

Minireview

Ribosomal Crystallography: Peptide Bond Formation, Chaperone Assistance and Antibiotics Activity

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(Received August 1, 2005; Accepted August 3, 2005)

The peptidyl transferase center (PTC) is located in a protein free environment, thus confirming that the ribosome is a ribozyme. This arched void has dimensions suitable for accommodating the 3'ends of the A-and the P-site tRNAs, and is situated within a universal sizable symmetry-related region that connects all ribosomal functional centers involved in amino-acid polymerization. The linkage between the elaborate PTC architecture and the A-site tRNA position revealed that the A-to P-site passage of the tRNA 3'end is performed by a rotatory motion, which leads to stereochemistry suitable for peptide bond formation and for substrate mediated catalysis, thus suggesting that the PTC evolved by gene-fusion. Adjacent to the PTC is the entrance of the protein exit tunnel, shown to play active roles in sequence-specific gating of nascent chains and in responding to cellular signals. This tunnel also provides a site that may be exploited for local co-translational folding and seems to assist in nascent chain trafficking into the hydrophobic space formed by the first bacterial chaperone, the trigger factor. Many antibiotics target ribosomes. Although the ribosome is highly conserved, subtle sequence and/or conformational variations enable drug selectivity, thus facilitating clinical usage. Comparisons of high-resolution structures of complexes of antibiotics bound to ribosomes from eubacteria resembling pathogens, to an archaeon that shares properties with eukaryotes and to its mutant that allows antibiotics binding, demonstrated the unambiguous difference between mere binding and therapeutical effectiveness. The observed variability in antibiotics inhibitory modes, accompanied by the elucidation of the structural basis to antibiotics mechanism justifies expectations for structural based improved properties of existing compounds as well as for the development of novel drugs.

Keywords: Antibiotics Selectivity; Elongation Arrest; Resistance; Ribosomal Antibiotics; Ribosomal Symmetrical Region; Trigger Factor.

Introduction

The ribosomes are giant ribonucleoprotein cellular assemblies that translate the genetic code into proteins. They are built of two subunits of unequal size that associate upon the initiation of protein biosynthesis to form a functional particle and dissociate once this process is terminated. The bacterial ribosomal subunits are of molecular weights of 0.85 and 1.45 Mega Dalton. The small subunit (called 30S in prokaryotes) contains an RNA chain (called 16S) of ~1500 nucleotides and 20–21 proteins, and the large one (called 50S in prokaryotes) has two RNA chains (23S and 5S RNA) of about 3000 nucleotides in total, and 31–35 proteins.

Protein biosynthesis is performed cooperatively by the two ribosomal subunits and several non-ribosomal factors, assisting the fast and smooth processivity of protein formation, required for cell vitality. While elongation proceeds, the small subunit provides the decoding-center and controls translation fidelity, and the large one contains the catalytic site, called the peptidyl-transferase-center (PTC), as well as the protein exit tunnel. mRNA carries the genetic code to the ribosome, and tRNA molecules bring the protein building block, the amino acids, to the ribosome. These L-shape molecules are built mainly of double helices, but their two functional sites, namely the anticodon loop and the CCA 3'end, are single strands. The ribosome posses three tRNA binding site, the A-(aminoacyl), the P-(peptidyl), and the E-(exit) sites. The tRNA anticodon loop interacts with the mRNA on the small subunit, whereas the tRNA acceptor stem, together with the aminoacylated or peptidylated tRNA 3'ends interacts with the large subunit. Hence, the tRNA molecules are the entities combining the two subunits, in addition to the intersubunit bridges, which are built of flexi-

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ble components of both subunits. The elongation cycle involves decoding, the creation of a peptide bond, the detachment of the P-site tRNA from the growing polypeptide chain and the release of a deacylated tRNA molecule and the advancement of the mRNA together with the tRNA molecules from the A- to the P- and then to the E-site. This motion is driven by GTPase activity.

