

Structural insights into species-specific features of the ribosome from the pathogen *Staphylococcus aureus*

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The emergence of bacterial multidrug resistance to antibiotics threatens to cause regression to the preantibiotic era. Here we present the crystal structure of the large ribosomal subunit from *Staphylococcus aureus*, a versatile Gram-positive aggressive pathogen, and its complexes with the known antibiotics linezolid and telithromycin, as well as with a new, highly potent pleuromutilin derivative, BC-3205. These crystal structures shed light on specific structural motifs of the *S. aureus* ribosome and the binding modes of the aforementioned antibiotics. Moreover, by analyzing the ribosome structure and comparing it with those of nonpathogenic bacterial models, we identified some unique internal and peripheral structural motifs that may be potential candidates for improving known antibiotics and for use in the design of selective antibiotic drugs against *S. aureus*.

antibiotic resistance | potential advanced pleuromutilin | species specificity | protein biosynthesis

The emergence of bacterial resistance to antibiotics threatens to cause regression to the preantibiotic era, as the treatment of infections with the available arsenal of clinically used antibiotics has been badly affected by the appearance of multidrug-resistant strains (1–3). Consequently, today many infections are caused by highly resistant bacterial strains of *Staphylococcus aureus*. These Gram-positive, versatile, and potentially aggressive strains are among the most worrisome pathogenic bacteria (3, 4).

A large number of antibiotics paralyze ribosomes, the universal cellular multicomponent RNA/PROTEIN particles that translate the genetic code into proteins in all living cells. Previous structural studies of the modes of action of antibiotics that inhibit bacterial ribosomes, performed using their complexes with then-available crystallizable ribosomal particles, yielded high-resolution crystal structures. These include the large ribosomal subunit of *Deinococcus radiodurans* (D50S) and the whole ribosome of *Thermus thermophilus* (T70S), as well as *Escherichia coli* (E70S) (5–7). These studies provided useful clues about the common traits of inhibitory mechanisms of antibiotics, namely binding at ribosomal functional sites, such as the peptidyl transferase center (PTC) or the protein exit tunnel (Fig. 1); illuminated the structural basis for the distinction between pathogenic bacteria and their mammalian hosts despite the high conservation of the ribosomal functional sites; shed light on antibiotic synergism; and highlighted the general principles of resistance and cross-resistance to antibiotics.

Based on their similar ribosomal RNA (rRNA) and ribosomal protein sequences, the structures of ribosomes from pathogens resemble those of ribosomes from other eubacteria. Nevertheless, species specificity in clinically relevant properties, particularly the modes of acquiring antibiotic resistance, have been identified (8). Given the knowledge that small structural differences between bacterial species can affect drug binding (9), for the progress of structure-based drug design, it is imperative to have a high-resolution crystal structure of the ribosome and its subunits from the pathogenic bacterial species. As a first step toward this goal, we determined the structure of large ribosomal subunit of the pathogen *S. aureus* (SA50S).

Infections caused by *S. aureus* have been treated with various drugs, including the ribosomal antibiotics oxazolidinones, pleuromutilins, macrolides, and ketolides, which bind to the large ribosomal

subunit. Eubacteria possess several copies of rRNA operons; thus, acquisition of single-nucleotide mutations and/or posttranscription modifications in the 23S rRNA, which are among the common resistance mechanisms with respect to ribosomal antibiotics that enter the cells, should occur after relatively long periods (10). Nevertheless, the epidemiology of *S. aureus* is increasing (11). Resistance mutations in *S. aureus* are also associated with ribosomal protein (rProtein) uL3, located in proximity to the PTC, as well as with uL4 and uL22 rProteins, whose segments are exposed in the exit tunnel (12–20).

Linezolid (Fig. 2), a synthetic antibiotic drug of the oxazolidinone class, binds at the PTC (21). It was approved by the Food and Drug Administration in 2000 to treat Gram-positive infections. As a synthetic drug, linezolid had no known preexisting resistance mechanisms, and resistance to it was expected to emerge rather slowly (22). Despite these expectations, however, *S. aureus*-acquired linezolid resistance was detected within a relatively short time. This resistance is conferred by a specific 23S rRNA point mutation (G2576U) (*E. coli* rRNA numbering is used throughout) (23). Additional resistant strains were identified later, including strains with rRNA mutations in U2500A (24), A2503G, U2504C, and G2447U (25), as well as those that have acquired a transmittable gene of rRNA methyltransferase *cfi* targeting A2503 (26). Within 10 years, linezolid resistance was detected in >1% of *S. aureus* clinical isolates (16).

Telithromycin (Fig. 2), a ketolide antibacterial drug that is structurally related to the macrolides, was developed specifically to provide optimal therapy for respiratory tract infections. All ketolides have two structural modifications compared with

Significance

Clinical use of the currently available antibiotics is severely compromised by the increasing resistance to them, acquired by the natural bacterial capability to manipulate their genomes. Many existing antibiotics target the fundamental process of protein biosynthesis, mainly by paralyzing the ribosome. Although antibiotics' modes of action are similar across most eubacteria, species specificity has been detected. We determined the structures of the large ribosomal subunit from *Staphylococcus aureus*, a pathogenic bacterial species with a known capacity to become multiresistant, and of its complexes with known antibiotic compounds, as well as with a novel potential pleuromutilin derivative. Our new insights provide unique chemical tools for enhanced distinction between pathogens and the useful benign microbiome, as well as for suggesting novel sites for potential future antibiotics.

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Data deposition: The atomic coordinates and structure factors have been deposited in the Protein Data Bank, www.rcsb.org (PDB ID codes 4WCE, 4WFB, 4WFF, and 4WFA).

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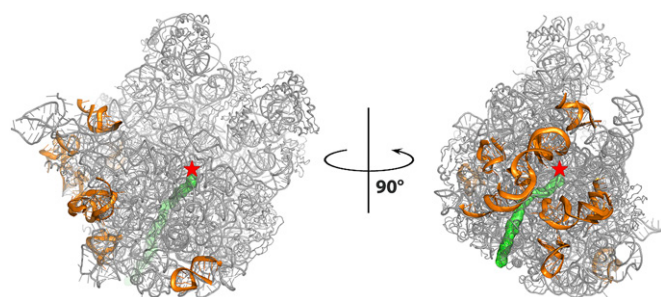


Fig. 1. The structure of the large ribosomal subunit of SA50S is shown in gray, the polypeptide exit tunnel is shown in green, and the PTC location is marked by a red star. The rRNA regions with fold variability compared with all other known structures (see below) on the SA50S are shown in orange and are rotated 90° to one another.

macrolides of previous generations: a C3-keto group and an additional long alkyl-aryl arm linked in telithromycin to the C11, C12-carbamate cycle. Importantly, telithromycin shows potent in vitro activity against *Streptococcus pneumoniae*, including strains resistant to macrolide-lincosamide-streptogramin B-ketolide (MLS_BK). Crystal structures of telithromycin in complex with the bacterial 50S subunits or 70S ribosomes reveal that telithromycin binds to the large ribosomal subunit at the macrolide-binding pocket in the ribosome's exit tunnel (27–30), and that in different species its flexible alkyl-aryl arm is stacked in various modes to different sites in the nascent protein exit tunnel.

Pleuromutilin (Fig. 2 and Fig. S1) is a natural product of the fungus *Pleurotus mutilus* (now called *Clitopilus scyphoides*) (31), which serves as a base for the synthesis of several semisynthetic antibacterial agents (32). All pleuromutilins consist of a tricyclic mutilin core, a C21 carboxyl group essential for antimicrobial activity (32), and various substituents at their C14 position, most of which are extensions with diverse chemical natures. Some pleuromutilins currently in clinical use in humans and veterinary medicine have elevated activity over a broad spectrum of pathogens, including multidrug-resistant staphylococci and streptococci. Among the group of C14-sulfanyl-acetate derivatives, retapamulin is a topical antibiotic that was approved for clinical use in 2007. All strains of *S. aureus* and *Streptococcus pyogenes* were susceptible to retapamulin with low minimum inhibitory concentrations, <0.5 μg/mL (33, 34). Other C14-sulfanyl-acetate derivatives, valnemulin and tiamulin, are approved for veterinary clinical use.

