A novel pleuromutilin antibacterial compound, its binding mode and selectivity mechanism

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The increasing appearance of pathogenic bacteria with antibiotic resistance is a global threat. Consequently, clinically available potent antibiotics that are active against multidrug resistant pathogens are becoming exceedingly scarce. Ribosomes are a main target for antibiotics, and hence are an objective for novel drug development. Lefamulin, a semi-synthetic pleuromutilin compound highly active against multi-resistant pathogens, is a promising antibiotic currently in phase III trials for the treatment of community-acquired bacterial pneumonia in adults. The crystal structure of the Staphylococcus aureus large ribosomal subunit in complex with lefamulin reveals its protein synthesis inhibition mechanism and the rationale for its potency. In addition, analysis of the bacterial and eukaryotes ribosome structures around the pleuromutilin binding pocket has elucidated the key for the drug’s selectivity.

Staphylococcus aureus is one of the leading causes of hospital and community associated infections such as skin and soft-tissue infections, sepsis, pneumonia and toxic shock syndrome1,2. These infections are treated by a variety of antibiotic classes including those targeting the large ribosomal subunit such as oxazolidinones, pleuromutilins, macrolides, lincosamides and ketolides. S. aureus possess 5–6 copies of ribosomal RNA (rRNA), operons, and hence acquisition of single nucleotide mutations in the 23S rRNA, which are among the common resistance mechanisms with respect to ribosomal active antibiotics, should have appeared after relatively long periods3. Nevertheless the epidemiology of S. aureus is increasing1.

Among other mechanisms, resistance in S. aureus is associated with post-transcription modifications in rRNA and mutations in ribosomal proteins (rProtein). In many cases methicillin-resistant Staphylococcus aureus (MRSA) is also resistant to other antibacterial agents such as aminoglycosides, macrolides, lincosamides, tetracyclines, cephalosporins, carbapenems, beta-lactamase inhibitor combinations, trimethoprim and sulphonamides4. The prevalence of MRSA strains has created an urgent demand for novel therapeutic agents effective against it.

The pleuromutilins, discovered as S. aureus inhibitors in 1951, are produced by the genus Pleurotus5. These natural products consist of a tricyclic muti-lin core, a C21 keto group and various C14 extensions. It was shown that pleuromutilin derivatives mediate inhibition of bacterial protein biosynthesis by binding to the peptidyl transferase center (PTC) at the A- and P-site6–8. Tiamulin was the first family member that was approved for veterinary use in 1979, followed by valnemulin in 1999. Retapamulin is a topically administered pleuromutilin that was approved for human use in 2007.

In the crystal structures of various pleuromutilins complexes with the large ribosomal subunit of Dienococcus radiodurans (D50S)9,10, Haloarcula marismortui (H50S)11 and Staphylococcus aureus (SA50S)12 the tricyclic muti-lin core is blocking the A-site and its 4-amino-2-hydroxy-cyclohexyl-sulfanyl-acetyl C14 extension is pointing into the P-site, thus perturbing tRNA accommodation at the A- and P-site.

Known ribosomal resistance mechanisms to pleuromutilins include post-transcription modification in the 23S ribosomal RNA by the Cfr enzyme which methylates the C8 position of the PTC nucleotide A2503 (E. coli rRNA numbering is used throughout)13. In addition, resistance occurs by mutations in the ribosomal protein (rProtein) uL3 G144-D159 loop and uL4 G69 residue (S. aureus rProtein numbering is used throughout)14,15, which is located in proximity to the PTC.

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Lefamulin (also known as BC-3781) developed by Nabriva Therapeutics, Vienna, Austria, is a highly active semi-synthetic pleuromutilin compound against pathogens that are commonly associated with community-acquired bacterial pneumonia (CABP), including multidrug resistant *S. pneumoniae*, *S. aureus*, and *M. pneumoniae*. Lefamulin has been evaluated in a phase II trial, which demonstrated the first proof of concept for the systemic use of a pleuromutilin antibiotic for the treatment of acute bacterial skin and skin structure infection (ABSSSI). Currently, it is in phase III development for treatment of community-acquired bacterial pneumonia (CABP).

