

Resistance to Linezolid Caused by Modifications at Its Binding Site on the Ribosome

Long, Katherine S.; Vester, Birte

Published in: Antimicrobial Agents and Chemotherapy

Link to article, DOI: 10.1128/AAC.05702-11

Publication date: 2012

Document Version Publisher's PDF, also known as Version of record

Link back to DTU Orbit

Citation (APA): Long, K. S., & Vester, B. (2012). Resistance to Linezolid Caused by Modifications at Its Binding Site on the Ribosome. *Antimicrobial Agents and Chemotherapy*, *56*(2), 603-612. https://doi.org/10.1128/AAC.05702-11

General rights

Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

• Users may download and print one copy of any publication from the public portal for the purpose of private study or research.

- You may not further distribute the material or use it for any profit-making activity or commercial gain
- You may freely distribute the URL identifying the publication in the public portal

If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

Antimicrobial Agents and Chemotherapy

Resistance to Linezolid Caused by Modifications at Its Binding Site on the Ribosome

Katherine S. Long and Birte Vester Antimicrob. Agents Chemother. 2012, 56(2):603. DOI: 10.1128/AAC.05702-11. Published Ahead of Print 5 December 2011.

	Updated information and services can be found at: http://aac.asm.org/content/56/2/603
REFERENCES	<i>These include:</i> This article cites 103 articles, 70 of which can be accessed free at: http://aac.asm.org/content/56/2/603#ref-list-1
CONTENT ALERTS	Receive: RSS Feeds, eTOCs, free email alerts (when new articles cite this article), more»

Information about commercial reprint orders: http://aac.asm.org/site/misc/reprints.xhtml To subscribe to to another ASM Journal go to: http://journals.asm.org/site/subscriptions/

Journals.ASM.org



Resistance to Linezolid Caused by Modifications at Its Binding Site on the Ribosome

Katherine S. Long^a and Birte Vester^b

Department of Systems Biology and Novo Nordisk Foundation Center for Biosustainability, Technical University of Denmark, Lyngby, Denmark,^a and Department of Biochemistry and Molecular Biology, University of Southern Denmark, Odense M, Denmark^b

Linezolid is an oxazolidinone antibiotic in clinical use for the treatment of serious infections of resistant Gram-positive bacteria. It inhibits protein synthesis by binding to the peptidyl transferase center on the ribosome. Almost all known resistance mechanisms involve small alterations to the linezolid binding site, so this review will therefore focus on the various changes that can adversely affect drug binding and confer resistance. High-resolution structures of linezolid bound to the 50S ribosomal subunit show that it binds in a deep cleft that is surrounded by 23S rRNA nucleotides. Mutation of 23S rRNA has for some time been established as a linezolid resistance mechanism. Although ribosomal proteins L3 and L4 are located further away from the bound drug, mutations in specific regions of these proteins are increasingly being associated with linezolid resistance. However, very little evidence has been presented to confirm this. Furthermore, recent findings on the Cfr methyltransferase underscore the modification of 23S rRNA as a highly effective and transferable form of linezolid resistance. On a positive note, detailed knowl-edge of the linezolid binding site has facilitated the design of a new generation of oxazolidinones that show improved properties against the known resistance mechanisms.

LINEZOLID

inezolid (Fig. 1A) is a synthetic drug and a member of the oxazolidinone class of antibiotics. It acts as a protein synthesis inhibitor by binding to the ribosomal peptidyl transferase center (PTC) and stopping the growth of bacteria. Linezolid appeared on the market in 2000 for treatment of serious infections caused by Gram-positive bacteria resistant to other antibiotics, including streptococci, vancomycin-resistant enterococci (VRE), and methicillin-resistant Staphylococcus aureus (MRSA) (9). It is not well suited for fighting Gram-negative pathogenic bacteria that are intrinsically resistant due to efflux pumps that force linezolid out of the cell faster than it can accumulate (1,74). Linezolid is one of the few truly new antibiotics that have been introduced in many years, as most newcomers are derivatives of existing drugs. The development of new antibiotics is apparently not an attractive business despite the expanding problem of antibiotic resistance in pathogenic bacteria. At the introduction of linezolid, it was claimed that there would be no cross-resistance to linezolid and that resistance would be rare and difficult for the bacteria to develop (20, 103), but resistance does appear (reviewed in references 61 and 85). This minireview will present in detail how several types of modifications at the linezolid binding site on the ribosome can cause resistance to linezolid and other oxazolidinones. The only clear nonribosomal linezolid resistance mechanism reported is related to mutations causing increased expression of ABC transporter genes in Streptococcus pneumoniae (3, 19). It has also been shown that Staphylococcus aureus possesses a gene for a majorfacilitator-superfamily-type multidrug efflux pump named LmrS that is capable of extruding linezolid (21).

Although linezolid is currently the only oxazolidinone antibiotic approved for treatment of bacterial infections, other oxazolidinones have been investigated and have entered development. Some examples are eperezolid, which behaves similarly to linezolid (reviewed in reference 9) and posizolid (AZD2563) (98). Some newer derivatives are tedizolid phosphate (previously

known as torezolid phosphate [TR-701]), an oxazolidinone drug currently in phase III trials for acute bacterial skin and skin structure infections, including those caused by MRSA (http://www .clinicaltrials.gov/ct2/show/NCT01170221?term=tedizolid &rank=4) (79), and radezolid (RX-1741), which has completed two phase II clinical trials, one for uncomplicated skin and skinstructure infections and the other for community-acquired pneumonia (86). There is an obvious interest in developing a new generation of oxazolidinone derivatives which can overcome the known resistance mechanisms. A recent paper (52) focused on the structure-activity relationships (SAR) of oxazolidinones, including some new derivatives, using a panel of clinical and laboratoryderived S. aureus strains possessing ribosomal resistance mutations or the Cfr methyltransferase gene. The data show that improvement regarding the known resistance mechanisms is possible for oxazolidinone derivatives.

THE LINEZOLID BINDING SITE ON THE RIBOSOME

Although some of the first reports on linezolid claimed that it was a unique drug with a unique mechanism of action, we now know that the mechanism of action is similar to those of other antibiotics binding to the PTC. The first indications of the exact binding site of linezolid on the ribosome came in 1999 from studies of resistance mutations in 23S rRNA and pointed to the PTC (41). The PTC is in the middle of the 50S ribosomal subunit in the bottom of the cleft where the 3' ends of aminoacyl-tRNA and peptidyl-tRNA are positioned for peptide transfer (Fig. 1B and C). Binding of linezolid to this area was subsequently supported by other mutagenesis studies but could not be verified by antibiotic

Published ahead of print 5 December 2011 Address correspondence to Birte Vester, b.vester@bmb.sdu.dk. Copyright © 2012, American Society for Microbiology. All Rights Reserved. doi:10.1128/AAC.05702-11