Two decades of experimentation (reviewed in Yonath, 2002) yielded high resolution structures of the small ribosomal subunit from *Thermus thermophilus*, T30S (Schluzen et al., 2000; Wimberly et al., 2000), of the large subunit from the archaeon *Haloarcula marismortui*, H50S (Ban et al., 2000) and from the eubacterium *Deinococcus radiodurans*, D50S (Harms et al., 2001). Together with additional structures of their complexes with substrate analogs (Bashan et al., 2003a; Hansen et al., 2002a; Nissen et al., 2000; Schmeing et al., 2002; Yusupov et al., 2001) and with a medium resolution structure of the whole ribosome from *T. thermophilus*, T70S in complex with three tRNA molecules (Yusupov et al., 2001), these structures shed light on the vast amount of biochemical knowledge accumulated in over five decades of ribosomal research.

The actual reaction of peptide bond formation is performed by a nucleophilic attack of the primary amine of the A-site amino acid on the carbonyl carbon of the peptidyl tRNA at the P-site. This reaction can be performed by tRNA 3'end analogs. Puromycin is a universal inhibitor mimicking the tip of the tRNA 3'end. Its binding to the ribosome in the presence of an active donor substrate can result in peptide-bond formation uncoupled from the translocation of the A-site tRNA, namely from the polymerization of the amino acids into polypeptides. Puromycin has been commonly used as a minimal substrate for investigating peptide bond formation, in a process called the "fragment reaction", which yields a single peptide bond. The finding that naked ribosomal RNA can catalyze the "fragment reaction" (Nitta et al., 1998; Noller et al., 1992); the localization of the PTC in an environment rich in conserved nucleotides (Harms et al., 2001; Yusupov et al., 2001) the usage of puromycin derivatives bound to the partially disordered large subunits, H50S (Nissen et al., 2000) together with a compound originally presumed to resemble the reaction intermediate (Moore and Steitz, 2003) led to the suggestion that ribosome catalysis resembled the reverse reaction of serine proteases, and that specific ribosome nucleotides participate in the chemical events of peptide bond formation, as a "general base" (Nissen et al., 2000).

Biochemical, kinetic and mutational results (Barta et al., 2001; Polacek et al., 2003; Sievers et al., 2004; Thompson et al., 2001; Weinger et al., 2004; Youngman et al., 2004) and the finding that the PTC conformation in crystalline H50S hardly resembles the active one (Bayfield et al., 2001) challenged this hypothesis, and indicated that there is no ground for the expectation that a complex as-

sembly such as the ribosome catalyzes protein biosynthesis by the reverse of a common enzymatic mechanism. Indeed, the well ordered structure of the large ribosomal subunit from *D. radiodurans*, D50S (Harms et al., 2001), determined under conditions resembling its optimal growth environment, revealed that the striking ribosomal architecture provides all structural elements enabling its function as a aminoacid polymerase that ensures proper and efficient elongation of nascent protein chains in addition to the formation of the peptide bonds (Agmon et al., 2003; 2004; 2005; Baram and Yonath, 2005; Bashan and Yonath, 2005; Bashan et al., 2003a; 2003b; Yonath, 2003a; 2003b; 2005; Zarivach et al., 2004).

Being a prominent key player in a vital process, the ribosome is targeted by many antibiotics of diverse nature. Consequently, since the beginning of therapeutic administration of antibiotics, ribosomal drugs have been the subject to numerous biochemical and genetic studies (reviewed in Auerbach et al., 2002; 2004; Courvalin et al., 1985; Gale et al., 1981; Gaynor and Mankin, 2003; Katz and Ashley, 2005; Knowles et al., 2002; Poehlsgaard and Douthwaite, 2003; Sigmund et al., 1984; Spahn and Prescott, 1996; Vazquez, 1979; Weisblum, 1995; Yonath, 2005; Yonath and Bashan, 2004). These findings were enforced by the lessons learned from the high resolution structures of their complexes with ribosomal particles (Berisio et al., 2003a; 2003b; Brodersen et al., 2000; Carter et al., 2000; Hansen et al., 2002b; 2003; Harms et al., 2004; Pfister et al., 2004; 2005; Pioletti et al., 2001; Schluzen et al., 2001; 2003; 2004; Tu et al., 2005). These were found indispensable for illustrating the basic mechanisms of antibiotics activity and synergism, provided the structural basis for mechanisms of antibiotic resistance and enlightens the principles of antibiotics selectivity, namely the discrimination between pathogens and humans, the key for therapeutical usefulness (Auerbach et al., 2004; Yonath, 2005; Yonath and Bashan, 2004).