The crystal structures of complexes of the large ribosomal subunit from a nonpathogenic bacterium, D50S, with various pleuromutilin compounds (namely tiamulin, retapamulin, SB-264128, and SB-571519) reveal that the pleuromutilins are bound to the large ribosomal subunit at the PTC (35). In all cases, their cores are placed in a similar fashion at the A-site and their C14 extensions are pointing toward the P-site; thus, they directly inhibit peptide bond formation. Because the PTC is almost fully conserved, the clinical efficacy of the pleuromutilins stems from their efficient inhibitory modes, which are attained by structural diversity in the second or third shells around the PTC, by exploiting the ribosomal intrinsic functional flexibility for induced fit and remote conformational rearrangements that result in a tightening up of the binding pockets (35, 36).

Recent advances in pleuromutilin chemistry have yielded several new potential drugs, among which are BC-3205 (Fig. 2 and Fig. S1) and lefamulin (BC-3781). These novel semisynthetic pleuromutilin derivatives were developed by Nabriva Therapeutics (Vienna, Austria) for i.v. or oral treatment of community-acquired bacterial pneumonia and skin infections. Being 16- to 32-fold more potent than linezolid against *S. aureus*, BC-3205 exhibits clinically relevant potent antibacterial activity (37). Nabriva's lead product, Lefamulin, is about to enter phase 3 of clinical trials on community-acquired bacterial pneumonia (38, 39).

Here we present the crystal structure of the large ribosomal subunit from *S. aureus* alongside the structures of its complex with linezolid, telithromycin, and BC-3205. By analyzing these structures and their comparisons with ribosomes from eubacteria in complex with various oxazolidinones, ketolides, and pleuromutilins, we have elucidated specific traits that may govern *S. aureus* inhibition. We have also provided structural definitions of ribosomal features that could be selectively used in designing novel antibiotic drugs to target the *S. aureus* ribosome.

Results and Discussion

We determined the crystal structures of the large ribosomal subunit from *S. aureus* (SA50S) and of its complexes with linezolid, telithromycin, and BC-3205 at near-atomic resolution (Table 1). Most of the amino acid residues of 26 of the SA50S rProteins and 93% of the rRNA 50S nucleotides were reliably traced in the electron density map. Electron density maps around linezolid, telithromycin, and BC-3205 are shown in Fig. S2A–F. In the vicinity of the rRNA, we also detected an electron density that may be interpreted as hydrated ions, presumably of Mg²⁺ and Mn²⁺.

Main Features of *S. aureus* rRNA and rProteins. Sequence alignment of the 23S rRNA shows 81%, 76%, and 73% identity between *S. aureus* and *D. radiodurans*, *E. coli*, and *T. thermophilus*, respectively. Despite the high sequence conservation, local variability was detected. To highlight the unique features of SA50S, we compared it with the available eubacterial ribosome structures D50S (40) and the large ribosomal subunits in the T70S (6) and E70S (7) ribosomes (29, 41).

As expected, we found significant similarity in the rRNA folds of all high-resolution structures examined here, yet also identified several prominent differences on the surface of the 50S subunit. These differences are located mainly at the subunit surface (Fig. 1), and some at the interface with the small subunit within the active ribosome. Most of these differences are not involved in crystal contacts (all being located >5 Å from their crystal neighbors), and thus seem to originate from sequence variability or from the difference between isolated and bound large subunits. Examples include H25 (uppercase H is used throughout to refer to specific 23S rRNA helices), which has different folds in each of the four structures (Fig. 3A and Fig. S3B). Similarly, H63, which is located in proximity to the intersubunit bridge B5 (42), has a different length in each of the four structures (longest in E70S, shorter in SA50S and T70S, and shortest in D50S) (Fig. 3B and Fig. S3D). Helices H10 and H79, which interact with each other and with rProteins bL28 and uL2, respectively, are extended in the pathogen's SA50S and E70S compared with their conformation in T70S and D50S (Fig. 3C and D). H16 of SA50S and E70S are longer than H16 in T70S and D50S (Fig. 3D). Of interest, H15 shows small structural diversity between SA50S and T70S that may result from different interactions with a symmetry-related molecule owing to crystal packing, indicating its flexibility. It could not be traced in D50S electron density maps and does not exist in E70S (Fig. 3D). Also of note, H68, which is involved in the binding of rProtein uL1, elongation factor-G, and ribosomal recycling factor, and belongs to the intersubunit bridge B7a (42) with the 30S subunit, is longer in SA50S than in T70S and D50S; however, it is not fully

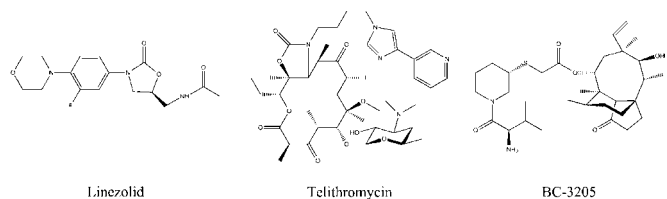


Fig. 2. Chemical structures of the oxazolidinone linezolid, the ketolide telithromycin, and the pleuromutilin BC-3205.

Table 1. Crystallographic data and refinement statistics of SA50S (native 50S), SA50S in complex with linezolid (SA50Slin), SA50S in complex with telithromycin (SA50Stel), and SA50S in complex with BC-3205 (SA50S-BC3205)

Statistic	SA50S	SA50Slin	SA50Stel	SA50S-BC3205
Crystal information				
Space group	P6 ₅ 22	P6 ₅ 22	P6 ₅ 22	P6 ₅ 22
a = b, Å	279.8	279.9	282.7	280.9
c, Å	872.7	870.6	877.0	875.6
α, β, γ, °	90, 90, 120	90, 90, 120	90, 90, 120	90, 90, 120
Diffraction data statistics				
X-ray source (ESRF)	ID23-2, ID23-1, ID23-1	ID23-1, ID23-1	ID23-1	ID23-2
Wavelength, Å	0.873, 0.972, 1.000	0.972, 1.000	0.973	0.873
Number of crystals	11	8	11	20
Resolution, Å*	50–3.51 (3.57–3.51)	200–3.4 (3.46–3.4)	50–3.45 (3.51–3.45)	50–3.43 (3.47–3.43)
Observed reflections	2,813,593	1,392,748	1,792,405	2,109,298
Unique reflections	236,855	246,474	257,382	253,918
Redundancy	12.0 (6.0)	5.7 (2.8)	7.0 (5.0)	8.3 (3.4)
Completeness, %	96.3 (84.3)	89.4 (64.8)	98.5 (96.5)	92.1 (78.9)
<I>/<σ(I)>	7.40 (1.43)	8.45 (1.16)	8.84 (1.21)	10.51 (1.21)
R-merge, %	25.3 (97.8)	15.6 (77.6)	16.5 (92.6)	15.2 (74.4)
Refinement statistics				
R-factor, %	20.18	20.18	19.19	20.42
R-free (5%), %	24.65	24.28	23.15	24.20
rmsd bonds, Å	0.006	0.006	0.006	0.006
rmsd angles, °	1.161	1.086	1.106	1.099

ESRF, European Synchrotron Radiation Facility.

*Values for the highest-resolution shells are in parentheses.

traced in the SA50S electron density map (Fig. 3E). Additional fold variability was observed for H28, which has a different orientation in SA50S compared with the other three structures (Fig. 3F and Fig. S3C).