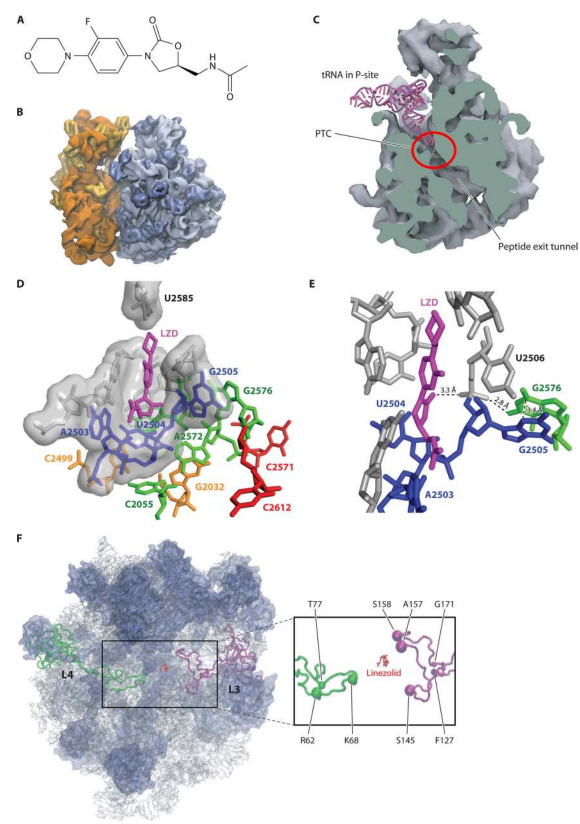


FIG 1 (A) The chemical structure of linezolid. (B) The *E. coli* 70S X-ray structure with 16S rRNA in orange, 30S ribosomal proteins in yellow, 23S rRNA in gray, and 50S ribosomal proteins in blue (PDB file from reference 75). (C) A cutaway view of the large ribosomal subunit with a red circle at the PTC (PDB file from reference 77). A P-site-bound tRNA is shown in magenta, and the PTC and peptide exit tunnel are indicated. (D and E) Close-up views of the linezolid binding site made with the PyMOL software program. The coloring of nucleotides indicates first-layer (blue), second-layer (green), third-layer (orange), and outer-layer (red) nucleotides with respect to linezolid. (D) The locations of mutated nucleotides with respect to bound linezolid. First-layer nucleotides are shown in surface representation. (E) The position of G2576 with respect to linezolid. (F) Illustration of how parts of the L3 (in purple) and L4 (in green) ribosomal proteins extend toward the PTC where linezolid (in red) is bound. Selected amino acids are marked with the corresponding *S. aureus* numbering. The coordinates are from PDB file 3DLL (95).

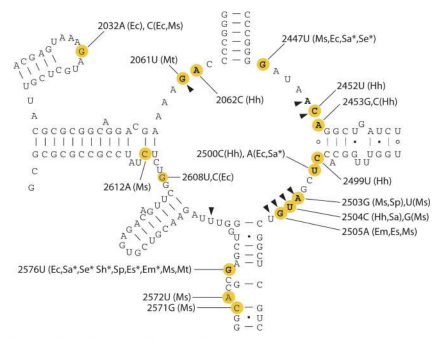


FIG 2 Secondary structure of the peptidyl transferase loop of domain V of 23S rRNA (*M. smegmatis* sequence in *E. coli* numbering). The nucleotides that form the linezolid binding pocket are indicated with black triangles. Nucleotide positions where mutations confer linezolid resistance are marked with yellow circles. The nucleotides where mutations have a significant effect on linezolid MIC (>4-fold MIC increase) are in bold type, whereas those where mutations have a small to moderate effect on the linezolid MIC (4-fold or less MIC increase) are in regular type. The mutations and corresponding organisms are indicated with two-letter abbreviations: Ec (*E. coli*), Sa (*S. aureus*), Se (*S. epidermidis*), Sh (*S. haemolyticus*), Sp (*S. pneumoniae*), Es (*E. faecalis*), Em (*E. faecium*), Ms (*M. smegmatis*), Mt (*M. tuberculosis*), and Hh (*H. halobium*). Asterisks indicate mutations found in clinical isolates. Only mutations where some evidence or strong indication of the mutation-resistance relationship has been published are marked on the figure (8, 19, 30, 41, 47, 50, 56, 59, 60, 62, 66, 69, 73, 97, 99, 100).

footprinting, a technique that has been used successfully to localize ribosomal binding sites for other antibiotics. Cross-linking studies with linezolid derivatives also pointed to a location at the PTC (12, 46). In 2008, this site was confirmed by crystal structures of linezolid bound to the 50S ribosomal subunit from the archaeon Haloarcula marismortui (36) and from the bacterium Deinococcus radiodurans (95). The structures reveal that linezolid binds to the A site of the PTC and interacts extensively with many 23S rRNA nucleotides in the neighborhood (Fig. 1D and E and Fig. 2). The PTC is almost exclusively composed of RNA, and the nucleotides in this region are phylogenetically highly conserved. The many crystal structures of 50S subunit-antibiotic complexes that have now been published also reveal that linezolid partially shares its binding site with other antibiotics that inhibit protein synthesis. Chloramphenicol, clindamycin, tiamulin, and streptogramin A (plus bruceantin, homoharringtonine, and anisomycin, which preferentially bind to eukaryotic and archaeal ribosomes) all occupy overlapping sites with linezolid at the A site of the PTC (original references and Protein Data Bank [PDB] identification [ID] are in references 10, 14, and 94), where the aminoacyl moiety of A-site tRNA has to be positioned for successful peptide transfer. This is underscored by the cross-resistance between linezolid and other drugs binding to this region. Thus, linezolid binds to a common antibiotic site on the ribosome and acts by blocking precise positioning of A-site tRNA in the PTC. The crystal structures have also shown that the nucleotides in the PTC adapt to the binding of antibiotics by slight changes in their relative positions, where especially nucleotide 2585 (Fig. 1D) seems very disposed to repositioning (95). The differences in modes of action of the antibiotics must be ascribed to their overlapping but not identical binding

sites and how they affect neighboring nucleotides in combination with their mode of access, affinity for the site, and association and dissociation rate constants.

Knowledge of the molecular details of linezolid binding to the PTC is highly advantageous for development of new oxazolidinone derivatives. It can indicate where there is room for drug derivatization, suggest sites for additional interactions, and thus facilitate the prediction of beneficial as well as detrimental interactions. Finally, the structures can be used to suggest drug modifications that will allow it to bind to the ribosome despite the presence of resistance determinants (22). This approach has been successfully exploited by the company Rib-X, who used knowledge about linezolid and sparsomycin binding to the 50S ribosomal subunit to create new drugs from parts of these antibiotics (86). They bridged and derivatized the two components, and one example is the radezolid mentioned above, where one ring of the linezolid (left side on Fig. 1A) is replaced and extended to obtain additional interaction in the PTC. Another example is presented by Trius Therapeutics, who modeled linezolid and tedizolid (previously known as torezolid [TR-700]) to the binding sites in the PTC with and without a methylation at the C-8 position of the adenine at position 2503 (52). This showed a better accommodation of TR-700 than linezolid in the presence of the methylation, which is the resistance determinant mediated by the Cfr methyltransferase (described below).

RESISTANCE CAUSED BY 23S rRNA MUTATIONS

The linezolid binding site at the PTC is composed entirely of RNA, and until recently, mutation of 23S rRNA was the only known linezolid resistance mechanism. The binding pocket is lined with

23S rRNA mutation ^a	D. radiodurans position ^b	LZD distance (Å) ^c	Origin ^d	Organism(s) ^e	Reference(s)
G2032A	G2015	9.3		Ec	99
G2032C	G2015	9.3		Ec, Ms	56, 99
G2061U	G2044	3.0	S	Mt	30
A2062C	A2045	8.1	S	Hh	41
G2447U	G2426	6.2	C, S	Ec, Ms, Sa, Se	50, 66, 73, 97, 99
C2452U	C2431	2.9	S	Hh	41
A2453G	A2432	6.1	S	Hh	41
A2453C	A2432	6.1	S	Hh	41
C2499U	C2478	8.1	S	Hh	41
U2500A	U2479	7.2	C, S	Ec, Sa	62, 66
U2500C	U2479	7.2	S	Hh	41
A2503G	A2482	2.2	S	Ms, Sp	19, 56
A2503U	A2482	2.2		Ms	47
U2504C	U2483	4.5	S	Hh, Sa	41, 50
U2504G	U2483	4.5		Ms	56
G2505A	G2484	2.4	S	Em, Es, Ms	8, 56, 69
C2571G	C2550	14.7		Ms	56
A2572U	A2551	6.6		Ms	56
G2576U	G2555	7.9	C, S	Ec, Em, Es, Ms, Mt, Sa, Se, Sh, Sp	19, 30, 50, 56, 59, 60, 66, 69, 97
G2608U	G2588	15.9		Ec	100
G2608C	G2588	15.9		Ec	100
C2612A	C2591	17.7		Ms	56

TABLE 1 23S rRNA linezolid resistance mutations and corresponding nucleotide-linezolid distances

^a The nucleotide positions of the mutations are listed according to *E. coli* numbering.