### Results

**Structure of the *S. aureus* 50S in complex with lefamulin.** We determined the crystal structure of the large ribosomal subunit of *S. aureus* with lefamulin (SA50S-lef) (Table 1), from which we could readily identify the lefamulin molecule. An electron density map around lefamulin is shown in Fig. 1.

In the SA50S-lef crystal structure, lefamulin is bound at the PTC (Fig. 1), which is in agreement with other pleuromutilin complexes of D50S, H50S, and SA50S. In the available ribosome-pleuromutilin complex structures, all pleuromutilins bind to ribosomes at the same pocket and form two hydrogen bonds between the drug's acetyl carbonyl with the NH and NH2 of G2061 and between the hydroxyl group with the phosphate group of G2505 (Fig. 1D). In the SA50S-lef complex, U2585 is stacked with the lefamulin's C14 extension, which is a non-planar cyclohexane moiety. This interaction is stabilized by a U-U-carbonyl N3 symmetric interaction between U2585 and U2506 that, upon binding, are shifted from the binding pocket (Fig. 2A). Interactions between these two nucleotides have been identified previously in several D50S pleuromutilin complexes (Fig. 2C). A unique hydrogen bond between 23S rRNA nucleotides and lefamulin is formed between NH2 group of lefamulin's C14 extension and the O2 of the A2062 ribose. All of the other interactions of lefamulin with the rRNA nucleotides C2063, U2506, A2503, U2504, G2505, A2453, C2452, A2425 and C2424 are either hydrophobic or based on Van der Waals forces.

In the SA50S-lef complex structure, the conformation adopted by the flexible nucleotide U2585 is different from that in the structures: apo-SA50S, SA50S-BC3205, H50S-tiamulin, and D50S-pleuromutilin (Fig. 2). Lefamulin seems to make the same number of hydrogen bonds with the PTC nucleotides as BC-3205. Stabilization of lefamulin in its binding pocket is achieved by U2585 and U2506 U-U interactions. This agrees with the IC50 values that were determined in *S. aureus* cell free in vitro transcription-translation assay for BC-3205, lefamulin and tiamulin: IC50 = 0.02 μg/ml of lefamulin, IC50 = 0.08 μg/ml of BC-3205, IC50 = 0.10 μg/ml of tiamulin (Fig. 3) Moreover, inhibition of in vitro transcription-translation show lower IC50 values for lefamulin than for other known pleuromutilins acting on *S. aureus*.

**Structural basis for pleuromutilins selectivity.** Selectivity, namely the distinction between bacterial pathogens and eukaryotes, is crucial for the clinical use of antibiotics. Ribosome inhibiting antibiotics, which are currently in clinical use, target bacterial ribosomes without hampering the activity of the eukaryote's ribosome.
By comparing the SA50S structure with archaeal and eukaryotic ribosome structures, namely those of Haloarcula marismortui 50S, Saccharomyces cerevisiae 80S, Tetrahymena thermophila 60S, Leishmania donovani 60S and human (Homo sapiens) 80S the previously suggested mechanism by which A-site cleft antibiotic selectivity could be extended.

As already suggested, in all known bacterial ribosome with pleuromutilins, nucleotides C2452 and U2504 are part of the pleuromutilin binding pocket. Nucleotides U2504 and C2452 hydrophobically interact with the tricyclic mutilin core. These nucleotides interact with each other by non-Watson-Crick C:U 4-carbonyl-amino N3-N3 interactions. In contrast, the respective nucleotides in eukaryotic and archaeal ribosomes of the same sequence identity are positioned in different orientations thus cannot interact with each other (Fig. 4A). As described previously, in archaeal and eukaryotic ribosomes, nucleotides U2504 forms pi stacking interaction to A2055 (in archaea and eukaryotes) that is pointing away from the PTC binding site (Fig. 4A and B). This interaction pushes U2504 out of the binding pocket that opens up, thus pleuromutilins binding is hampered.