Since X-ray crystallography requires diffracting crystals, and since so far no ribosomes from pathogenic bacteria could be crystallized, the crystallographic studies are confined to the currently available crystals. The findings that *E. coli* and *T. Thermophilus* are practically interchangeable (Gregory et al., 2005; Thompson and Dahlberg, 2004) and that both crystallizable ribosomes are from eubacteria that resemble pathogens, permit considering them as suitable pathogen models for ribosomal antibiotics. Genetically engineered pathogen models, such as *Mycobacterium smegmatis*, can also serve as pathogen models. These should be advantageous, as they can provide isogenic mutations (Pfister et al., 2004). Similarly, for mutagenesis studies species with single rRNA operon chromosomal copy, such as *Halobacterium halobium* (Mankin and Garrett, 1991; Tan et al., 1996) are beneficial.

Additional concern relates to the relevance of the crystallographic results. The ability to rationalize biochemical,

functional and genetics observations by the crystallographic structures demonstrate the inherent reliability of the crystallographic results. The consistencies of drug locations with biochemical and resistance data, alongside the usage of crystalline complexes obtained at clinically relevant drug concentrations, manifest further the reliability of the crystallographic results. Last, the similarities of the structures of T30S wild type as well as of its complexes with antibiotics, elucidated by two independent laboratories (Brodersen *et al.*, 2000; Carter *et al.*, 2000; Pioletti *et al.*, 2001; Schlutzenzen *et al.*, 2000; Wimberly *et al.*, 2000), indicate that dissimilarities observed crystallographically reflect genuine variability in drug binding modes.

This article focuses on the ribosomal architectural elements that govern both the positional and the chemical contributions to the catalysis of peptide bond formation, sheds light on the essentiality of accurate substrate placement and portrays the parameters dictating it; points at evolution aspects implicated by the ribosomal symmetry; describes how the first chaperon to be encounter by the nascent chain contributes to the mature protein correct folding; and points at a possible correlation between peptide bond formation, nascent protein progression, cotranslational folding and cellular regulation. It also relates the structural findings associated with ribosomal antibiotics action and highlights the unique achievements of these studies as well as their shortcoming. Full coverage of the vast amount of biochemical, structural and medical knowledge is beyond the scope of this article. Instead, it emphasizes the structural finding associated with antibiotics selectivity and synergism, and describes current issues concerning to the acute problem of resistance to antibiotics.

Symmetry within the asymmetric ribosome

The recently determined three-dimensional structures of ribosomal particles from eubacteria and archaea revealed that the interface surfaces of both subunits are rich in RNA (Fig. 1), and localized the PTC in a protein-free environment the middle of the large subunit, thus confirming that the ribosome is a ribozyme. Further analysis, based on the structure of a functional complex of D50S, showed that the peptide bond is being formed within a universal sizable symmetrical region (Fig. 2), containing about 180 nucleotides (Agmon *et al.*, 2003; 2004; 2005; Baram and Yonath, 2005; Bashan and Yonath, 2005; Bashan *et al.*, 2003a; 2003b; Yonath, 2003a; 2003b; 2005; Zarivach *et al.*, 2004). The symmetrical region is located in and around the PTC, and its symmetry axis, which is directed into the protein exit tunnel, passes through the peptidyl transferase center, midway between the RNA features shown to host the 3'ends of the A- and the P- sites tRNA. Although first identified in D50S, this symmetrical region seems to be a universal ribosomal feature, as it is present in

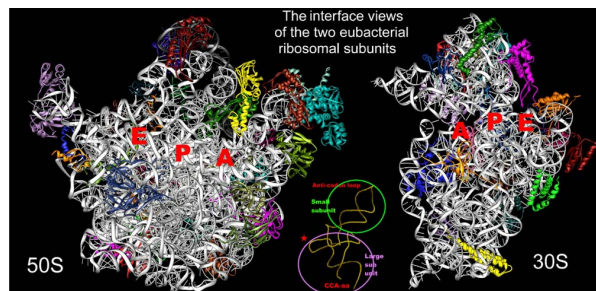


Fig. 1. The two ribosomal subunits. The small (30S) and the large (50S) ribosomal subunit, from *T. thermophilus* (Schlutzenzen *et al.*, 2000) and *D. radiodurans* (Harms *et al.*, 2001), respectively, showing their intersubunit interfaces. In both, the ribosomal RNA is shown as silver ribbons, and the ribosomal proteins main chains in different colors. A,P,E designate the approximate locations of the A-,P-, and E- tRNA anticodons on the small subunit, and the tRNA beginning of the tRNA acceptor stems (the red star on the inserted figure) on the large one. The regions of tRNA interactions with each subunit are shown on the tRNA molecule, inserted in the middle. The red star indicates the position at which the tRNA acceptor stem meets the large subunit.

all known structures of the large ribosomal subunit (Fig. 2A).