^b The corresponding nucleotide positions in D. radiodurans 23S rRNA.

^c The closest approximate distances are given between the corresponding *D. radiodurans* nucleotides and linezolid (LZD) from the *D. radiodurans* 50S-linezolid complex X-ray structure (PDB ID 3DLL) (95).

^d Indicates whether a mutation was obtained via *in vitro* selection (S) or found in clinical isolates (C).

^e Abbreviations: Ec, E. coli; Sa, S. aureus; Se, S. epidermidis; Sh, S. haemolyticus; Sp, S. pneumoniae; Es, E. faecalis; Em, E. faecium; Ms, M. smegmatis; Mt, M. tuberculosis; Hh, H. halobium.

the universally conserved nucleotides G2061, A2451, C2452, A2503, U2504, G2505, U2506, and U2585, which interact directly with linezolid (95) (Fig. 1D and 2). Strains selected for linezolid resistance have mutations in 23S rRNA nucleotides G2061, C2452, A2503, U2504, and G2505, which abut the bound linezolid molecule (Fig. 1D and E and Table 1), but also at nucleotides that are located more distally, such as A2062, G2447, A2453, C2499, U2500, and G2576 (11, 19, 30, 41, 50, 51, 66, 69, 73) (Fig. 1D and E and Table 1). The pattern of resistance is organism specific in that the obtained mutation sites differ, with only little overlap, between Halobacterium halobium, Escherichia coli, Enterococcus faecalis, Enterococcus faecium, Mycobacterium smegmatis, Mycobacterium tuberculosis, S. aureus, and S. pneumoniae (Fig. 2 shows detailed information). This specificity is likely related to singlebase identity differences in the RNA nucleotides that are more distant from and do not directly abut the bound linezolid molecule and variations in the fitness costs associated with specific mutations in different organisms. The degree of resistance conferred by the mutations is not a simple function of the nucleotidelinezolid distance, and mutations at distal nucleotides that do not interact with linezolid directly, in particular, G2576U (19, 30, 56, 66, 69, 97) (Fig. 1D and E) and G2447U (66, 73, 97, 99), confer significant resistance (Table 1).

Although the effect of a few engineered 23S rRNA mutations on linezolid susceptibility has been investigated in *E. coli* by expressing the mutated rRNA on a multicopy plasmid (99), the multiplicity of rRNA operons and heterogeneous ribosome populations complicates the study of rRNA mutations in many bacteria.

Additional mutations have been investigated in M. smegmatis strains modified to contain a single rRNA operon (31, 32, 47, 57), where one model system allows for the introduction of mutations regardless of whether the mutations confer antibiotic resistance or not (32). This genetic system has been used to test whether rRNA mutations that confer linezolid resistance in other organisms (G2576U) also do so in M. smegmatis and to validate rRNA mutations associated with linezolid resistance in the literature without genetic proof (A2503G, G2505A, C2571G, and C2612A) (56). The study revealed that mutations at some distance from an antibiotic binding site can confer resistance and that the same rRNA mutation can have significantly different effects in different bacteria. Moreover, the system has been used to introduce the double rRNA mutations G2032A-C2499A, G2032A-U2504G, C2055A-U2504G, and C2055A-A2572U and show that there are strong synergistic effects on linezolid resistance relative to the effects of the corresponding single rRNA mutations (56).

The most frequently reported mutation in linezolid-resistant clinical isolates is 23S RNA G2576U. This mutation has been reported in both staphylococci and enterococci (6, 15, 34, 40, 59, 61, 70, 71, 87, 97, 101), and a clear correlation between the number of mutated rRNA operons and the linezolid MIC has been found (2, 35, 59). Most reports of the G2576U mutation in clinical isolates are associated with some form of increased or prolonged linezolid treatment or with high linezolid usage at the local hospital (29, 84), underscoring the importance of judicious use of linezolid in clinical settings. Although some studies have documented reversion of the G2576U mutation in the absence of linezolid selection

L3: 1	MTKGILGRKI	GMTQVFGENG	ELIPVTVVEA	KENVVLQKKT	VEVDGYNAIQ
51	VGFEDKKAYK	KDAKSNKYAN	KPAEGHAKKA	DAAPKRFIRE	FRNVDVDAYE
101	VGQEVSVDTF	VAGDVIDVTG	VSKGKGFQGA	IKRHGQSRGP	MSHGSHFHRA
151	PGSVGMASDA	SRVFKGQKMP	GRMGGNTVTV	QNLEVVQVDT	ENKVILVKGN
201	VPGPKKGLVE	IRTSIKKGNK			
L4: 1	MANYDVLKLD	GTKSGSIELS	DAVFGIEPNN	SVLFEAINLQ	RASLRQGTHA
51	VKNRSAVSGG	GRKPWKQKGT	GRARQGTIRA	PQWRGGGIVF	GPTPRSYAYK
101	MPKKMRRLAL	RSALSFKVOE	NGLTVVDAFN	FEAPKTKEFK	NVLSTLEOPK

151 KVLVVTENED VNVELSARNI PGVQVTTAQG LNVLDITNAD SLVITEAAAK

201 KVEEVLG FIG 3 The sequences of 50S ribosomal proteins L3 and L4 from *Staphylococcus aureus* MRSA252. The amino acids highlighted in purple and green correspond to those shown as balls in Fig. 1F. For the L3 sequence, A157 is the

equivalent of E. coli L3 N149.

(62, 87), a recent report showed that the G2576U mutation was retained in a *Staphylococcus haemolyticus* isolate even after 30 passages on antibiotic-free medium (60).

The U2500A (62) and G2447U (97) 23S RNA mutations have been reported in linezolid-resistant clinical isolates of staphylococci, and these mutations have been shown to confer linezolid resistance in *in vitro*-selected mutants of *E. coli* and/or *M. smegmatis* (66, 73). Although the U2504A mutation has been reported in clinical staphylococcal isolates (16, 48, 97), the only U2504 mutation isolated from *in vitro* selection with linezolid to date is U2504C (41, 50). Additional mutated positions of 23S rRNA at G2603U (49, 78, 83) and C2534U (97) have been reported in clinical isolates with reduced linezolid susceptibility, but a direct relationship between these mutations and linezolid resistance is unclear.

LINEZOLID RESISTANCE AND A TENTATIVE RELATIONSHIP TO MUTATIONS IN RIBOSOMAL PROTEINS L3 AND L4

Another type of linezolid resistance determinant has received attention recently, namely, mutations in the ribosomal L3 protein. The main part of ribosomal protein L3 is positioned on the surface of the 50S subunit, but a loop ending in two tips extends into the PTC (Fig. 1F and Fig. 3). Bacterial L3 mutations have, since 2003, been associated with resistance to linezolid, tiamulin/valnemulin, and anisomycin, which all bind to overlapping sites at the PTC. The first L3 resistance mutation in bacteria was detected by us in *E. coli* by selection with tiamulin, and its role in resistance was verified by genetic evidence (7). Many studies have since associated L3 mutations with linezolid resistance in *S. aureus, Staphylococcus cohnii*, and *Staphylococcus epidermidis*, and the data are summarized in Table 2.

Locke and colleagues (54) performed a selection study with both methicillin-sensitive and MRSA strains and the oxazolidinones linezolid and TR-700. They obtained various mutations in 23S RNA (see above) as well as mutations in L3 and L4 (Table 2), and apparently only the 23S RNA mutations give high resistance. As the authors note, directed mutagenesis and heterologous expression studies are needed to conclusively link these mutations to oxazolidinone resistance. Nevertheless, the positions of the mutations are correlated with the linezolid binding site in an X-ray structure and a possible mechanism of structural perturbations is explained, which, together with the absence of mutations in other remote peptide regions and in L22 (also sequenced), points to a role in resistance development.