Additional observations indicate that for bacterial ribosome, the PTC 2nd shell base-pair A2543-U2500, is an important determinant for the bacterial drug binding. This base-pair keeps the pleuromutilin binding pocket in a "closed" form by stacking to C2452 and U2504 which interact with the tricyclic mutilin core. These nucleotides interact with each other by non-Watson-Crick C:U 4-carbonyl-amino N3-N3 interactions. In contrast, the respective nucleotides in eukaryotic and archaeal ribosomes of the same sequence identity are positioned in different orientations thus cannot interact with each other (Fig. 4A). As described previously, in archaeal and eukaryotic ribosomes, nucleotides U2504 forms pi stacking interaction to A2055 (in archaea and eukaryotes) that is pointing away from the PTC binding site (Fig. 4A and B). This interaction pushes U2504 out of the binding pocket that opens up, thus pleuromutilins binding is hampered.

The double layer nucleotides around the pleuromutilin tricyclic mutilin core create a chemically and structurally different binding pocket in prokaryotes vs. eukaryotes and archaea. The tricyclic mutilin core, which is the common moiety for all the pleuromutilins, is used as an anchor for the drug binding. Although the tricyclic mutilin core doesn’t form any hydrogen bonds with the PTC nucleotides, it is stabilized by hydrophobic and Van der Waals interactions. Compared to the pocket part that binds the mutilin core in eukaryotes and archaea, in eubacteria this part of the pocket is more rigid and chemically stabilized by the C:U interaction in the 1st shell stacked to the A:U base pair in the 2nd shell. Also, since in eukaryotes and archaea, the walls of the binding pocket are open, the tricyclic moiety cannot be properly anchored and consequently no binding or inhibition by pleuromutilins can be stabilized. In this way selectivity towards prokaryotes is achieved.
Figure 2. Comparative analysis of the lefamulin binding pocket. (A) A superposition of the PTC in apo SA50S structure (blue) and in the SA50S-lef complex (lefamulin in green 23S rRNA in orange) reveals relocations of nucleotides U2585 and U2506 in the bound vs. apo structure. (B) A superposition of the PTC in the two S. aureus pleuromutilins complexes [SA50S-BC3205 (teal and pink) (4WFB) and SA50S-lef (orange and green) - this work. The dash line represents the unique hydrogen bond BC-3205 forms with the 23S rRNA. (C) A superposition of pleuromutilins within their binding pockets of S. aureus and D. radiodurans [SA50S-lef (orange and green) - this work, D50S-retapamulin (dark grey) (2OGO)\textsuperscript{10}, D50S-tiamulin (yellow) (1XBP)\textsuperscript{9}].
Figure 3. Inhibition of bacterial protein synthesis. The inhibitory effect of BC-3205, lefamulin and tiamulin on protein expression in \(S.\) \(aureus\) in vitro transcription-translation cell-free system. The activity of the reporter protein (luciferase) in the presence of various concentrations of BC-3205, lefamulin and tiamulin is shown as arbitrary unit of luminescence \([\text{a.u.}]\). The IC_{50} values calculated by the plotted data showed better inhibition of protein synthesis by lefamulin than by BC-3205 and tiamulin.