Sequence alignment of the large ribosomal subunit proteins showed an overall identity of ~50% between *S. aureus* and *E. coli*, *T. thermophilus*, and *D. radiodurans*. Consequently, the rProteins show a greater diversity in their detailed structures compared with

the differences observed for the rRNA structure. Typically, the main features of the rProtein folds, such as the globular domains and the secondary structural elements that interact with the rRNA, are rather conserved; however, several SA50S rProteins contain extensions that were not seen in any other published ribosome structure. (We use *S. aureus* rProtein residues numbering throughout, to minimize confusion owing to the significant variability in the rProtein sequences and folds of the four organisms.)

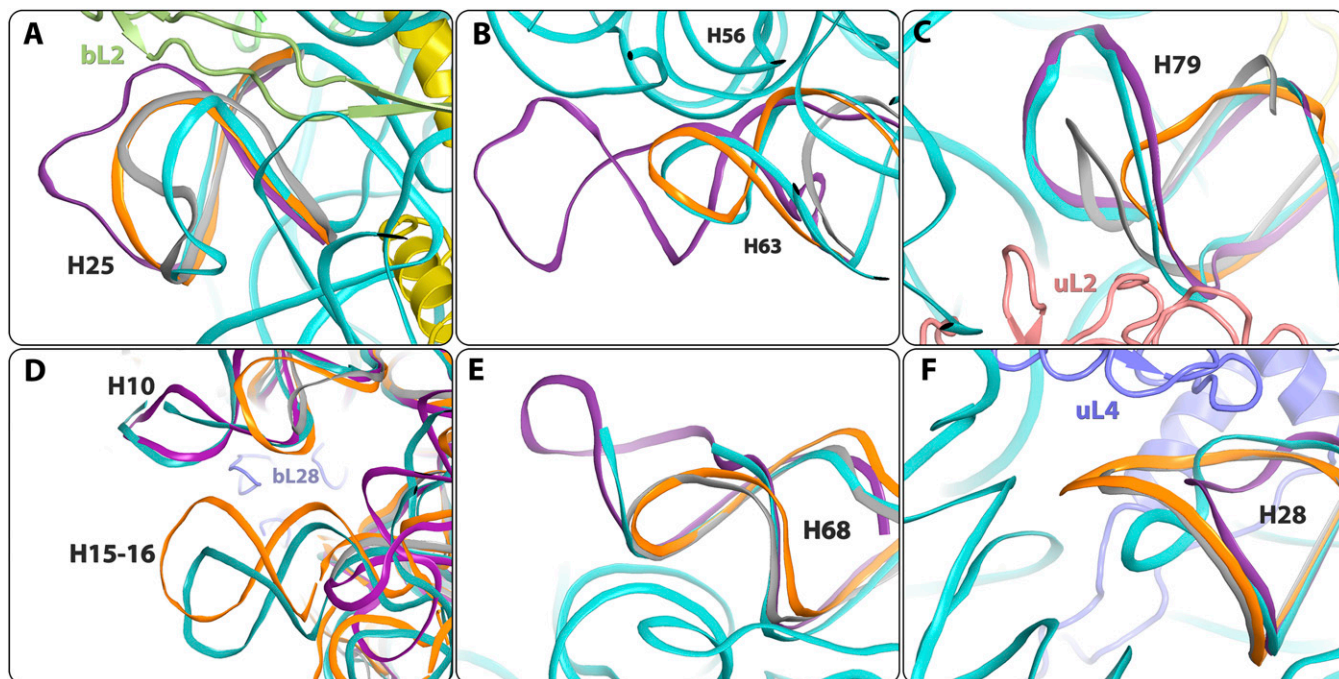


Fig. 3. Details of structural variability in the rRNA backbone among the four known eubacteria structures. SA50S is in teal, *E. coli* (E70S) is in purple, *D. radiodurans* (D50S) is in gray, and *T. thermophilus* (T70S) is in orange. (A) H25. (B) H63. (C) H79. (D) H10 and H15-16. (E) H68. (F) H28.

Important differences in *S. aureus* were detected in the N-terminal region of bL32 rProtein, resulting in the creation of a unique void at the rims of the macrolide-binding site. Thus, in D50S and T70S, the N-terminal tails of bL32 reach the erythromycin-binding pocket, whereas in the ribosome from SA50S and E70S, they are distal to this pocket (Fig. 4A and Figs. S3H and S4A). Sequence alignment of bL32 to other 721 eubacterial species (43) confirms that the elongated N-terminal end of bL32

is unique to *Deinococcus* and *Thermus* species. In addition, in all known structures of eubacterial ribosome nucleotide A2058 is stacked to G2057, which interacts with residue 5 of the bL32 N-terminal tail. The sequence and structure variability of this residue (Lys in SA50S, T70S, and D50S and Gln in E70S) creates different environments around G2057. Of interest, this protein does not exist in the archaeon *Haloarcula marismortui* or in eukaryotes.

