Structures of Triacetyloleandomycin and Mycalamide A Bind to the Large Ribosomal Subunit of *Haloarcula marismortui*[∇]

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Structures have been obtained for the complexes that triacetyloleandomycin and mycalamide A form with the large ribosomal subunit of *Haloarcula marismortui*. Triacetyloleandomycin binds in the nascent peptide tunnel and inhibits the activity of ribosomes by blocking the growth of the nascent peptide chain. Mycalamide A binds to the E site and inhibits protein synthesis by occupying the space normally occupied by the CCA end of E-site-bound tRNAs.

Since 2001, crystal structures have been reported for the complexes that ribosomes form with scores of antiribosomal antibiotics and other inhibitors of protein synthesis (3, 4, 11-14, 20-22, 26, 27, 32-34). Many of them have been obtained using the large ribosomal subunit from Haloarcula marismortui (5, 11-14, 26, 27, 32), which forms crystals that diffract to higher resolution than do the ribosomal crystals obtained from any other organism thus far. Although H. marismortui, which is an archaeal halophile, is not pathogenic, the structures that it forms with protein synthesis inhibitors generally explain why those inhibitors block protein synthesis and are consistent with the relevant genetic and biochemical data, almost all of which derive from experiments done with either eukaryotes or eubacteria. Furthermore, there is ample evidence that protein synthesis inhibitors bind to the ribosomes from *H. marismortui* at the same sites where and in the same manner that they bind to the ribosomes from pathogenic organisms but (often) with different affinity (32). Thus, inhibitor structures obtained with H. marismortui have provided important insights into the mechanisms of action of many clinically important compounds, as well as the structural basis of both their species specificities and the source of ribosomal drug resistance. Here we present crystal structures of the complexes that the large ribosomal subunit of *H. marismortui* forms with triacetyloleandomycin and mycalamide A. The triacetyloleandomycin complex structure presented below differs significantly from that reported previously for the same compound bound to the large ribosomal subunit of Deinococcus radiodurans (4). The structure of mycalamide A complex is interesting because it binds to the E site, where normally the CCA end of an E-site tRNA would bind, and there are only two other structures available for inhibitors bound at or near the same site in the ribosome (27).

Triacetyloleandomycin is a semisynthetic derivative of the macrolide oleandomycin (Fig. 1a) (7). Both its structure and its

mode of action closely resemble those of other 14-membered macrolides like erythromycin (10), and it is used to treat diseases caused by gram-positive bacteria in some parts of the world. As expected, triacetyloleandomycin binds in the peptide exit tunnel and therefore should inhibit the growth of nascent peptide chains but not the peptidyl transferase reaction as such (17).

Mycalamide A was first isolated from a marine sponge of the genus *Mycale* that is found in the waters of New Zealand, and it is chemically related to pederin, onnamides A to F, and theopederins A to L (Fig. 1c) (29). Mycalamide A disrupts protein synthesis in cell-free protein synthesis systems from rabbit reticulocyte lysates with a 50% inhibitory concentration in the micromolar range. It possesses strong antitumor activity both in vitro and in vivo (8).

Binding experiments demonstrating that pederins compete with 13-deoxytedanolide (18), which is known to be an E-site inhibitor (27), and the chemical similarity between mycalamide A and pederins suggest that mycalamide A is likely to be an E-site inhibitor. The structure presented here confirms that inference and explains why compounds that inhibit protein synthesis by binding to the E site are likely to discriminate very strongly between eukaryotic and bacterial cells. It remains to be seen if an E-site inhibitor specific for bacteria exists in nature or can be synthesized de novo.

MATERIALS AND METHODS

Inhibitors. Triacetyloleandomycin was purchased from Sigma. Mycalamide A was provided by Sarath Gunasekera of the Harbor Branch Oceanographic Institution in Fort Pierce, FL (Florida Atlantic University).

Preparation of crystals. Crystals of the large ribosomal subunit from *H. marismortui* were prepared as previously described (1, 2, 24). Preformed crystals were soaked in solutions containing inhibitors dissolved in buffer B (12% polyethylene glycol 6000, 20% ethylene glycol, 1.7 M NaCl, 0.5 M NH₄Cl, 1 mM CdCl₂, 100 mM KCH₃COO, 6.5 mM CH₃COOH [pH 6.0], and 5 mM β-mercaptoethanol and with 30 mM MgCl₂ or 21 mM MgCl₂ and 30 mM SrCl₂) at a 1 mM concentration for several hours at 4°C, with one exchange of soaking solution.