Figure 4. The binding pocket of lefamulin. (A) SA50S-lef (lefamulin in green and 23S rRNA in orange) superimposed on the structure of human 80S (grey) (4UG0). In S. \(aureus\) nucleotides C2452 and U2504 interact with each other (marked with dashed lines) while the same nucleotides in eukaryotes' structures have the same identity but different orientations, so they cannot interact with each other. Nucleotides A2543 and U2500 are paired (marked with dashed lines) and stacked to C2452 and U2504 in the bacterial ribosomes. In eukaryotes' structures, nucleotides 2453 and 2500 are both uridines and no interaction between them occurs owing to their orientation. In eukaryotes, U2504 has different orientation than in bacteria which allows pi stacking to A2055 (C in bacteria, A in eukaryotes). This interaction stabilizes the “open” conformation of the PTC which is the key for the pleuromutilins' selectivity. (B) A superposition of the rRNA H50S-tiamulin structure (light blue) (3G4S) on that of SA50S-lef (orange). In archaea, nucleotides A2543 and U2500 cannot form Watson-Crick base-pair due to the stacking between U2504 and A2055. (C) A superposition of the rRNA apo SA50S structure (blue) (4WCE) on the SA50S-lef (orange) shows that the position of these nucleotides is the same in the bound and the unbound 50S. (D) Multiple sequence alignment of part of the pleuromutilin binding site, comparing the bacterial, archaeal and eukaryotes large ribosomal subunit rRNA. Helix 89 nucleotides, which are part of the PTC, are marked in a box. Arrows are indicating on the nucleotides C2452, A/U2453, U2504 and U2500 which are 1\(^{st}\) and 2\(^{nd}\) shell from the mutilin moiety. The lines represent Watson-crick base pairs, dashed lines are base pairs only in bacteria.
Discussion
The binding of lefamulin to the S. aureus large ribosomal subunit seems to be tighter than those produced by other pleuromutilins. Its binding triggers an induced fit rearrangement similar to that observed for other pleuromutilins, but in SA50S-lef complex an additional specific U:U interaction is formed. This interaction serves as a physical barrier that maintains a rather tight binding pocket conformation and might be the reason for the lefamulin higher potency over BC-3205 in S. aureus cell free in vitro transcription-translation assay.

Even though the PTC rRNA nucleotides are conserved across all kingdoms of life, the pleuromutilin family targets selectively bacterial ribosome, without inhibiting eukaryotic ribosomes. This selectivity originates from sequence differences. In Eukaryotic ribosome, the 2nd shell nucleotides are arranged such that they induce a structurally different conformation for the tricyclic mutilin binding site, namely a more “open” conformation, hence hampering the pleuromutilins binding and facilitating their clinical usage. Our current and previous studies revealed the factors governing the inhibition and highlighted additional aspects of pleuromutilins selectivity, and thus may extend the design of additional derivatives of these potent antibacterial drugs for the treatment against multi-drug resistant bacterial strains.

Materials and Methods

S. aureus growth and cell wall disruption. S. aureus strain RN4220 (American Type Culture Collection 35556)38 was grown and disrupted as described previously.12

Ribosome Purification, Crystallization, and Compound Soaking Experiments. Ribosomes were purified as described previously.12 SA50S was crystallized at 20 °C by the hanging-drop vapor diffusion technique. The crystallization drop contained 0.166% MPD, 0.333% EtOH, 20 mM Hepes pH range 6.8–7.8, 10 mM MgCl2, 60 mM NH4Cl, 15 mM KCl, 5 mM spermidine, 0.5 mM MnCl2, and 1–1.6 mg/ml SA50S subunits. The reservoir solution contained 15% of 1:2 ethanol-MPD and 110 mM Hepes pH range 6.8–7.8, 10 mM MgCl2, 60 mM NH4Cl, 15 mM KCl (pH range, 6.8–7.8) as previously described.12 For obtaining SA50S-lefamulin complex, SA50S crystals were soaked in stabilization solutions containing 44.9 μM (22.8 μg/ml) lefamulin in the stabilization solution for 6 hrs prior to flash freezing and data collection.

Data collection and processing. Before data collection the crystals were immersed in cryoprotectant solution of 20% MPD, 15% EtOH, 110 mM Hepes pH range 6.8–7.8, 10 mM MgCl2, 60 mM NH4Cl, 15 mM KCl and 0.5 mM MnCl2. Crystallographic data were collected at the ID23-1 beamlines, at the European Synchrotron Radiation Facility (ESRF), Grenoble, France. X-ray diffraction data were collected from the hexagonal crystals at 100 K. Up to 15 crystals were needed for yielding complete datasets of SA50S using 0.1° oscillations. Data were processed with HKL200029 and CCP4 package suite30.