The symmetrical region extends far beyond the vicinity of the peptide synthesis location and interacts, directly or through its extensions, with all ribosomal functional features that are relevant to the elongation process: the tRNA entrance and exit regions, namely the L7/L12 stalk and the L1 arm, respectively, the peptidyl transferase center, and the bridges connecting the two subunits (Fig. 2B), among which bridge B2a resides on the PTC cavity and reaches the vicinity of the decoding center in the small subunit (Yusupov *et al.*, 2001). The 3'ends of the A- and the P-tRNAs bind to the PTC, and even the 3' end of the E-site tRNA contacts the neighborhood of the symmetry region edge in the T70S complex (Yusupov *et al.*, 2001) but not in H50S complexed with a fragment of the E-site tRNA (Schmeing *et al.*, 2003). Hence, the spatial organization of this region and its central location may enable signal transmission between the remote locations on the ribosome (Agmon *et al.*, 2003).

Peptide bond formation and aminoacid polymerization

Located at the bottom of a V-shaped cavity (Fig. 3A), the PTC is an arched void with dimensions suitable for accommodating the 3'ends of the A- and the P-site tRNAs. Each of the symmetry related sub-regions contains half of the PTC, namely either the A- or the P-site, and the axis relating them by $\sim 180^\circ$ rotation, is located in the middle of the PTC, midway between the two tRNA binding sites. In a complex of D50S with a 35-nucleotides oligomers

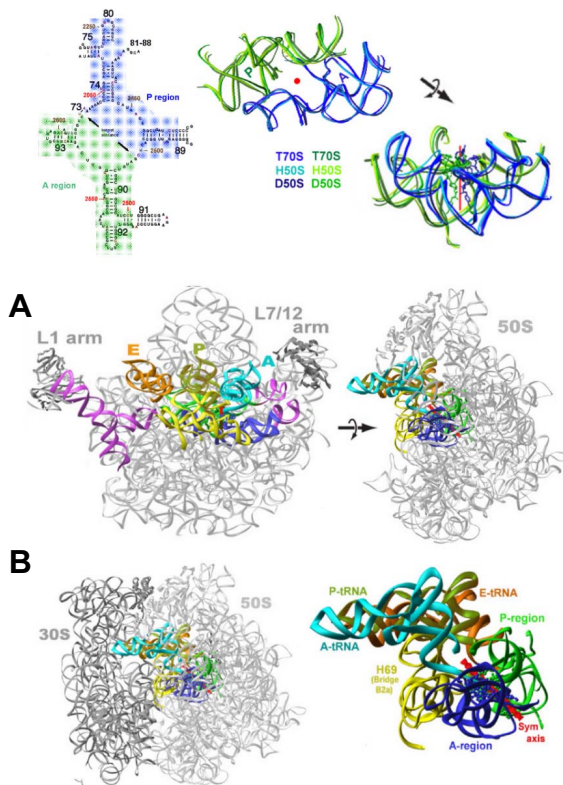


Fig. 2. The symmetrical region within the large ribosomal subunit. Throughout, the part containing the A-loop (namely the site of A-site tRNA 3'end) is blue (called: A-region), and the corresponding one, containing the P-site tRNA 3'end, is green. Similarly, the A-site tRNA mimic is shown in blue and the derived P-site tRNA is green. The symmetry axis is shown in red. **A.** Left: The secondary structure scheme of the symmetrical region in D50S, shown in a manner that exhibits the 2-fold symmetry of the region (*E. coli* and D50S numbering in gray and red, respectively). Middle and right: Two orthogonal views (top and side, respectively) of the superposition of the backbone of the symmetrical regions in all known structures: the entire ribosome from T70S (PDB 1GIY), D50S (PDB 1NKW) and H50S (PDB 1JJ2). Note that the A-site mimic and the derived P-site are incorporated into the side view. **B.** The symmetry related region within the large subunit (upper panel) and the entire ribosome (bottom left). The direct extensions of the symmetrical region are shown in purple. Ribosomal RNA is shown in gray ribbons. The positions of the docked three tRNA molecules, as seen in the complex of T70S (PDB 1GIY) are also shown, to indicate their relationship to the symmetry related area. The gold feature is the intersubunit bridge (B2a) that combines the two ribosomal active sites. An enlarged view of the symmetry related region is shown in right bottom corner. Note the strategic location of H69, which bridges the two subunits, and plays a major role in A-site tRNA accurate placement.