A search for similar mutations by the same authors in clinical staphylococci revealed a linezolid-resistant *S. aureus* strain with an

L3 mutation and a linezolid-resistant S. epidermidis strain with an L3 mutation plus the G2447U 23S rRNA mutation (53). Another study identified L3 mutations in the region near the PTC in linezolid-resistant S. aureus strains from a hospital outbreak in Spain that also harbored the cfr gene (55). Similarly, sequencing of selected genes from three clinical isolates from Mexico, with cfr and linezolid resistance, also revealed L3 and L4 mutations (63) (Table 2), but the effects of the mutations were not confirmed. Two of these strains harbored an L101V L3 mutation that has also been detected in a sensitive control strain and is therefore not considered relevant for linezolid resistance. Ten S. epidermidis strains from Italy with reduced linezolid susceptibility were investigated by sequencing the 23S rRNA gene and ribosomal protein L3, L4, and L22 genes, as well as looking for the *cfr* gene (65). It was concluded that the L3 mutations F147L and/or A157R appear to be responsible for the elevated linezolid MIC values, as adjacent alterations have been associated with resistance in other strains. Another recent study (43) reported multiple L3 mutations, but in most cases, the mutations were together with the 23S rRNA mutation G2576T, which is known to confer linezolid resistance, and with an L4 insertion. Finally, two new L3 mutations have also been found in linezolid-resistant S. aureus strains from a cystic fibrosis patient after prolonged drug treatment, where one strain also possessed the G2576U 23S rRNA mutation (15). It is worth mentioning that mutations of L3 at some of the positions mentioned above, as well as at amino acids close by, are associated with resistance to the pleuromutilins retapamulin and tiamulin (24, 42, 43, 66). As linezolid and the pleuromutilins binds at overlapping sites at the PTC, these findings support the relationship between L3 mutations and PTC antibiotic resistance in general.

Part of ribosomal protein L4 is also placed relatively close to the PTC (Fig. 1F) but in the tunnel through which nascent peptides exit the ribosome. A surveillance study found a slightly linezolidresistant S. pneumoniae isolate with a six-nucleotide deletion in the L4 gene (Δ W65-R66) but with no genetic proof presented (18). Another surveillance study identified an S. pneumoniae strain with the same deletion plus a strain with a neighboring six-nucleotide deletion (Δ K68-G69) in the L4 gene. The deletions caused a slightly reduced susceptibility to linezolid (MIC change from 1 μ g/ml to 4 μ g/ml) as evidenced by transformations and were associated with a fitness cost (96). The amino acid deletions are in the same region as mutations known to be involved in macrolide resistance (26), and as macrolide antibiotics bind to a site neighboring but not directly overlapping the linezolid binding site, the effect of the deletions is probably caused by an allosteric mechanism.

Four different mutations were found in the L4 ribosomal protein in another study of linezolid-resistant *S. epidermidis* isolates (97), but two of these mutations occurred in isolates that also harbored 23S rRNA mutations. The N158S mutation has been found among linezolid-sensitive *S. epidermidis* isolates and is therefore probably a clonal marker rather than a resistance mutation (97). Also, it has been concluded that L4 K68R is not responsible for resistance in an *S. epidermidis* study, as it did not increase resistance when present together with other mutations (65). In contrast, the L4 mutation K68Q found in the *S. aureus* selection study mentioned above was assumed to play a role in resistance (54), as was an L4 gene mutation in *Clostridium perfringens* (33). The L4 mutations related to linezolid resistance are summarized in Table 2, but the data presented do not present a consistent

Mutation	Organism	D. radiodurans L3 ^a	LZD distance ^b	Remarks ^c	Reference
L3					
ΔF127-H146	S. aureus	T113-K132			54
G139R	S. aureus	G125	>25	С, Т, 2576Т	15
Δ S145	S. aureus	S131	>25	С	53
Δ S145/H146Y	S. aureus	S131/K132	>25/19.3	C, cfr	55
H146R/M156T	Staphylococcus	K132/G143	19.3/15.2	C, T, 2215A, 2576T, —, d L4-ins70G	43
H146Q/V154L/A157R	Staphylococcus	K132/I141/R144	19.3/22.1/7.0	C, T, $-,^d$ L4-ins70G, $-e^e$	43
F147I	Staphylococcus	K133 or W134	22.4 or >25	C, T, 2215A, 2576T, —, d L4-ins70G	43
F147L/A157R	S. epidermidis	K133 or W134/R144	22.4 or >25/7.0	C, —, ^d —, ^e L4-K68R/ ^e	65
G152D	S. aureus	G139	20.4	2447T	54
G152D	S. aureus	G139	20.4	С, Т	15
G155R	S. aureus	G142	17.5		54
G155R/M169L	S. aureus	G142/M156	17.5/>25		54
A157R	S. epidermidis	R144	7.0	С, 2447Т	53
S158F/D159Y	S. epidermidis	K145/T146	13.3/5.4	C, cfr, L3-L101V	63
S158Y/D159Y	S. cohnii	K145/T146	13.3/5.4	C, cfr, L4-N20S/A133T/V155I	63
Δ M169-G174	S. aureus	DM156-G161		C, cfr	55
L4					
N20S/A133T/V155I	S. cohnii	f		C, cfr, L3-S158F/D159Y	63
Δ W65-R66	S. pneumoniae	Y59-G60	24.2-21.6		18
Δ W65-R66	S. pneumoniae	Y59-G60	24.2-21.6	C, RE	96
Δ K68-G69	S. pneumoniae	K62-Q63	11.2-15.1	C, RE	96
K68N	S. epidermidis	K62	11.2	2576T	97
K68Q	S. aureus	K62	11.2		54
G71D	Clostridium perfringens	G65	15.6		33
ins71GGR72	S. epidermidis	G65/N66	15.6/16.1	2576T	97
L108S/ins71GGR72	S. epidermidis	L102	>75	2534T, — ^e	97

TABLE 2 Mutations in L3 and L4 that have been associated with linezolid resistance in staphylococci (and one case in *Clostridium perfringens*) and corresponding amino acid-linezolid distances^g

^a The L3 sequence from the *D. radiodurans* 50S X-ray structure (95) used for distance determination has been aligned with L3 from *S. aureus*, and the amino acids corresponding to the mutations are listed.

^b The distance is the smallest distance from linezolid (LZD) to the corresponding D. radiodurans amino acid.

^c Selected additional information. C, from clinical isolates; T, treatment with linezolid; cfr, containing the *cfr* gene; RE, resistance evidence. Additional resistance determinants are shown (xxxxN refers to 23S RNA positions corresponding to *E. coli* 23S RNA, and L3- and L4- indicate additional mutations). See the original references for more details. ^d —, L3-L101V, which is not expected to influence linezolid resistance.

e -, L4-N158S, which is not expected to influence linezolid resistance.

^f No data for alignment were available.

^g Mutations with indications that they are merely strain markers and not relevant for antibiotic resistance are not included, except where they are found together with other mutations and are then posted under Remarks. The resistance effects vary from very minor effects to substantial resistance, and only in a few cases has there been direct evidence linking the mutations to resistance. The slash is used when there is more than one mutation. "K133 or W134" indicates that the alignment did not reveal which of these two positions corresponds to F147.

pattern and it is not definitively established which changes contribute directly to linezolid resistance.

As sequencing of genes from bacteria exhibiting resistance becomes common, many mutations are revealed, but they might not all be related to the observed resistance. Some mutations could be just random changes without significant effect, while others are likely related to either causing the resistance or accommodating or sustaining resistance in a synergistic way. This is supported by the fact that several of the L3 and L4 mutations observed in relation to linezolid resistance are found together with mutations in 23S rRNA at or near the PTC as mentioned above. Similar possible synergistic effects have also been reported for other PTC antibiotics in other bacteria such as M. smegmatis (56) and Brachyspira spp. (28, 68), indicating an interplay between multiple mutations in relation to resistance, accommodation of mutations, and fitness cost. More specific information about the effects of the single and combined mutations is needed to clarify their detailed interactions.