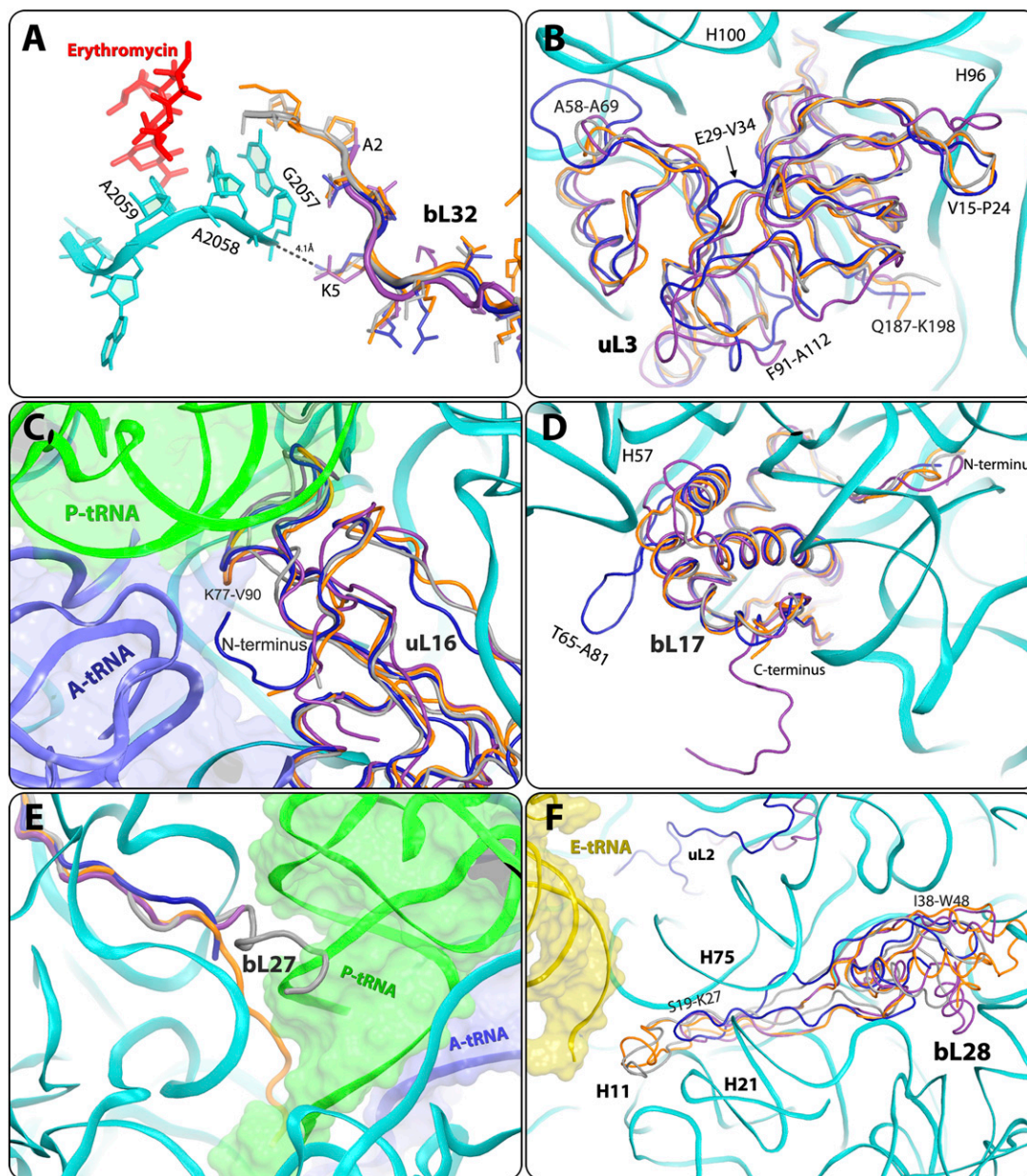


Fig. 4. Structural differences in rProteins. SA50S is shown in navy, *E. coli* (E70S) is in purple, *D. radiodurans* (D50S) is in gray, and *T. thermophilus* (T70S) is in orange. (A) The N-terminal end of protein bL32 that resides in the second shell around the erythromycin-binding pocket is shorter in *S. aureus* (SA50S, navy) and *E. coli* (E70S, purple) compared with in *D. radiodurans* (D50S, gray) and *T. thermophilus* (T70S orange), and thus may provide a space for a specific extension of erythromycin (Fig. S5). (B) SA50S rProtein bL3 has a unique extended A58-A69 loop compared with D50S, T70S, and E70S. (C) SA50S uL16 points opposite to E70S and thus may provide a potential specific binding site (Fig. S5) for a compound that may interfere with binding of the A-site tRNA acceptor stem (blue). P-site tRNA (green) interacts with the loop K77-V90, which has some structural variability. (D) SA50S bL17 has an extended (T65-A81) loop that is unique to SA50S. Because it is exposed on the surface of the SA50S subunit, it may provide a potential specific binding site (Fig. S5) for a compound that may interfere with the two subunits association. Its C terminus is ~10 aa longer in E70S relative to the other three organisms. (E) In D50S, bL27 reaches the position of the acceptor stem of the P-site tRNA (green), whereas in T70S it reaches the PTC, in the proximity of the CCA-end of the P-site tRNA. (F) bL28 is located near the 50S surface, close to the position of the CCA 5' of E-site tRNA (yellow). In D50S and T70S, it has a 15-residue extended loop (S19-K27) This loop is shorter in SA50S and E70S and does not exist in D50S and T70S, and thus may provide a potential site for a compound that may interfere with the binding of the E-site tRNA acceptor stem (yellow) (Fig. S5).

SA50S rProtein uL3 has an extended (A58-A69) loop next to h100 of 23S rRNA. Sequence alignment of uL3 to other 756 eubacterial species (43) confirms that this loop is unique to staphylococci and thus may serve as a new antibiotic target site against staphylococci ribosomes. In addition, variability in the SA50S uL3 backbone fold compared with D50S, T70S, and E70S was observed. Among these, E29-V34 and F91-A112 and loops V15-P24, S142-P151, G152-P170, and Q187-K198 have a different fold and orientation in SA50S than in T70S and E70S (Fig. 4B and Fig. S3E).

uL16 protein is positioned in the intersubunit surface and interacts with the A- and P-site transfer RNAs (tRNAs) (superimposed on PDB ID code 2WDK) (6). The N terminus of uL16 SA50S is pointing in the opposite direction compared with E70S, and thus may form a unique network of interactions with the acceptor stem of A-site tRNA (Fig. 4C and Figs. S3F and S5C).

rProtein bL17 of SA50S has a unique extended (T65-A81) loop. Sequence alignment of bL17 to other 788 eubacterial species (43) confirms that this loop has a unique conserved sequence in staphylococci and thus may provide a potential specific binding site, because it may interfere with subunit association or with nonribosomal factor interactions. The N terminus of SA50S bL17, which is located in proximity to H100-H101, H61, and H96, is four residues shorter relative to that in the other three structures. Its C terminus is ~10 aa longer in E70S relative to the other three organisms (Fig. 4D and Figs. S3G and S5D).

The bL27 N-terminal has various conformations in the four known structures. In SA50S, it is traced only from residue 19 (based on sequence alignment with PDB ID code 2ZJR) (Fig. 4E). It was recently reported that in *S. aureus*, bL27 has an extended N terminus that is cleaved posttranslationally after amino acid 9 by a specific essential protease, before or concurrent with ribosome assembly (44). In D50S, it reaches the acceptor stem of the P-site tRNA (40), whereas in T70S, it reaches the PTC and interacts with the CCA 3' end of the P-site tRNA (6). In the E70S structure, its five N-terminal residues are missing in the electron density map, and the N-terminal tail is shorter than in the other large subunits. In *E. coli*, rProtein bL27 appears to be essential for cell survival and for peptide bond formation (45), and the variations in its length and orientation apparently do not hamper its functional key role.

bL28 rProtein has a 15-residue extended loop (S19-K27) that reaches H11 of the 23S rRNA and interacts with the CCA 5' of E-site tRNA in the T70S and D50S structures. This loop is shorter in SA50S and E70S (Fig. 4F) and, although it interacts with H21 and H75, it does not reach H11 or the E-site tRNA. In addition, the core of bL28, which is situated in proximity to the outer surface of the large subunit, is folded differently in the four structures. Moreover, the bL28 I38-W48 loop adopts a different orientation in SA50S than in E70S, T70S, and D50S (Fig. 4F and Fig. S5F).

A detailed comparative structural analysis presenting the main structural differences found in SA50S rProteins and rRNA is summarized in Tables S1 and S2 and shown in Figs. S4, S6, and S7.

Structure of the SA50S-Linezolid Complex. The SA50Slin crystal structure indicates that linezolid is bound at the PTC, blocking the A-site in an orientation clearly similar to that observed in other ribosome linezolid complexes with a D50S (46), H50S (47), and E70S linezolid model (48). However, in the SA50Slin complex, the flexible nucleotide U2585 (49) undergoes significant rotation and forms a hydrogen bond with the O4 of the linezolid morpholino ring, leading to a nonproductive conformation of the PTC (Fig. 5A and B).

The linezolid 1,3-oxazolidin-2-one moiety and acetamide group form additional hydrogen bonds with G2505 and A2451, respectively. The fluorophenyl moiety of linezolid is located in a heteroaromatic crevice formed by the PTC residues C2451 and C2452, the so-called "A-site cleft." All of its other interactions with rRNA nucleotides G2061, C2501, U2504, U2506 and G2447 are either van der Waals or hydrophobic.

Comparing the SA50Slin structure (Fig. 5A) with other available crystal structures or models of large ribosomal subunits in complex

with linezolid, namely of H50S (H50Slin) (47), D50S (D50Slin) (46) and E70S (E70Slin) (48) (Fig. 5B) revealed that linezolid binds in the same pocket in all structures. However, there are subtle differences between the conformations of the linezolid acetamide group in SA50Slin and in the other structures. These include a 100°–120° rotation of the acetamide group, which in SA50Slin enables fixation of A2451 by a hydrogen bond. Of note, the H50Slin complex was crystallized in the presence of CCA-Phe, a tRNA-3' end substrate analog, assuming that in such an environment it should interact strongly with the 50S subunit P-site (47). Indeed, it seems that this P-site analog altered linezolid conformation compared with other linezolid complex structures that were determined with an empty P-site. Of importance, a hydrogen bond between the morpholino ring of linezolid and nucleotide U2585 was observed in D50Slin and in SA50Slin, but not in the complex of the archaeal H50S with linezolid.

Structure of the SA50S-Telithromycin Complex. The SA50Stel structure shows that telithromycin is bound at the MLS_BK-binding site and forms the typical ketolide (and macrolide) hydrogen bond between its desosamine sugar and A2058. At this position, it is partially blocking the protein exit tunnel, as has been found in other ribosome-MLS_BK complex structures (5, 27, 50). In the SA50Stel complex, the flexible nucleotide A2062 is rotated compared with its conformation in native SA50S, and its conformational range is minimized by the hydrogen bond with A2503 (Fig. 5C). All of the other interactions of telithromycin with the rRNA nucleotides G2505, A2059, C2611, and U746 are either hydrophobic or based on van der Waals distances.