mimicking the aminoacylated-tRNA acceptor stem, called ASM (Fig. 3A), the bond connecting the 3'end with the

acceptor stem was found to roughly coincide with the symmetry axis (Bashan *et al.*, 2003a), suggesting that tRNA A → P-site passage is a combination of two independent, albeit synchronized motions: a sideways shift of most of the tRNA molecules, performed as a part of the overall mRNA/tRNA translocation, and a rotatory motion of the tRNA 3'end within the PTC (Fig. 3B). The path provided by the rotatory motion is confined by the PTC rear wall and by two nucleotides that bulge from the front wall into the PTC center.

Simulation of the rotatory motion (Fig. 3C) revealed that it is navigated and guided by striking architectural design of the PTC, and that it terminates in a stereochemistry appropriate for a nucleophilic attack of the A-site amino acid on the carbonyl carbon of the peptidyl tRNA at the P-site (Agmon *et al.*, 2003; 2005; Bashan *et al.*, 2003a; 2003b). The spatial match between the PTC rear wall and the contour of the tRNA aa-3'end, formed by the rotatory motion, indicates that it provides the template for the translocation path. From the other side of the PTC, two universally conserved nucleotides A2602 and U2585 (*Escherichia coli* nomenclature, throughout), bulge towards the PTC center (Figs. 3B and 3C) and seem to anchor and/or propel the rotatory motion (Agmon *et al.*, 2003; 2004; 2005; Baram and Yonath, 2005; Bashan *et al.*, 2003a; 2003b; Polacek *et al.*, 2003; Zarivach *et al.*, 2004).

Importantly, the derived P-site tRNA 3'end forms all interactions found biochemically (e.g. Bocchetta *et al.*, 1998; Green *et al.*, 1997) and the orientation of the so-created peptide bond is adequate for the ribosomal subsequent tasks, including the release of the peptidyl-tRNA and the entrance of the nascent protein into the exit tunnel. Hence, it appears that the ribosome provides a striking architectural frame, ideal for amino acid polymerization. Thus, the ribosome functions as an enzyme, a ribozyme, responsible not only to peptide bond formation, but also for the successive reactions, namely the creation of polypeptides that can eventually acquire their functional fold (Agmon *et al.*, 2003; 2004; 2005; Bashan *et al.*, 2003a; Zarivach *et al.*, 2004).

Accurate substrate positioning

Positioning reactants in orientation suitable for chemical reactions is performed by almost all bio-catalysts (Jencks, 1969, reissued 1987). Different from enzymes catalyzing a single chemical reactions, such as proteases, and similar to other polymerases, the ribosome provides the means not only for the chemical reaction (peptide bond formation), but also for substrates motions required for the processivity of peptide bond formation, namely for amino acid polymerization. However, a prerequisite for achieving the ribosome contribution is accurate substrate placement (Yonath, 2003a; 2003b).