Map calculation, model building and refinement. The apo-SA50S structure (PDB code: 4WCE) was used as a starting model for rigid body and positional refinement as implemented in PHENIX31. Densities for the antibiotics were located using a standard difference Fourier maps. For R-free calculations during refinement cycles, random 5% of the data were omitted during refinement cycles. Modelling of the ribosomal RNA and the ribosomal proteins according to the electron density maps was performed using Coot32,33, Rosetta ERRASER34 was used to facilitate further building and to improve the quality of the rRNA geometry. Figures were generated using Pyml35 and Chimera36. Sequence alignments were performed using BLAST37 Structure alignments were done using LSQMAN38 and Coot.

Sequence alignment. Sequence alignment was done with MAFFT version 739 and its figure was done by Jalview software.60

Ribosome Inhibition assay. The inhibition effect of BC-3205 and lefamulin on S. aureus ribosomes was tested in a bacterial coupled transcription/translation assay system, in the presence of increasing concentrations of the compounds, which measures the expression of the luciferase gene.41 The luciferase gene was inserted into plasmid with T7 RNA polymerase promoter. The reaction mixture contained: 160 mM Hepes-KOH (pH 7.5), 6.5% PEG 8 K, 0.074 mg/ml tyrosine, 1.3 mM ATP, 0.86 mM CTP, GTP and UTP, 208 mM potassium glutamate, 83 mM creatine phosphate, 28 mM NH4OAc, 0.663 mM cAMP, 1.8 mM DTT, 0.036 mg/ml folinic acid, 0.174 mg/ml E. coli tRNA mix, 1 mM amino acid, 0.25 mg/ml creatine kinase, 0.027 mg/ml T7 RNA polymerase, ribosome free E. coli cell free extract, 300 nM of S. aureus ribosomes, 0.003 μg/μl luciferase plasmid and pleuromutilin compound diluted to 0.46 mM to 0.98 μM (233 μg/ml to 0.05 ng/ml). The reaction mixture was incubated at 37 °C for 1 hr and terminated by the addition of erythromycin at a final concentration of 8 μg/ml. To quantify the reaction's products, Luciferin Assay Reagent (LAR, Promega) at 5:3 (luciferase: reaction mix) volume ratio was added to the mixture and luminescence was measured. The results were plotted and IC50 values were calculated with the program GraFit.42

Data deposition. The atomic coordinates and structure factors have been deposited in the Protein Data Bank, www.rcsb.org (PDB ID code: 5HL7).

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Acknowledgements
We thank Dr. Moshe Peretz, Shoshana Tel- Or, Miriam Lachever and Maggie Kessler for their interest and experimental support; Astrid Gruss for S. aureus ribosome preparation; staff members at Beamlines ID23-1 of the European Synchrotron Radiation Facility/European Molecular Biology Laboratory for their assistance during data collection. Funding was provided by European Research Council Grants 322581 (NOVRIB) and the Kimmel Center for Macromolecular Assemblies. M.K. is a recipient of an Adams Fellowship. A.Y. holds the Martin S. and Helen Kimmel Professorial Chair at the Weizmann Institute of Science.

Author Contributions
A.B. and A.Y. designed research; Z.E., D.M., M.K., E.Z., H.R., and A.B. performed research; S.P. and R.R. contributed new reagents/analytic tools; Z.E., D.M., M.K., E.Z., H.R., and A.B. analyzed data; and Z.E., D.M., A.B., and A.Y. wrote the paper.

Additional Information
Competing financial interests: This study was performed in collaboration with Nabriva Therapeutics AG (Vienna, Austria and King of Prussia, PA, United States). Dr. Paukner and Dr. Riedl are employees of Nabriva Therapeutics and own Nabriva stocks.

How to cite this article: Eyal, Z. et al. A novel pleuromutilin antibacterial compound, its binding mode and selectivity mechanism. Sci. Rep. 6, 39004; doi: 10.1038/srep39004 (2016).

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