RESISTANCE CAUSED BY ALTERATIONS IN 23S rRNA MODIFICATION

It is well established that RNA modifications placed at or near an antibiotic binding site can affect drug binding to the ribosome. rRNA is intrinsically modified with methyl groups and pseudouridine residues, mediated by methyltransferase and pseudouridine synthase enzymes, respectively. The modifications are clustered at functional centers on the ribosome and have a collective importance in optimizing different steps of protein synthesis, although the exact roles of each modification are not well described. Modifications can also be acquired, but to date, the only type of modification that provides acquired antibiotic resistance is methylation. Thus, resistance generally occurs either by the inactivation of an indigenous methyltransferase or by the acquisition of an antibiotic resistance methyltransferase.

Several housekeeping modifications at the peptidyl transferase center are known to affect linezolid susceptibility. The pseudouridylation of 23S rRNA nucleotide 2504 confers resistance to linezolid, clindamycin, and tiamulin, suggesting that this modification may have evolved as an intrinsic resistance mechanism to protect bacteria from PTC antibiotics (88). Inactivation of the spr0333 methyltransferase targeting G2445 in 23S rRNA results in decreased susceptibility to linezolid in *S. pneumoniae* (3, 19). Likewise, mutations inactivating the methyltransferase RlmN, which methylates the C-2 position of A2503 of 23S rRNA, also lower linezolid susceptibility in *S. aureus* (23, 44, 88).

The only known transferable form of linezolid resistance is conferred by the multiresistance gene cfr, which encodes an rRNA methyltransferase (39). Cfr adds a methyl group at the C-8 position of 23S rRNA nucleotide A2503 (25), the only example of this modification found in natural RNAs so far. The methylation confers combined resistance to five different classes of antibiotics that bind at overlapping nonidentical sites at the PTC (58). The resistance is substantial and functions in both Gram-positive and Gram-negative bacteria. The phenotype is called $PhLOPS_A$, for resistance to phenicols, lincosamides, oxazolidinones, pleuromutilins, and streptogramin A antibiotics. In addition, Cfr confers significant resistance to selected 16-membered ring macrolide antibiotics such as josamycin and spiramycin, but not tylosin (81). The methylation is positioned in the PTC, and its direct interference with drug binding is supported by X-ray structures of linezolid bound to the D. radiodurans and H. marismortui 50S subunits (36, 95).

Cfr is related to the RlmN methyltransferases, which add a methyl group at the C-2 position of 23S rRNA nucleotide A2503 (90). Although the primary activity of Cfr is C-8 methylation, analysis of a $cfr^+ \Delta rlmN$ strain by mass spectrometry showed that Cfr has a secondary C-2 methylation activity, which is the primary activity of RlmN (25). Phylogenetic analysis of the RlmN/Cfr family of proteins suggests that the RlmN subfamily is the ancestral subfamily and that Cfr likely arose through duplication and horizontal gene transfer (37). Both RlmN and Cfr are radical S-adenosylmethionine (SAM) enzymes, a superfamily that catalyzes a diverse set of reactions that involve cleavage of unreactive C-H bonds by a 5'-deoxyadenosyl radical generated by reductive cleavage of SAM (82, 92). A recent crystal structure of the E. coli RlmN enzyme (4) and studies on the reaction mechanisms of RlmN and Cfr show that the methyl group is not transferred directly from SAM to the RNA but rather through a two-step sequence involving intermediate methylation of a conserved cysteine residue in the C-terminal domain (27).

The cfr gene was originally discovered on multiresistance plasmids isolated during surveillance studies of florfenicol resistance in Staphylococcus isolates of animal origin (38, 76). The first cfrpositive clinical strain of methicillin-resistant S. aureus was isolated in 2005, and it had cfr on the chromosome together with the ermB gene on a transposable genetic element (89). This strain is notable because these two rRNA methyltransferase genes are located on the same operon and their coexpression confers resistance to all clinically relevant antibiotics that target the large ribosomal subunit (81). A number of staphylococcal clinical isolates containing cfr in different genetic contexts and parts of the world have subsequently been reported (5, 6, 16, 17, 65, 67, 72, 80). In some instances, a connection between the resistant isolates and prior linezolid treatment (6, 64) or extensive use of linezolid (72) can be documented. In addition to staphylococci, the cfr gene has also been identified in two Bacillus strains isolated from swine feces (13, 102). Moreover, a recent report detects cfr in an isolate of *Proteus vulgaris*, a naturally occurring Gram-negative bacterium of pigs. The gene is found in a region with homology to a staphy-lococcal plasmid that is flanked by two IS26 elements and inserted into the chromosomal *fimD* gene (93). The presence of *cfr* on mobile genetic elements such as plasmids and transposons in different geographical locations strongly suggests that it can be disseminated within the microbial community and spread among pathogenic bacteria.

DOES LINEZOLID RESISTANCE COME WITH A COST AND HOW IS IT RELATED TO CROSS-RESISTANCE?

The maintenance and spread of resistance genes are directly related to their fitness cost. Expression of Cfr in a laboratory strain has only a small effect on growth rate. Competition experiments involving wild-type and inactivated Cfr indicate that the small fitness cost is due not to the C-8 methylation but rather to expression of the protein (45). Similar experiments with strains coexpressing the Erm methyltransferase showed that dimethylation of 23S rRNA position A2058 increases the fitness cost of Cfrmediated methylation of A2503 (45). The low fitness cost of cfr is troubling, as it suggests that cells can maintain the gene even in the absence of antibiotic selection. The available data on the fitness cost of RlmN and the effect of RlmN-mediated methylation on linezolid susceptibility are confusing and contradictory. It is concluded from one study that an RlmN-deletion strain is outcompeted by wild-type cells in the absence of antibiotic selection and that RlmN methylation leads to a 2-fold decrease in linezolid susceptibility in S. aureus (90). In a follow-up study, the authors concede that the earlier MIC data are inconclusive (44). Subsequent competition experiments show that cells with an inactivated *rlmN* gene outcompeted wild-type cells under linezolid selection (44). This result is corroborated by a study of clinical S. aureus strains (23) and has implications for resistance development in patients undergoing prolonged therapy with linezolid.

The fitness cost of ribosomal mutations varies enormously from one position to another and is also dependent on the specific organism. Some bacteria accept mutations at positions in 23S rRNA that are lethal in others. As mentioned above, a resistance mutation may be accompanied by other mutations that compensate for deleterious effects or act synergistically to enhance resistance (3). A decrease in growth rates for 23S rRNA mutations at the PTC is expected because many of the nucleotides are phylogenetically conserved and are considered functionally important. Among the engineered single mutations in *M. smegmatis* that have the most significant effects on linezolid resistance, the mutations lead to either moderate (A2503G/U and G2447U) or large (U2504G and G2576U) decreases in growth rate, where the G2576U mutation has the largest effect and results in 3-foldslower growth (47, 56, 57). This is consistent with the fact that although both the G2447U and G2576U mutations lead to 32-fold increases in linezolid MIC values, only the G2447U mutation was isolated by selection in the presence of linezolid (56, 73). Engineered double mutations in M. smegmatis (G2032A-C2499A, G2032A-U2504G, C2055A-U2504G, and C2055A-A2572U) led to significantly slower growth compared to the corresponding single mutations, suggesting that multiple changes together have detrimental effects in this functionally important region of the ribosome (56). The G2576U mutation has been studied extensively in S. aureus, where a progressive decrease in growth rate is observed with each additional copy of the mutation (2). However, the ability of the mutation to persist in one copy in the absence of antibiotic selection and the rapid reemergence of multiple mutated copies upon reexposure to linezolid suggest that a single copy has a minimal fitness cost (91).

Cross-resistance between PTC antibiotics resulting from 23S rRNA mutations is not uncommon, but there is no straightforward relationship between overlapping binding sites and crossresistance. Although the different sets of specific mutations, bacteria, and antibiotics examined in the literature preclude a detailed analysis, some patterns have emerged. There is a correlation between linezolid and chloramphenicol resistance for the single G2447U, A2503G, U2504G, G2505A, and G2576U mutations in M. smegmatis (56, 57). However, this correlation does not hold for the G2032A-U2504G and C2055A-U2504G double mutations, which both include the U2504G mutation (56). In the same studies, no relationship between linezolid, clindamycin, and valnemulin resistance could be observed (56, 57). Cross-resistance between linezolid and tiamulin has been documented for the G2447U and U2500A mutations in E. coli and the G2576U mutation in E. coli and S. aureus (66). The complexity of cross-resistance patterns between PTC antibiotics is likely due to the unique set of interactions that each bound antibiotic makes with the PTC cavity.