Crystallography. Diffraction data were collected at the National Synchrotron Light Source (Brookhaven National Laboratory) on beam line X29 and at the Advanced Photon Source (Argonne National Laboratory) on beam line 24-ID-C, as previously described (1, 2, 23). After data processing with HKL2000 (19), electron density maps were calculated for each complex using ($|F_o(hkl)| + drug - |F_o(hkl)| - drug$) differences as amplitudes and phases derived from the structure

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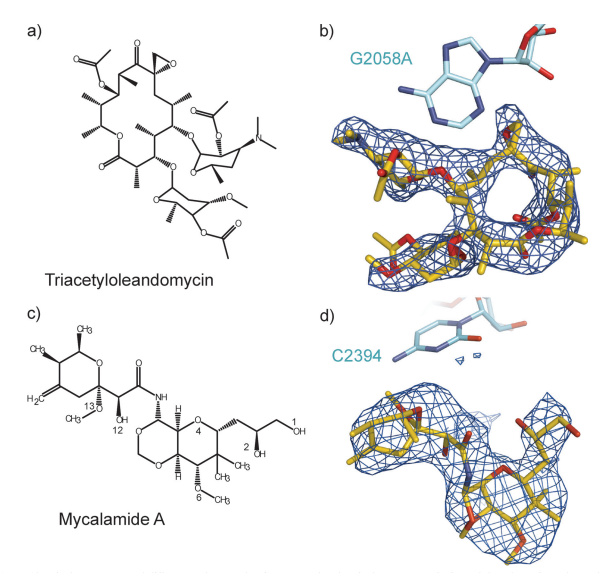


FIG. 1. Chemical structures and difference electron density maps. The chemical structures of triacetyloleandomycin and mycalamide A are shown in panels a and c, respectively. The atoms involved in hydrogen bonding interactions with the ribosome are labeled for mycalamide A. The features in the corresponding difference electron density maps that were assigned to triacetyloleandomycin and mycalamide are shown in panels b and d, respectively, with the structures of the drugs superimposed on them. Difference electron density maps are contoured at 3 σ .

of the large ribosomal subunit with no drug bound (Protein Data Bank accession no. 3CC2) (5). The starting models for both drugs were created using the Dundee PRODRG2 server (http://davapc1.bioch.dundee.ac.uk/prodrg/). The structures of the ribosome complexes with drugs were refined in CNS (6). Data at resolution shell 3.00 to 2.90 Å, at which I/ σ was < 1.2, were included in both data processing and refinement for triacetyloleandomycin. Structures were visualized using the program O (9), and figures were generated using PyMOL (15). Buried surface areas were calculated in CNS (6) using the Lee and Richards algorithm (16).

Protein structure accession numbers. Coordinates and structure factors have been deposited in the Protein Data Bank with the following accession numbers: 3I56 for triacetyloleandomycin and 3I55 for mycalamide A.

RESULTS

Crystal structures. The structures of the complexes that mycalamide A and triacetyloleandomycin form with the large

TABLE 1. Crystallographic statistics of data collection	TABLE 1.	Crystallographic	statistics of	data collection ^a
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Crystal soak ^b	Resolution (Å)	I/σ	$R_{ m merge}$ (%)	Completeness (%)	Redundancy
Triacetyloleandomycin	50.0–3.00 (3.11–3.00)	6.06 (1.2)	15.9 (73.9)	99.8 (99.7)	6.9 (3.8)
Mycalamide A	50.0–3.10 (3.21–3.10)	12.11 (1.9)	17.1 (92.7)	100.0 (100.0)	7.5 (6.7)

^a All data were collected at wavelengths ranging from 1.07 Å to 1.1 Å. Numbers in Parentheses refer to the value of the statistic in question in the highest-resolution shell of data. The redundancy is the average number of times each reflection was measured. ^b The concentration of both antibiotics was 1 mM.

TABLE 2. Statistics of model refinement									
Crystal soak ^a	Resolution (Å)	$R_{ m cryst}$ (%)	$R_{\rm free}$ (%)	Bond (Å)	Angle (°)	Protein Data Bank accession no.			
Triacetyloleandomycin Mycalamide A	50.0–2.90 50.0–3.10	20.44 19.94	24.51 24.73	0.007 0.005	1.1 1.0	3156 3155			

^a The concentration of both antibiotics was 1 mM.

ribosomal subunit from *H. marismortui* were determined by X-ray crystallography using the apo structure for that subunit (2) as the source of initial phase information. Table 1 summarizes the crystallographic statistics that characterize these structures. Their resolutions lie between 3.00 and 3.10 Å.

