL gene were obtained using the QuikChange XL kit (Stratagene) and, as the template for the PCR, the plasmid vector pET11a carrying the
coding sequence of S12, derived from E. coli MC4100 (rpsL150 [Str]). The DNA sequences of the
oligonucleotide primers used to introduce codon variants are indicated in Table 1. E. coli DHS cells
were transformed with the mutagenized plasmids, and Sanger DNA sequencing confirmed the
presence of mutated nucleotides.

Preparation of E. coli and Streptomyces S30 cell extracts. E. coli MRE600 cells were grown at 37°C
in LB medium until they reached mid-log phase, while Streptomyces cells were grown at 28°C in T1
medium for 45 h. Both cell types were disrupted by grinding with precooled alumina in a chilled mortar
and processed following a previously described protocol (45). Both S30 extracts were extensively dialyzed against a buffer containing 10 mM Tris-Cl (pH 7.7), 10 mM Mg acetate, 60 mM NH₄Cl, and 0.5 mM dithiothreitol (DTT) and stored in small aliquots at −80°C.

**Isolation of ribosomes and ribosomal subunits.** The crude extract S30 was subjected to ultracentrifugation at 100,000 × g for 17 h at 4°C. The supernatant, called S100, was stored at −80°C, while the pellet, consisting of 70S ribosomes, was resuspended and split in two aliquots, one for *in vitro* assays and one for preparation of 30S and 50S ribosomal subunits. 70S ribosomes were dissociated into subunits at 1 mM Mg²⁺ concentration by ultracentrifugation at 25,000 rpm for 17 h at 4°C in a swing bucket rotor (SW28), on a 10% to 30% sucrose density gradient (45).

**In vitro translation assays.** Cell-free systems programmed with either poly(U) or 027 model mRNA were used to investigate the effect of dityromycin on the bacterial translational apparatus. Translation activity tests were performed using either S30 extracts or S100 extracts and 70S ribosomes, or criss-crossed combinations of *E. coli* and *Streptomyces* sp. S100 extracts and 70S ribosomes (46).

**Ribosome pulldown assay.** The binding of dityromycin to either *Streptomyces* or *E. coli* 30S ribosomal subunits was monitored by an ultracentrifugation-based assay. Increasing concentrations of 30S subunits were incubated for 10 min at room temperature with a fixed amount of antibiotic (1 μM) in 90 μl of a solution consisting of 10 mM Tris-Cl (pH 7.7), 10 mM Mg acetate, 60 mM NH₄Cl, and 0.5 mM DTT. After ultracentrifugation in a 5-100 AT-3 rotor at 100,000 rpm for 1 h at 4°C, aliquots of 20 μl of the supernatant were removed and tested in poly(U)-dependent *in vitro* translation assays.

**Determination of MIC.** To verify the incorporation of S12 mutants in *E. coli* ribosomes, a MIC plate (5 rows × 12 wells) was set up with increasing concentrations of streptomycin, ranging from 0.0 μg/μl to 12.0 μg/μl. Each well of the plate contained 190 μl of *E. coli* BL21(DE3)/pLysS cell culture. The plate rows consisted of the following: (i) no-plasmid control, (ii) pET11a expression vector with rpsL150, (iii) mutant V32/I32, (iv) mutant R55/K55, and (v) mutant V78/I78.

A MIC plate (4 rows × 12 wells) with the four tester strains and a gradient of dityromycin (from 0.0 μg/μl to 400 μg/μl) was used to estimate the ability of the mutant proteins to counteract the presence of the antibiotic. Further details are provided in the legend to Fig. 4.

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at https://doi.org/10.1128/mSphere.00554-19.

**FIG S1, JPG file, 2.2 MB.**

**FIG S2, TIF file, 2.6 MB.**

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