ACKNOWLEDGMENTS

We thank Jacob Poehlsgaard for preparation of Fig. 1B and C and Can Ali Tomruk for Fig. 1F.

REFERENCES

- Barbachyn MR, Ford CW. 2003. Oxazolidinone structure-activity relationships leading to linezolid. Angew. Chem. Int. Ed. Engl. 42: 2010–2023.
- Besier S, Ludwig A, Zander J, Brade V, Wichelhaus TA. 2008. Linezolid resistance in *Staphylococcus aureus*: gene dosage effect, stability, fitness costs, and cross-resistances. Antimicrob. Agents Chemother. 52: 1570–1572.
- Billal DS, Feng J, Leprohon P, Legare D, Ouellette M. 2011. Whole genome analysis of linezolid resistance in *Streptococcus pneumoniae* reveals resistance and compensatory mutations. BMC Genomics 12:512.
- 4. Boal AK, et al. 2011. Structural basis for methyl transfer by a radical SAM enzyme. Science 332:1089–1092.
- Bongiorno D, et al. 2010. DNA methylase modifications and other linezolid resistance mutations in coagulase-negative staphylococci in Italy. J. Antimicrob. Chemother. 65:2336–2340.
- Bonilla H, et al. 2010. Multicity outbreak of linezolid-resistant *Staphylococcus epidermidis* associated with clonal spread of a *cfr*-containing strain. Clin. Infect. Dis. 51:796–800.
- 7. Bosling J, Poulsen SM, Vester B, Long KS. 2003. Resistance to the peptidyl transferase inhibitor tiamulin caused by mutation of ribosomal protein L3. Antimicrob. Agents Chemother. 47:2892–2896.
- Bourgeois-Nicolaos N, et al. 2007. Dose dependence of emergence of resistance to linezolid in *Enterococcus faecalis* in vivo. J. Infect. Dis. 195: 1480–1488.
- Brickner SJ, Barbachyn MR, Hutchinson DK, Manninen PR. 2008. Linezolid (ZYVOX), the first member of a completely new class of antibacterial agents for treatment of serious gram-positive infections. J. Med. Chem. 51:1981–1990.
- Bulkley D, Innis CA, Blaha G, Steitz TA. 2010. Revisiting the structures of several antibiotics bound to the bacterial ribosome. Proc. Natl. Acad. Sci. U. S. A. 107:17158–17163.
- 11. Carsenti-Dellamonica H, et al. 2005. In vitro selection of mutants of *Streptococcus pneumoniae* resistant to macrolides and linezolid: relationship with susceptibility to penicillin G or macrolides. J. Antimicrob. Chemother. 56:633–642.
- Colca JR, et al. 2003. Cross-linking in the living cell locates the site of action of oxazolidinone antibiotics. J. Biol. Chem. 278:21972–21979.
- 13. Dai L, et al. 2010. First report of the multidrug resistance gene cfr and the

phenicol resistance gene *fexA* in a Bacillus strain from swine feces. Antimicrob. Agents Chemother. **54**:3953–3955.

- Dunkle JA, Xiong L, Mankin AS, Cate JH. 2010. Structures of the *Escherichia coli* ribosome with antibiotics bound near the peptidyl transferase center explain spectra of drug action. Proc. Natl. Acad. Sci. U. S. A. 107:17152–17157.
- Endimiani A, et al. 2011. Emergence of linezolid-resistant *Staphylococcus aureus* after prolonged treatment of cystic fibrosis patients in Cleveland, Ohio. Antimicrob. Agents Chemother. 55:1684–1692.
- Farrell DJ, Mendes RE, Ross JE, Jones RN. 2009. Linezolid surveillance program results for 2008 (LEADER Program for 2008). Diagn. Microbiol. Infect. Dis. 65:392–403.
- Farrell DJ, Mendes RE, Ross JE, Sader HS, Jones RN. 2011. LEADER program results for 2009: an activity and spectrum analysis of linezolid using 6,414 clinical isolates from 56 medical centers in the United States. Antimicrob. Agents Chemother. 55:3684–3690.
- Farrell DJ, Morrissey I, Bakker S, Buckridge S, Felmingham D. 2004. In vitro activities of telithromycin, linezolid, and quinupristindalfopristin against *Streptococcus pneumoniae* with macrolide resistance due to ribosomal mutations. Antimicrob. Agents Chemother. 48: 3169–3171.
- Feng J, et al. 2009. Genome sequencing of linezolid-resistant *Streptococcus pneumoniae* mutants reveals novel mechanisms of resistance. Genome Res. 19:1214–1223.
- Fines M, Leclercq R. 2000. Activity of linezolid against Gram-positive cocci possessing genes conferring resistance to protein synthesis inhibitors. J. Antimicrob. Chemother. 45:797–802.
- Floyd JL, Smith KP, Kumar SH, Floyd JT, Varela MF. 2010. LmrS is a multidrug efflux pump of the major facilitator superfamily from *Staphylococcus aureus*. Antimicrob. Agents Chemother. 54:5406–5412.
- 22. Franceschi F, Duffy EM. 2006. Structure-based drug design meets the ribosome. Biochem. Pharmacol. 71:1016–1025.
- Gao W, et al. 2010. Two novel point mutations in clinical *Staphylococcus aureus* reduce linezolid susceptibility and switch on the stringent response to promote persistent infection. PLoS Pathog. 6:e1000944.
- 24. Gentry DR, Rittenhouse SF, McCloskey L, Holmes DJ. 2007. Stepwise exposure of *Staphylococcus aureus* to pleuromutilins is associated with stepwise acquisition of mutations in *rplC* and minimally affects susceptibility to retapamulin. Antimicrob. Agents Chemother. 51:2048–2052.
- Giessing AM, et al. 2009. Identification of 8-methyladenosine as the modification catalyzed by the radical SAM methyltransferase Cfr that confers antibiotic resistance in bacteria. RNA 15:327–336.
- 26. **Gregory ST, Dahlberg AE.** 1999. Erythromycin resistance mutations in ribosomal proteins L22 and L4 perturb the higher order structure of 23 S ribosomal RNA. J. Mol. Biol. **289**:827–834.
- 27. Grove TL, et al. 2011. A radically different mechanism for S-adenosylmethionine-dependent methyltransferases. Science 332: 604-607.
- Hidalgo A, et al. 2011. Trends towards lower antimicrobial susceptibility and characterization of acquired resistance among clinical isolates of *Brachyspira hyodysenteriae* in Spain. Antimicrob. Agents Chemother. 55: 3330–3337.
- Hill RL, et al. 2010. Linezolid-resistant ST36 methicillin-resistant *Staphylococcus aureus* associated with prolonged linezolid treatment in two paediatric cystic fibrosis patients. J. Antimicrob. Chemother. 65: 442–445.
- Hillemann D, Rusch-Gerdes S, Richter E. 2008. In vitro-selected linezolid-resistant *Mycobacterium tuberculosis* mutants. Antimicrob. Agents Chemother. 52:800–801.
- 31. Hobbie SN, et al. 2006. A genetic model to investigate drug-target interactions at the ribosomal decoding site. Biochimie 88:1033–1043.
- Hobbie SN, et al. 2007. Engineering the rRNA decoding site of eukaryotic cytosolic ribosomes in bacteria. Nucleic Acids Res. 35:6086–6093.
- 33. Holzel CS, Harms KS, Schwaiger K, Bauer J. 2010. Resistance to linezolid in a porcine *Clostridium perfringens* strain carrying a mutation in the *rplD* gene encoding the ribosomal protein L4. Antimicrob. Agents Chemother. 54:1351–1353.
- Hong T, Li X, Wang J, Sloan C, Cicogna C. 2007. Sequential linezolidresistant *Staphylococcus epidermidis* isolates with G2576T mutation. J. Clin. Microbiol. 45:3277–3280.
- Ikeda-Dantsuji Y, et al. 2011. Linezolid-resistant Staphylococcus aureus isolated from 2006 through 2008 at six hospitals in Japan. J. Infect. Chemother. 17:45–51.