Structural analysis began for both drugs with the computation of difference electron density maps which were obtained using the differences in the amplitudes between data from crystals into which the drug had been soaked and data obtained from crystals lacking the drug, i.e., $[|F_o(hkl)| + drug - |F_o(hkl)| - drug]$ (see Materials and Methods). Initial phases were derived from the apo 50S structure. In the difference maps for both drugs, there was only one positive feature large enough to accommodate the relevant drug, and in both cases that feature had an appropriate shape. Figures 1b and 1d show the difference density assigned to each drug with the structure of the drug superimposed. Once the location and orientation of each inhibitor were determined, models were constructed for the corresponding inhibitor-ribosome complex and refined. Table 2 summarizes the statistics of model refinement.

(i) Triacetyloleandomycin. The interaction of triacetyloleandomycin with the ribosome from H. marismortui was investigated using *H. marismortui* DT917. This strain carries a single rRNA operon with the mutation G2058A (2099), which greatly increases its affinity for macrolide antibiotics (32). (The convention used here for numbering 23S rRNA nucleotides is Escherichia coli number followed by the corresponding H. marismortui number in parentheses.) Triacetyloleandomycin binds to the large ribosomal subunit at the entrance of the nascent peptide tunnel in exactly the same way as erythromycin does. When the two macrolide-ribosome complexes are superimposed using the phosphorus atoms of the rRNA surrounding the binding site, their 14-membered lactone rings align almost perfectly (Fig. 2a). Like erythromycin, triacetyloleandomycin is inserted into the hydrophobic pocket formed by residues A2058 (2099), A2059 (2100), and G2611 (2646), and upon binding, 71% of its surface area is buried.

There are some minor differences between the two complexes. In the erythromycin complex, the 2' OH group of the desosamine forms an H-bond with the N-1 of A2058 (2099). That oxygen is esterified with an acetate group in triacetyloleandomycin, and it is within hydrogen bonding distance from the N-6 of A2058 (2099) (distance, 3.5 Å). Finally, a major reorientation of A2062 (2103) accompanies triacetyloleandomycin binding (not shown). Since the positioning of this base is highly variable, the significance of this observation is uncertain.

(ii) Mycalamide A. Mycalamide A binds in the E site of the large ribosomal subunit, where it makes extensive interactions both with the 23S rRNA and with the ribosomal protein L44e (Fig. 3a). Of its surface, 64% becomes solvent inaccessible upon binding to the ribosome, and it forms several hydrogen

bonds with the ribosome. The O-13 of mycalamide A is 3.37 Å away from the N-4 of C2394 (2431). Its O-12 is within hydrogen bonding distance of the N-2 of G2421 (2459) and the O-2 of C2395 (2432), two bases that form a Watson-Crick base pair. In addition, the O-4 of mycalamide A is 3.21 Å away from the 2' hydroxyl of the sugar of C2395 (2432), and the O-1 of mycalamide is hydrogen bonded to the backbone carbonyl oxygens of Lys51 and Lys54 of L44e (2.88 Å and 3.44 Å, respectively). Finally, the O-6 of mycalamide A may hydrogen bond with the NH₂ of Gln44 of L44 (3.57 Å).

DISCUSSION

As typical of the macrolides, triacetyloleandomycin binds in the peptide exit tunnel (17). Since the mode of binding of triacetyloleandomycin to the *H. marismortui* ribosome is very similar to that of erythromycin and its structural relatives, oleandomycin and erythromycin, both inhibit the progression of nascent peptide chains down the exit tunnel, triacetyloleandomycin presumably inhibits translation in the same way (30–33).

The position and orientation of triacetyloleandomycin in the ribosome complex reported here are quite different from those reported for the drug in the complex that it forms with the large ribosomal subunit from *D. radiodurans* (Fig. 2b) (4). In addition, the rearrangement of L22 reported to occur in *D. radiodurans* when triacetyloleandomycin binds is not evident in the *H. marismortui* complex. Since the strain of *H. marismortui* used for the study reported here carries the mutation G2058A

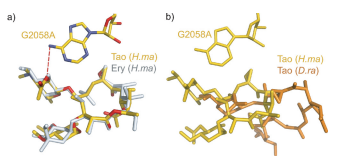


FIG. 2. Interaction of triacetyloleandomycin with the *H. marismortui* ribosome compared with other ribosome-macrolide complexes. (a) Comparison of the position and orientation of triacetyloleandomycin (gold) with those of erythromycin (gray) when both are bound to the large ribosomal subunit from *H. marismortui* (Protein Data Bank accession no. 1YI2) (31). The heteroatoms of triacetyloleandomycin are color coded, with N shown in blue and O in red. Hydrogen bonds are shown as red dashes. (b) The triacetyloleandomycin complex with the *H. marismortui* ribosome (gold) compared with the triacetyloleandomycin complex with the *D. radiodurans* ribosome (orange) (Protein Data Bank accession no. 10ND) (4).