Comparing the SA50Stel structure with other available crystal structures of ribosomal particles in complex with telithromycin (27–30) revealed that in all structures, the drug is bound at the same pocket but with distinct differences in orientations of the alkyl-aryl moiety. The orientation of this moiety was similar in SA50Stel and in H50Stel. In both, the arm is folded back over the macrolactone ring, creating a rather compact structure of the drug; however, in SA50Stel, the alkyl-aryl arm reaches the center of the tunnel ~5 Å closer to the PTC compared with its location in H50Stel, and may block nascent protein progression at an earlier stage. In this orientation, the alkyl-aryl arm is almost overlapping the location of the cladinose sugar of erythromycin in its complex with D50S (Fig. 6F) (5). In contrast, in T70Stel and E70Stel, this arm is stacked to A752 and U2609, a base pair located on the tunnel wall further away from the PTC, so that it can block nascent protein progression ~10 Å away from the point of blockage observed in H50Stel. In D50Stel, the alkyl-aryl arm is extended, and thus it blocks the tunnel by interacting with U790 located across the tunnel, creating a barrier located 15 Å further along the tunnel compared with SA50Stel (Fig. 5D). It has been suggested that the structure of E70Stel reflects the telithromycin binding mode to the ribosomes of medically relevant (namely pathogenic) eubacteria species, because the A752-U2690 base pair is conserved among all eubacteria (29). In addition, telithromycin partially protects A752 from chemical modification (29), and the telithromycin-resistant *S. pneumoniae* ΔA752 mutant has been isolated (51). Nevertheless, in SA50Stel, the alkyl-aryl arm of telithromycin does not interact with the A752-U2690 base pair. This observation clearly demonstrates that the general description of the overall antibiotic-binding properties, which is the main outcome from the previous structures, cannot be extrapolated safely to other species.

Structure of the SA50S-BC3205 Complex. In the SA50S-BC3205 crystal structure, BC-3205 is bound at the PTC, so that the tricyclic mutilin core is blocking the A-site, and its C14 extension is pointing into the P-site, thus perturbing A- and P-site tRNA accommodation, as was seen in the ribosome-pleuromutinin complexes with D50S (35, 36). In the SA50S-BC3205 complex, the conformation of the flexible nucleotide U2585 is different from that of the unbound SA50S, and its conformational range is reduced because of partial overlap by the BC-3205 (Fig. 5E). In addition, U2506 is shifted

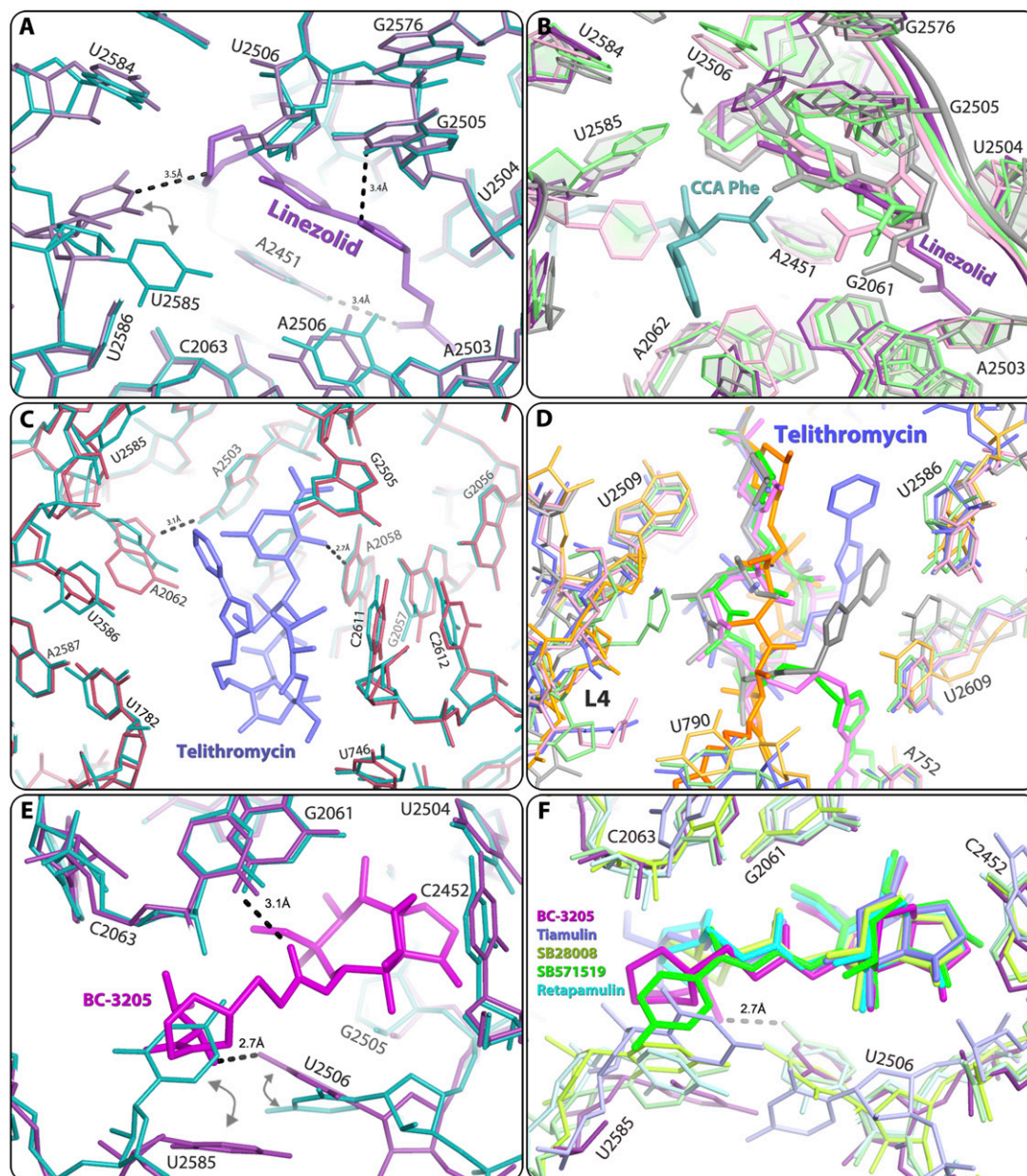


Fig. 5. The linezolid-binding site as identified in the SA50Slin complex structure (A and B), with hydrogen bonds between the bound drugs and 23S rRNA shown in black dashes, the telithromycin-binding site as identified in the SA50Stel complex structure (C and D), and the BC3205-binding site as identified in the SA50S-BC3205 complex structure (E and F). (A) Comparison of native SA50S PTC (teal) and SA50Slin complex (pale purple) structures. (B) Overlay of the structures of various ribosome-linezolid complexes, including SA50Slin (pale purple; this study), H50Slin (green) + CCA-Phe substrate analog (teal) (PDB ID code 3CPW) (47), and D50Slin (gray) (PDB ID code 3DLL) (46), and of the model of E70Slin (pink) (48). The color coding of the rRNA components of the various linezolid-binding pockets is the same as that of the corresponding linezolid molecules. (C) Comparison of the structures of native SA50S PTC (teal) and SA50Stel complex (red). (D) Structural overlay of telithromycin conformations observed in various ribosome-telithromycin complex structures: SA50Stel (slate; this study), D50Stel (orange) (PDB ID code 1P9X) (27), H50Stel (gray) (PDB ID code 1YUJ) (28), E70S-tel (pink) (PDB ID code 3OAT) (29), and T70Stel (green) (PDB ID code 3OIB) (30). The color coding of the rRNA components of the various telithromycin-binding pockets is in a brighter tone than the corresponding telithromycin molecules. (E) Comparison of the PTC structure in native SA50S (teal) and in SA50S-BC3205 complex (purple). The arrows indicate the movements of nucleotides U2585 and U2506 in the bound vs. native structure. (F) Structural overlay of various pleuromutilins in their binding pockets: SA50S-BC3205 (violet; this study), D50S-SB571519 (green) (PDB ID code 2OGM), D50S-retapamulin (cyan) (PDB ID code 2OGO), D50S-tiamulin (slate) (PDB ID code 1XBP), and D50S-SB280080 (lemon) (PDB ID code 2OGN). Only one hydrogen bond between BC-3205 and 23S rRNA is shown here (as black dashes) for increased clarity.

toward the walls of the binding pocket, forming a hydrogen bond between its O5 carbonyl and the NH₂ of the valyl moiety of BC-3205. An additional hydrogen bond is formed between the acetyl carbonyl and the NH₂ of G2061. All of its other interactions with the rRNA nucleotides A2063, A2503, U2504, G2505, A2451, C2452, and U2585 are either hydrophobic or based on van der Waals forces.