- 36. Ippolito JA, et al. 2008. Crystal structure of the oxazolidinone antibiotic linezolid bound to the 50S ribosomal subunit. J. Med. Chem. 51: 3353-3356.
- 37. Kaminska KH, et al. 2010. Insights into the structure, function and evolution of the radical-SAM 23S rRNA methyltransferase Cfr that confers antibiotic resistance in bacteria. Nucleic Acids Res. 38:1652-1663.
- 38. Kehrenberg C, Schwarz S. 2006. Distribution of florfenicol resistance genes fexA and cfr among chloramphenicol-resistant Staphylococcus isolates. Antimicrob. Agents Chemother. 50:1156-1163.
- 39. Kehrenberg C, Schwarz S, Jacobsen L, Hansen LH, Vester B. 2005. A new mechanism for chloramphenicol, florfenicol and clindamycin resistance: methylation of 23S ribosomal RNA at A2503. Mol. Microbiol. 57:1064-1073.
- 40. Kelly S, et al. 2008. An outbreak of colonization with linezolid-resistant Staphylococcus epidermidis in an intensive therapy unit. J. Antimicrob. Chemother. 61:901-907.
- 41. Kloss P, Xiong L, Shinabarger DL, Mankin AS. 1999. Resistance mutations in 23 S rRNA identify the site of action of the protein synthesis inhibitor linezolid in the ribosomal peptidyl transferase center. J. Mol. Biol. 294:93-101.
- 42. Kosowska-Shick K, et al. 2006. Single- and multistep resistance selection studies on the activity of retapamulin compared to other agents against Staphylococcus aureus and Streptococcus pyogenes. Antimicrob. Agents Chemother. 50:765–769.
- 43. Kosowska-Shick K, Julian KG, McGhee PL, Appelbaum PC, Whitener CJ. 2010. Molecular and epidemiologic characteristics of linezolidresistant coagulase-negative staphylococci at a tertiary care hospital. Diagn. Microbiol. Infect. Dis. 68:34-39.
- 44. LaMarre JM, Howden BP, Mankin AS. 2011. Inactivation of the indigenous methyltransferase RlmN in Staphylococcus aureus increases linezolid resistance. Antimicrob. Agents Chemother. 55:2989-2991.
- 45. LaMarre JM, Locke JB, Shaw KJ, Mankin AS. 2011. Low fitness cost of the multidrug resistance gene cfr. Antimicrob. Agents Chemother. 55: 3714-3719.
- 46. Leach KL, et al. 2007. The site of action of oxazolidinone antibiotics in living bacteria and in human mitochondria. Mol. Cell 26:393-402.
- 47. Li BB, Wu CM, Wang Y, Shen JZ. 2011. Single and dual mutations at positions 2058, 2503 and 2504 of 23S rRNA and their relationship to resistance to antibiotics that target the large ribosomal subunit. J. Antimicrob. Chemother. 66:1983-1986.
- 48. Liakopoulos A, et al. 2009. A T2504A mutation in the 23S rRNA gene responsible for high-level resistance to linezolid of Staphylococcus epidermidis, I. Antimicrob, Chemother, 64:206-207.
- 49. Lincopan N, de Almeida LM, Elmor de Araujo MR, Mamizuka EM. 2009. Linezolid resistance in Staphylococcus epidermidis associated with a G2603T mutation in the 23S rRNA gene. Int. J. Antimicrob. Agents 34: 281-282.
- 50. Livermore DM, Warner M, Mushtaq S, North S, Woodford N. 2007. In vitro activity of the oxazolidinone RWJ-416457 against linezolidresistant and -susceptible staphylococci and enterococci. Antimicrob. Agents Chemother. 51:1112–1114.
- 51. Lobritz M, Hutton-Thomas R, Marshall S, Rice LB. 2003. Recombination proficiency influences frequency and locus of mutational resistance to linezolid in Enterococcus faecalis. Antimicrob. Agents Chemother. 47:3318-3320.
- 52. Locke JB, et al. 2010. Structure-activity relationships of diverse oxazolidinones for linezolid-resistant Staphylococcus aureus strains possessing the cfr methyltransferase gene or ribosomal mutations. Antimicrob. Agents Chemother. 54:5337–5343.
- 53. Locke JB, Hilgers M, Shaw KJ. 2009. Mutations in ribosomal protein L3 are associated with oxazolidinone resistance in staphylococci of clinical origin. Antimicrob. Agents Chemother. 53:5275-5278.
- 54. Locke JB, Hilgers M, Shaw KJ. 2009. Novel ribosomal mutations in Staphylococcus aureus strains identified through selection with the oxazolidinones linezolid and torezolid (TR-700). Antimicrob. Agents Chemother. 53:5265-5274
- 55. Locke JB, et al. 2010. Elevated linezolid resistance in clinical cfr-positive Staphylococcus aureus isolates is associated with co-occurring mutations in ribosomal protein L3. Antimicrob. Agents Chemother. 54:5352-5355.
- 56. Long KS, et al. 2010. Mutations in 23S rRNA at the peptidyl transferase center and their relationship to linezolid binding and cross-resistance. Antimicrob. Agents Chemother. 54:4705-4713.
- 57. Long KS, et al. 2009. Single 23S rRNA mutations at the ribosomal

peptidyl transferase centre confer resistance to valnemulin and other antibiotics in Mycobacterium smegmatis by perturbation of the drug binding pocket. Mol. Microbiol. 71:1218-1227.