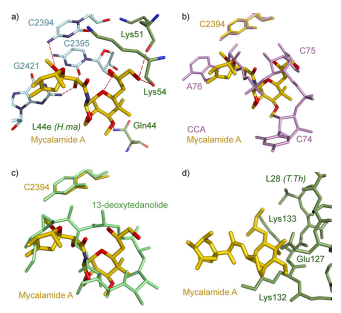


FIG. 3. Interaction of mycalamide A with the ribosome. (a) The complex that mycalamide A forms with the H. marismortui ribosome is displayed. Mycalamide A is shown with C in gold, N in blue, and O in red. 23S rRNA residues involved in hydrogen bonding are shown with C in light cyan, N in blue, and O in red. L44e residues involved in drug interactions are shown with C in green, N in blue, and O in red. Hydrogen bonds are shown as red dashes. E. coli base numbering is used for 23S rRNA residues, and H. marismortui numbering is used for L44e protein. (b) Superposition of structures of mycalamide A bound to H. marismortui and the deacylated tRNA mimic CCA oligonucleotide bound to the same ribosome (Protein Data Bank accession no. 1QVG) (25). Mycalamide A is shown with C in gold, N in blue, and O in red. CCA residues are shown in pink. (c) Superposition of the mycalamide A structure with the structure of the complex that 13deoxytedanolide forms with the H. marismortui ribosome (Protein Data Bank accession no. 20TJ) (27). Mycalamide A is shown with C in gold, N in blue, and O in red, and 13-deoxytedanolide is shown in green. (d) Superposition of the mycalamide A structure (gold) with the structure of the 70S ribosome from T. thermophilus (Protein Data Bank accession no. 2J03) (28). T. thermophilus protein L28 is shown in green, and T. thermophilus numbers are used for labeling of the protein L28.

(2099), the difference in the modes of triacetyloleandomycin binding to the ribosomes of the two species cannot be ascribed to a sequence difference at that position, which is known to be critical for macrolide binding.

Interestingly, other macrolides like erythromycin and telithromycin, which are close relatives of triacetyloleandomycin structurally, have also been reported to bind to the *D. radiodurans* ribosome in a manner different from what is seen in *H. marismortui* (4, 22, 32). However, a more recent publication, which includes a description of the redetermination of the structures of both erythromycin and telithromycin bound to the *D. radiodurans* ribosome, states that the discrepancies between the structures of drug-ribosome complexes reported for *H. marismortui* and the structures of those reported for *D. radiodurans* are in fact much smaller than they originally appeared (33). Indeed, the revised coordinates of the *D. radiodurans* erythromycin-ribosome complex are superimposed almost exactly on those of the *H. marismortui* erythromycin-ribosome complex, whereas the original erythromycin model was nearly orthogonal. Thus, it would be interesting to know if the structure first reported for triacetyloleandomycin bound to the *D. radiodurans* ribosome can be confirmed.

The structure reported here for mycalamide A bound to the ribosome offers a simple explanation for why it inhibits protein synthesis. When bound to the ribosome, it completely fills the space that is normally occupied by C75 and A76 of E-site-bound tRNAs (Fig. 3b). In this regard it is similar to 13-deoxytedanolide, another inhibitor of eukaryotic protein synthesis that binds to the E site (Fig. 3c) (27). Thus, this structure is in perfect accord with the competitive binding experiments reported by Nishimura and colleagues demonstrating that pederin, theopederin, and onnamide A, compounds that are structurally related to mycalamide A, all compete with 13-deoxytedanolide for binding in *Saccharomyces cerevisiae* ribosomes (18).

The structure reported here also explains the species specificity of mycalamide A. As shown in Fig. 3a, the drug interacts extensively with protein L44e, which is found at the E site of archaea and eukaryotes. In eubacteria, L28, a protein that is structurally unrelated to L44e, is found in about the same location. In order to investigate what might happen if mycalamide A were to bind to a eubacterial ribosome, the structure described above was superimposed onto 70S ribosome structure from *Thermus thermophilus* (Protein Data Bank accession no. 2J01) (28). This superposition reveals that mycalamide A is unlikely to be able to bind to eubacterial ribosomes. Residues Glu127, Lys132, and Lys133 of the eubacterial protein L28 extend too far into the drug's binding site (Fig. 3d).

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