Comparing the crystal structure of SA50S-BC3205 with those of D50S in complex with various pleuromutilins (35, 36) (Fig. 5F) revealed that all pleuromutilins bind to the same pocket, albeit by creating somewhat different interaction networks, including a shift of U2585. This shift is larger in SA50S-BC3205 compared with all other pleuromutilin complexes, so that the base of U2585 is located ~6 Å away from its position in unbound SA50S.

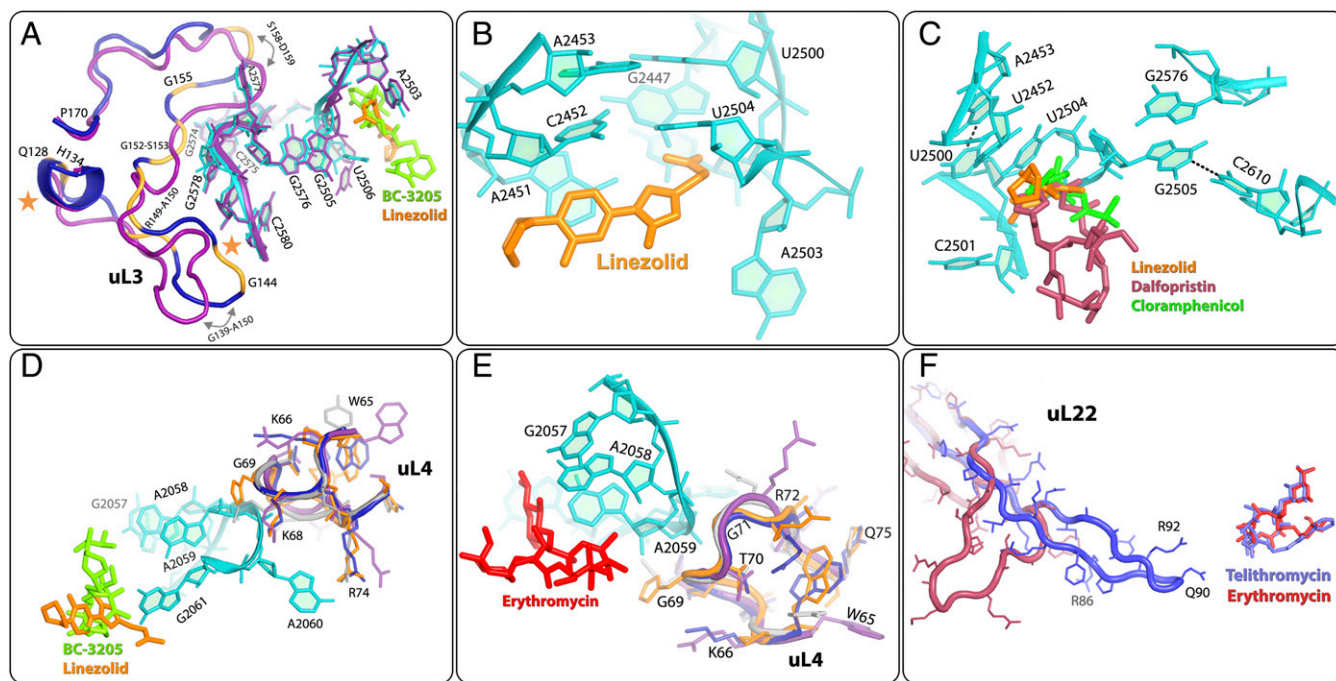


Fig. 6. Resistance and cross-resistance mechanisms in *S. aureus*. SA50S rRNA and rProteins are colored in navy. D50S (gray), E70S (purple), and T70S (orange) structures are superimposed on the SA50S structure for comparative analysis. rRNA nucleotides of *S. aureus* are shown only in regions in which they can be well aligned with the corresponding nucleotides in all other structures used for the comparisons. (A) SA50S rRNA nucleotides of the linezolid-binding sites (orange) and BC-3205-binding sites (green) are superimposed on the corresponding E70S rRNA (purple). G2576 is located in the second shell around the linezolid- and BC-3205-binding sites, in proximity to the first shell nucleotides G2505 and U2506; thus, its mutations may cause alterations in them. The locations of the uL3 mutations acquiring resistance to *S. aureus* are marked on the protein chain (yellow). For comparison, the structure of *E. coli* uL3 (purple) is superposed on SA50S uL3 (navy). Key structural differences are marked by arrows. The orange stars indicate deletions. (B and C) Linezolid (orange), chloramphenicol (green), and dalfopristin (streptogramins A, in raspberry) are shown in the SA50S rRNA-binding pocket. (D) uL4 rProtein is in the vicinity of *S. aureus* linezolid-binding (orange) and *S. aureus* BC-3205-binding (green) pockets. The structural variability of its loop (W65-Q75) in four species is shown. (E) SA50S rRNA A2058 and A2059 are main binding determinants of MLS_BK family of antibiotics, represented here by erythromycin (PDB ID code 3OFR) (red). uL4 and bL32 rProteins form a second shell around the erythromycin-binding pocket next to A2058 and A2059. Structural variability of uL4 among D50S, E70S, T70S, and SA50S is shown. (F) *S. aureus* telithromycin (slate) conformation within its *S. aureus* complex, where its alkyl-aryl arm is folded such that it overlaps the desosamine sugar of erythromycin (red) in its complex with D50S. SA50tel uL22 (blue) is superimposed on the same rProtein in *H. marismortui* large ribosomal subunit with a deletion mutation $\Delta 82-84$ in uL22 (red) that allows protein progression even though it binds erythromycin (PDB ID code 1YJ9).

Similarly, a movement of U2506 toward the bound drug was observed in pleuromutilin complexes with D50S. Notably, in the SA50S-BC3205 complex, U2506 forms a hydrogen bond with the NH₂ of the valyl moiety of the compound, and thus its shift is the largest. Consequently, the two sides of the BC-3205 molecule are held within its binding pocket by hydrogen bonds, compared with the single hydrogen bond created in the other pleuromutilin complexes with G2061, indicating a better fit of BC-3205 to its binding site. This additional interaction of BC-3205 seems to account for its greater potency against methicillin-resistant *S. aureus*-resistant strains and its low IC₅₀ value in a *S. aureus* cell-free translation system.

Structural Analysis of Known *S. aureus* Resistance Mechanisms. *S. aureus* linezolid resistance is caused mainly by the G2576U mutation (23), a nucleotide that is ~98% conserved throughout all kingdoms of life (52). The G2576U resistance mechanism is in accord with all crystal structures of ribosome-linezolid complexes, as well as with a model of such a complex (46–48). These structures indicate that linezolid binds to the PTC, and that linezolid blocks the A-site within the PTC. Of interest is the G2576U mutation, which is located in the second shell of the PTC and is also associated with linezolid resistance in the multidrug-resistant bacteria *Enterococcus faecium*, *Enterococcus faecalis* (53), and *S. pneumoniae*, as well as in *Mycobacterium smegmatis*, the species used in the diagnosis of *Mycobacterium tuberculosis* (54). G2576 is located in the second shell around the linezolid-binding pocket in the PTC, in the vicinity of the uL3 rProtein. Compared with E70S, the uL3 loop G152-P170

in SA50S is bent toward the minor groove of H72 away from G2576, thereby permitting more flexibility (Fig. 6A).