- 58. Long KS, Poehlsgaard J, Kehrenberg C, Schwarz S, Vester B. 2006. The Cfr rRNA methyltransferase confers resistance to phenicols, lincosamides, oxazolidinones, pleuromutilins, and streptogramin A antibiotics. Antimicrob. Agents Chemother. 50:2500-2505.
- 59. Marshall SH, Donskey CJ, Hutton-Thomas R, Salata RA, Rice LB. 2002. Gene dosage and linezolid resistance in Enterococcus faecium and Enterococcus faecalis. Antimicrob. Agents Chemother. 46:3334-3336.
- 60. Mazzariol A, et al. 27 July 2011. Outbreak of linezolid-resistant Staphylococcus haemolyticus in an Italian intensive care unit. Eur. J. Clin. Microbiol. Infect. Dis. [Epub ahead of print.] doi:10.1007/s10096-011-1343-6.
- 61. Meka VG, Gold HS. 2004. Antimicrobial resistance to linezolid. Clin. Infect. Dis. 39:1010-1015.
- 62. Meka VG, et al. 2004. Linezolid resistance in sequential Staphylococcus aureus isolates associated with a T2500A mutation in the 23S rRNA gene and loss of a single copy of rRNA. J. Infect. Dis. 190:311-317.
- 63. Mendes RE, et al. 2010. First report of Staphylococcal clinical isolates in Mexico with linezolid resistance caused by cfr: evidence of in vivo cfr mobilization. J. Clin. Microbiol. 48:3041-3043.
- 64. Mendes RE, et al. 2008. First report of cfr-mediated resistance to linezolid in human staphylococcal clinical isolates recovered in the United States. Antimicrob. Agents Chemother. 52:2244-2246.
- 65. Mendes RE, et al. 2010. Assessment of linezolid resistance mechanisms among Staphylococcus epidermidis causing bacteraemia in Rome, Italy. J. Antimicrob. Chemother. 65:2329-2335.
- 66. Miller K, Dunsmore CJ, Fishwick CW, Chopra I. 2008. Linezolid and tiamulin cross-resistance in Staphylococcus aureus mediated by point mutations in the peptidyl transferase center. Antimicrob. Agents Chemother. 52:1737-1742.
- 67. Morales G, et al. 2010. Resistance to linezolid is mediated by the cfr gene in the first report of an outbreak of linezolid-resistant Staphylococcus aureus. Clin. Infect. Dis. 50:821-825.
- 68. Pringle M, Poehlsgaard J, Vester B, Long KS. 2004. Mutations in ribosomal protein L3 and 23S ribosomal RNA at the peptidyl transferase centre are associated with reduced susceptibility to tiamulin in Brachyspira spp. isolates. Mol. Microbiol. 54:1295–1306.
- 69. Prystowsky J, et al. 2001. Resistance to linezolid: characterization of mutations in rRNA and comparison of their occurrences in vancomycinresistant enterococci. Antimicrob. Agents Chemother. 45:2154-2156.
- 70. Raad II, et al. 2004. Clinical-use-associated decrease in susceptibility of vancomycin-resistant Enterococcus faecium to linezolid: a comparison with quinupristin-dalfopristin. Antimicrob. Agents Chemother. 48: 3583-3585.
- 71. Roberts SM, et al. 2006. Linezolid-resistant Staphylococcus aureus in two pediatric patients receiving low-dose linezolid therapy. Pediatr. Infect. Dis. J. 25:562-564.
- 72. Sanchez Garcia M, et al. 2010. Clinical outbreak of linezolid-resistant Staphylococcus aureus in an intensive care unit. JAMA 303:2260–2264.
- 73. Sander P, et al. 2002. Ribosomal and non-ribosomal resistance to oxazolidinones: species-specific idiosyncrasy of ribosomal alterations. Mol. Microbiol. 46:1295-1304.
- 74. Schumacher A, et al. 2007. Intracellular accumulation of linezolid in Escherichia coli, Citrobacter freundii and Enterobacter aerogenes: role of enhanced efflux pump activity and inactivation. J. Antimicrob. Chemother. 59:1261-1264.
- 75. Schuwirth BS, et al. 2005. Structures of the bacterial ribosome at 3.5 A resolution. Science 310:827-834.
- 76. Schwarz S, Werckenthin C, Kehrenberg C. 2000. Identification of a plasmid-borne chloramphenicol-florfenicol resistance gene in Staphylococcus sciuri. Antimicrob. Agents Chemother. 44:2530–2533.
- 77. Selmer M, et al. 2006. Structure of the 70S ribosome complexed with mRNA and tRNA. Science 313:1935-1942.
- 78. Seral C, et al. 2011. Nosocomial outbreak of methicillin- and linezolidresistant Staphylococcus epidermidis associated with catheter-related infections in intensive care unit patients. Int. J. Med. Microbiol. 301:354-358.
- 79. Shaw KJ, et al. 2008. In vitro activity of TR-700, the antibacterial moiety of the prodrug TR-701, against linezolid-resistant strains. Antimicrob. Agents Chemother. 52:4442-4447.
- 80. Shore AC, et al. 2010. Identification and characterization of the multidrug resistance gene cfr in a Panton-Valentine leukocidin-positive se-

quence type 8 methicillin-resistant *Staphylococcus aureus* IVa (USA300) isolate. Antimicrob. Agents Chemother. **54**:4978–4984.

- Smith LK, Mankin AS. 2008. Transcriptional and translational control of the *mlr* operon, which confers resistance to seven classes of protein synthesis inhibitors. Antimicrob. Agents Chemother. 52:1703–1712.
- 82. Sofia HJ, Chen G, Hetzler BG, Reyes-Spindola JF, Miller NE. 2001. Radical SAM, a novel protein superfamily linking unresolved steps in familiar biosynthetic pathways with radical mechanisms: functional characterization using new analysis and information visualization methods. Nucleic Acids Res. 29:1097–1106.
- Sorlozano A, et al. 2010. Detection of new mutations conferring resistance to linezolid in glycopeptide-intermediate susceptibility Staphylococcus hominis subspecies hominis circulating in an intensive care unit. Eur. J. Clin. Microbiol. Infect. Dis. 29:73–80.
- Souli M, et al. 2009. Colonisation with vancomycin- and linezolidresistant *Enterococcus faecium* in a university hospital: molecular epidemiology and risk factor analysis. Int. J. Antimicrob. Agents 33:137–142.
- Stefani S, Bongiorno D, Mongelli G, Campanile F. 2010. Linezolid resistance in staphylococci. Pharmaceuticals 3:1988–2006.
- Steitz TA. 2010. From the structure and function of the ribosome to new antibiotics (Nobel lecture). Angew. Chem. Int. Ed. Engl. 49:4381–4398.
- Swoboda S, et al. 2005. Varying linezolid susceptibility of vancomycinresistant *Enterococcus faecium* isolates during therapy: a case report. J. Antimicrob. Chemother. 56:787–789.
- Toh SM, Mankin AS. 2008. An indigenous posttranscriptional modification in the ribosomal peptidyl transferase center confers resistance to an array of protein synthesis inhibitors. J. Mol. Biol. 380:593–597.
- 89. Toh SM, et al. 2007. Acquisition of a natural resistance gene renders a clinical strain of methicillin-resistant *Staphylococcus aureus* resistant to the synthetic antibiotic linezolid. Mol. Microbiol. **64**:1506–1514.
- 90. Toh SM, Xiong L, Bae T, Mankin AS. 2008. The methyltransferase YfgB/RlmN is responsible for modification of adenosine 2503 in 23S rRNA. RNA 14:98–106.
- Tsakris A, et al. 2007. Persistence of rRNA operon mutated copies and rapid re-emergence of linezolid resistance in *Staphylococcus aureus*. J. Antimicrob. Chemother. 60:649–651.

- 92. Wang SC, Frey PA. 2007. S-adenosylmethionine as an oxidant: the radical SAM superfamily. Trends Biochem. Sci. 32:101–110.
- 93. Wang Y, et al. 2011. Detection of the staphylococcal multiresistance gene *cfr* in *Proteus vulgaris* of food animal origin. J. Antimicrob. Chemother. **66**:2521–2526.
- Wilson DN. 2009. The A-Z of bacterial translation inhibitors. Crit. Rev. Biochem. Mol. Biol. 44:393–433.
- Wilson DN, et al. 2008. The oxazolidinone antibiotics perturb the ribosomal peptidyl-transferase center and effect tRNA positioning. Proc. Natl. Acad. Sci. U. S. A. 105:13339–13344.
- Wolter N, et al. 2005. Novel mechanism of resistance to oxazolidinones, macrolides, and chloramphenicol in ribosomal protein L4 of the pneumococcus. Antimicrob. Agents Chemother. 49:3554–3557.
- 97. Wong A, et al. 2010. Polyphyletic emergence of linezolid-resistant staphylococci in the United States. Antimicrob. Agents Chemother. 54: 742–748.
- Wookey A, et al. 2004. AZD2563, a novel oxazolidinone: definition of antibacterial spectrum, assessment of bactericidal potential and the impact of miscellaneous factors on activity in vitro. Clin. Microbiol. Infect. 10:247–254.
- Xiong L, et al. 2000. Oxazolidinone resistance mutations in 23S rRNA of Escherichia coli reveal the central region of domain V as the primary site of drug action. J. Bacteriol. 182:5325–5331.
- Xu J, et al. 2005. Protected nucleotide G2608 in 23S rRNA confers resistance to oxazolidinones in *E. coli*. Biochem. Biophys. Res. Commun. 328:471–476.
- Yoshida K, et al. 2009. Linezolid-resistant methicillin-resistant *Staphylococcus aureus* isolated after long-term, repeated use of linezolid. J. Infect. Chemother. 15:417–419.
- 102. Zhang WJ, et al. 2011. The new genetic environment of *cfr* on plasmid pBS-02 in a *Bacillus* strain. J. Antimicrob. Chemother. 66: 1174–1175.
- Zurenko GE, et al. 1996. In vitro activities of U-100592 and U-100766, novel oxazolidinone antibacterial agents. Antimicrob. Agents Chemother. 40:839-845.