S. aureus linezolid resistance is also caused by rRNA mutations U2500A (24), A2503G, U2504C, and G2447U (25) of nucleotides that are ~98% conserved throughout all kingdoms of life (52). In SA50S, D50S, and H50S linezolid complexes, linezolid interacts with U2504 (46, 47). Mutating it to C will abolish the C:U non-canonical base pair interactions (55) with C2452, and thus likely change the shape of the linezolid-binding pocket. G2447 interacts with A2451; thus, the mutated G2447U may form a new base pair with A2451, which may limit A2451 flexibility and thereby influence drug binding. U2500 is base-paired with A2453. Its mutation to A will abolish the base pair interactions and consequently change the PTC environment (Fig. 6C) and prevent drug binding (56). The G2447U mutation, which renders *M. smegmatis*, *M. tuberculosis* (57), and *E. coli* (58) linezolid-resistant, has been suggested to be involved in functional differences between Gram-positive and Gram-negative bacteria (59). Of interest, significant structural similarity was observed in this region of ribosomes from T70S, D50S, E70S, and SA50S, which represent both Gram-positive and Gram-negative species.

Cross-resistance to linezolid, chloramphenicol, and dalfopristin (the streptogramin A component of Synercid) is associated with the second shell nucleotide G2576U mutation (10). Linezolid, chloramphenicol, and streptogramin A bind at the PTC via the first shell nucleotide G2505, which is stacked to U2576. In addition, cross-resistance of *S. aureus* to linezolid and chloramphenicol, of which the binding pockets partially overlap, may be caused by

G2505A and U2500A mutations (60), even though the conservation level of these nucleotides is ~98% throughout all kingdoms of life (52). These mutations indicate that despite its high conservation, this nucleotide is not essential for ribosomal function. Nucleotide G2505 is base-paired to C2610, and its ribose and phosphate are located in the vicinity of the antibiotic. The mutation G2505A leads to a mismatch in this base pair, thereby altering the antibiotic binding pocket. Similarly, U2500 is base-paired with A2453 and has hydrophobic interactions with U2504. The mutation U2500A disrupts the base pair U2500-A2453, and thus may allow U2504 to tilt away from the linezolid binding pocket (56) (Fig. 6B).

S. aureus-resistant mutations are also associated with the region of the uL3 rProtein located in proximity to the PTC (12–17, 61). The fold of uL3 loop G139-A150 is similar in SA50S to its folds in D50S and T70S, all bending toward H90 and H62, but is different in E70S, where it bends toward the H61 major groove (Fig. 6A). All of the mutated amino acids found in clinically resistant isolates are located in the uL3 region, which is adjacent to the third shell of nucleotides around the PTC, and appear to reshape the antibiotic-binding pocket (marked in yellow in Fig. 6A). Mutations in the uL3 loops in the vicinity of G2578 and C2575, which are part of the second shell nucleotides, may affect G2576 and its interactions with nucleotides G2505-U2506, which are part of the pleuromutilin- and linezolid-binding pockets (35, 46).

The uL3 loop F127-P170 is also located in the vicinity of A2572, which interacts with the flexible nucleotide U2504. In D50S and *S. aureus*, this nucleotide is located in the first shell around the linezolid- and tiamulin-binding pockets, and interacts with them (Fig. 6A). The nucleotide A2572 has been previously implicated in PTC antibiotic resistance acquired by induced-fit alterations that, although not directly interacting with the bound drugs, reshape the binding pocket via networks of remote interactions, most of which occur through the flexible nucleotide U2504 (56). In another *S. aureus* linezolid-resistant strain, a deletion of F127-H146 occurs in the *rplC* gene, which encodes uL3 rProtein (12). In wild-type uL3, this loop is located within the area described above, in the vicinity of 23S rRNA H90 nucleotides G2576-G2580, which reside in the second shell around the linezolid-binding pocket. Loop deletion may eliminate or alter interactions of uL3 with its proximal rRNA, and thus may reshape the drug-binding pocket and hinder antibiotic binding (Fig. 6A). Similar mutations in locations somewhat distal from the actual antibiotic-binding pockets that confer resistance by allosteric rearrangements have been detected in other systems as well (54, 62, 63).

A2058 and A2059, known as the main determinants for binding MLS_BK antibiotics by direct interaction (5, 64), are in direct contact with the W65-Q75 loop of uL4 rProtein, which has diverse sequences in SA50S, E70S, T70S, and D50S (Fig. 6D and E). Mutations in uL4 rProtein, G69A, and T70P produce *S. aureus* resistance to linezolid (65), whereas Q67K, G69E, G69A, and T70P mutations produce *S. aureus* resistance to erythromycin (15). In addition, ΔK68 and ΔG69 are susceptible to linezolid, and K68Q mutation produces *S. aureus* resistance to linezolid, pleuromutilins, chloramphenicol, and tedizolid (17). Because the uL4 W65-Q75 loop is located in the vicinity of the phosphate of A2059, which belongs to the MLS_BK-binding pocket, the aforementioned mutations may alter the W65-Q75 loop conformation, affecting A2059 conformation (Fig. 6D and E). In E70S, T70S, and SA50S structures, residue 72 of uL4 is Arg, whereas in D50S it is Asn. Each of these four side chains of residue 72 point to a different direction, creating a different environment around nucleotide A2059 in the four crystal structures compared. Residue K68 of uL4 is located in the vicinity of the phosphate of nucleotide G2061 and the sugar-phosphate backbone of A2059, which are in direct contact with linezolid (46). It has a similar orientation in E70S, SA50S and T70S, but different from that in D50S (Fig. 6D). The mutation K68Q replaces a long, positively charged side chain with a shorter, uncharged side chain with no electrostatic interactions with G2061 and A2059. G69A and T70P point mutations in uL4 introduce hydrophobic residues, which may increase the flexibility of the binding pocket, thereby reducing the efficiency of binding and explaining

how resistance to linezolid is gained in these cases. All available National Center for Biotechnology Information sequences of uL4 rProtein indicate that its residue 68 lysine is highly conserved; however, in H50S, the respective residue is serine, a short polar uncharged residue, which seems to reduce binding efficiency. This may be an additional feature of the weaker binding mode of linezolid to H50S (46, 47).

Point mutations, deletions, and insertions in the uL22 hairpin loop that is part of the exit tunnel wall in residues R80-S108 in *S. aureus* confer resistance to erythromycin, Synercid, and telithromycin, although its location is too far to allow direct chemical interactions with the bound drugs (18, 19). Importantly, similar resistance mutations were also observed in additional bacterial species. The crystal structure of one of these mutants of a *H. marismortui* large ribosomal subunit with a deletion mutation Δ82–84 in uL22 demonstrates that the deletion triggers a shift in the position of the uL22 hairpin loop and it is located deeper in the tunnel (28). It is conceivable that similar mechanisms play roles in acquiring resistance in *S. aureus* deletion/insertion mutants.

Methylation of the C8 position of A2503 in *S. aureus* by the *Cfr* enzyme leads to multidrug resistance to phenicols, lincosamides, oxazolidinones, pleuromutilins, and streptogramin A (PhLOPS_A) (26, 66, 67). The A2503 nucleotide interacts with linezolid and is ~98% conserved throughout all kingdoms (52); nevertheless, it can be modified without interfering with protein biosynthesis. A2503 methylation of C8 by *Cfr*, which increases its size, interferes with PhLOPS_A drug binding, thereby conferring resistance to these drugs. In addition, hindering the C2 position of A2503 indigenous methylation by inactivating the gene of the methyltransferase *RlmN* renders resistance to linezolid, because a lack of methyl substitution may change its orientation with respect to the drug (25, 68) (Fig. 6A and B).

Summary and Concluding Remarks. The sequence and structure alignments of *S. aureus* large ribosomal subunit rRNA with other eubacteria indicate high conservation. Nevertheless, diversity was identified in several locations, some of which may serve as antibiotic target sites. Furthermore, structural characterization of this ribosomal particle and of its complexes with linezolid, telithromycin, and BC-3205 revealed minute but nonetheless significant differences. In this respect, the superiority of the new pleuromutilin compound BC-3205 over known veterinary pleuromutilins or retapamulin, as demonstrated by its strong binding to *S. aureus* ribosome and low minimum inhibitory concentration, is clearly supported by the structural elements of its binding mode.

Our structural analysis identified structural motifs within the particle core and on its periphery that do not belong to the known binding sites of antibiotics but may be candidates for the design of novel selective drugs against *S. aureus*, thus enriching the pool of potential antibiotic compounds. Several of these features are shown in Fig. 4 and highlighted in Fig. S5. Some of these features may provide inhibitory mechanisms similar to that of the periphery binding site of RsfS that inhibits protein synthesis by interfering with the association of the two subunits (69, 70).

Analysis of the structures of the unbound and drug-bound ribosomes vs. ribosome structures from other known eubacteria identified specific structural motifs that may indicate possible features involved in species-specific responses to antibiotics. Thus, our novel insights provide unique structural tools for discriminating between pathogenic bacteria and the microbiome, namely the useful bacterial species within the human body, mainly in the gut flora.

Methods

***S. aureus* Growth and Cell Wall Disruption.** *S. aureus* strain RN4220 (American Type Culture Collection 35556) (71) was grown overnight at 37 °C. Cells were harvested at an OD₆₀₀ of ~1.5. The bacterial culture was centrifuged three times at 5,000 rpm using a GS-3 rotor at 4 °C for 15 min, after which the supernatant was discarded. The pellet was resuspended and centrifuged in a tabletop centrifuge at 4,000 rpm using a GS-3 rotor at 4 °C for 10 min. The supernatant was then discarded, and the wet pellets (cells) were weighed. To disrupt the cell membranes, the samples were resuspended in 10 mM

Tris-acetate buffer pH 8.0, 14 mM Mg-acetate, 1 M KCl, 1 mM DTT, and 50 $\mu\text{g}/\text{mL}$ Lysozymin, then incubated at 37 °C for 1 h and periodically inverted during incubation. The lysates were centrifuged at 16,000 rpm using a SS34 rotor, at 4 °C for 30 min to remove cell debris. The supernatants were incubated in 670 mM Tris-acetate buffer pH 8.0, 20 mM Mg-acetate, 7 mM DTT, 7 mM $\text{Na}_3\text{-phosphoenolpyruvate}$, 5.5 mM ATP, 70 mM of each amino acid, and 1.9 mg of pyruvatekinase at 37 °C for 30 min and then dialyzed overnight at 4 °C against dialysis in 10 mM Tris-acetate buffer pH 8.0, 14 mM Mg-acetate, 60 mM K-acetate and 1 mM DTT. The extract was then flash-frozen and stored at -80 °C until use.

S. aureus Ribosome Purification. Cell extract was layered on a 1.1 M sucrose cushion (72), $\text{H}_{10}\text{M}_{15}\text{N}_{100}\text{K}_{50}\beta\text{Me}_6$ pH 8.0 (10 mM HEPES pH 8.0, 15 mM MgCl_2 , 100 mM NH_4Cl , 50 mM KCl, and 6 mM $\beta\text{-mercaptoethanol}$), and ultracentrifuged twice, each time at 55,000 rpm using a Ti-70 rotor at 4 °C for 17 h. The supernatant was then discarded, and the pellet was dissolved in $\text{H}_{10}\text{M}_{15}\text{N}_{150}\beta\text{Me}_6$ buffer pH 8.0 at 4 °C. Ribosomal subunits were then separated by zonal ultracentrifugation, using a Ti-15 zonal rotor with a gradient of 8–40% sucrose, at low Mg^{2+} concentration (1 mM MgCl_2) for 17.5 h at 27,000 rpm. After separation, the Mg^{2+} concentration was adjusted to 10 mM, and the ribosomal subunits fractions were concentrated using sequential centrifugations. The samples were kept in $\text{H}_{10}\text{M}_{10}\text{N}_{60}\text{K}_{15}$ at pH 7.6, brought to a final concentration not exceeding 1,000 $\text{A}_{260} \text{ mL}^{-1}$, and then flash-frozen for storage at -80 °C.

Ribosome Activity Assay. *S. aureus* ribosome activity was determined in a bacterial-coupled transcription/translation assay system, which measures the expression of the luciferase gene (73).

Crystallization of the SA50S Subunit. Crystals of SA50S were obtained at 20 °C by the hanging-drop vapor diffusion technique. The crystallization drop contained 0.166% MPD, 0.333% EtOH, $\text{H}_{20}\text{M}_{10}\text{N}_{60}\text{K}_{15}$ (pH range, 6.8–7.8), 5 mM spermidine, 0.5 mM MnCl_2 , and 1–1.6 mg/mL SA50S subunits. The reservoir solution contained 15% of 1:2 ethanol-MPD and $\text{H}_{10}\text{M}_{10}\text{N}_{60}\text{K}_{15}$ buffer (pH range, 6.8–7.8). The SA50S subunits were heat-activated for 30 min at 37 °C before crystallization. These conditions usually yield ~60- to 300- μm hexagonal crystals, which appear as hexagons. High-resolution diffracting crystals were obtained by macro seeding, using crystals that had been fished out of the crystallization drop; washed in 10 μL of 7.5% EtOH, 7.5% MPD, $\text{H}_{10}\text{M}_{10}\text{N}_{60}\text{K}_{15}$ buffer, and 0.5 mM MnCl_2 ; and seeded for 24 h in a preequilibrated crystallization drop. SA50S crystals were kept in

stabilization solution consisting of 15% MPD, 15% EtOH, $\text{H}_{10}\text{M}_{10}\text{N}_{60}\text{K}_{15}$ buffer (pH range, 6.8–7.8), and 0.5 mM MnCl_2 .

Crystals of SA50S Complexes. For obtaining SA50S antibiotic complexes, SA50S crystals were soaked in solutions containing 11.4–22.7 $\mu\text{g}/\text{mL}$ BC-3205 in the stabilization solution for 3–6 h before flash-freezing and data collection.

Data Collection and Processing. Before data collection, the crystals were immersed in cryoprotectant solution of 20% MPD, 15% EtOH, $\text{H}_{10}\text{M}_{10}\text{N}_{60}\text{K}_{15}$ buffer, and 0.5 mM MnCl_2 . Crystallographic data were collected at the ID23-1, ID23-2, and ID-29 beamlines at the European Synchrotron Radiation Facility in Grenoble, France. X-ray diffraction data were collected from the hexagonal crystals at 100K. Up to 15 crystals were needed to yield complete datasets of SA50S using 0.1° oscillations. Data were processed with HKL2000 (74) and the CCP4 package suite (75).

Map Calculation, Model Building, and Refinement. The structures were determined using PHASER for molecular replacement, implemented in PHENIX, with the D50S structure (PDB ID code 2ZJR) used as a starting model. Once initial phases were obtained, rigid body and positional refinement were performed using Phenix.refine (76) and CNS (77). For R-free calculations during refinement cycles, a random 5% of the data were omitted during refinement cycles. Tracing the ribosomal RNA and remodeling the ribosomal proteins with *S. aureus* strain NCTC8325 sequence (78) according to the electron density maps was performed using Coot (79, 80), Rosetta ERRASER (81) was used to facilitate further building and to improve the quality of the rRNA geometry. Figures were generated using Pymol (82) and Chimera (83). Sequence alignments were performed using BLAST (84), and structure alignments were done using LSQMAN (85) and Coot.

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