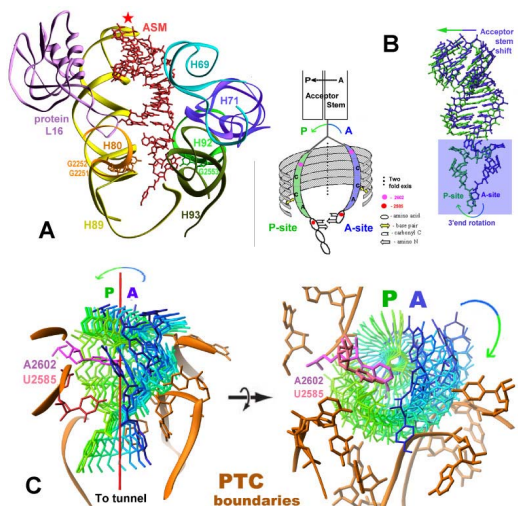


Fig. 3. The rotatory motion. Throughout, the part containing the A-loop (namely the site of A-site tRNA 3' end) is blue (called: A-region), and the corresponding one, containing the P-site tRNA 3' end, is green. Similarly, the A-site tRNA mimic is shown in blue and the derived P-site tRNA is green. The symmetry axis is shown in red. **A.** The PTC pocket, including ASM, an A-site substrate analog, which is represented by atoms in red. The red star indicates the position at which the tRNA acceptor stem meets the large subunit, as in Fig. 1. The RNA components of the PTC pocket are numbered according to *E. coli* nomenclature (also shown in Fig. 5D) and colored differently. Note the remote interactions positioning the substrate, as well as the universal contributors to the 3' end base pairs (a single basepair at the A-site, and two in the P-site). **B.** Left: A schematic cartoon of the rotatory mechanism. The tRNA 3' ends are represented by banana shaped objects, divided by dotted lines into the four nucleotides composing them. The PTC rear wall is drawn as ribs. The locations of the interactions of the anchoring nucleotides, A2602 and U2585 with the tRNA 3' ends are marked by colored circles. Right: Translocating tRNA from A- (blue) to P- (green) site. The dark green line is the division between the shifted double helical region (above) and the rotating 3' end (below). The tRNA regions interacting with the small subunit are represented by blue-green boxes. Straight arrows show the shifting direction. The round one represents the rotatory motion. **C.** Two orthogonal snapshots (sideways and from the tunnel into the PTC) of intermediate stages (represented by gradual transformation from blue to green) in the motion of the A-site tRNA CCA from the A- to the P-site. The two front-wall bulged nucleotides are shown in pink and magenta. The simulation was performed by rotating the ASM aminoacylated 3' end by (10 times 18° each) within D50S PTC, around the bond connecting the ASM 3' end with its acceptor stem, accompanied by a 2 \AA shift in the direction of the tunnel, as implied by the overall spiral nature of the PTC template. The blue-green round arrows show the rotation direction. The ribosomal components belonging to the PTC rear wall, that confine the exact path of the rotatory motion, are shown in gold. The two front wall flexible nucleotides, A2602 and U2585, are colored in magenta and pink, respectively.

The universal Watson-Crick basepair between C75 of A-site tRNA terminus and G2553 (Fig. 3A) (Kim and Green, 1999), and its symmetrical mate at the P-site, namely the basepair between C75 and G2251, exist in all known structures (Bashan *et al.*, 2003a; Hansen *et al.*, 2002a; Nissen *et al.*, 2000; Schmeing *et al.*, 2002; Yusupov *et al.*, 2001). Positioning governed solely by this basepair is sufficient for entropy driven peptide bond formation (Gregory and Dahlberg, 2004; Sievers *et al.*, 2004). However, it may not suffice for allowing smooth amino acid polymerization, as shown by the correlation found between the rates of peptide bond formation and the substrate type. Thus, compared to the reaction rate with full size tRNAs, when using the minimal substrate puromycin, such as the “fragment reaction” reactants, the peptide bond is being formed at significantly reduced rates (Moore and Steitz, 2003). Consistently, the locations and orientations of all “fragment reaction” reactants in ribosomal crystals indicate a need to undergo repositioning and/or rearrangements in order to participate in peptide bond formation (Moore and Steitz, 2003; Yonath, 2003b). This time-consuming process can be responsible to the slowness of the “fragment reaction”.

These observations indicate that the A-site basepairing is not sufficient for accurate tRNA placement, essential for performing the rotatory motion. As the main structural difference between fragment reaction reactants and full size tRNA is the substrates relative sizes, it appears that accurate positioning is achieved by remote interactions of the A-site tRNA acceptor stem with the upper part of the PTC cavity (Agmon *et al.*, 2003; Yonath, 2003a; 2003b). Remote interactions cannot be formed by substrate analogs that are too short to reach the PTC cavity upper part, as “fragment reaction” participants, or when Helix H69, the remote interactions mate at the PTC upper end (Figs. 2B and 3A) is disordered, as in H50S structure (Ban *et al.*, 2000; Nissen *et al.*, 2000). It appears, therefore, that the CCA basepairing contributes to the overall positioning of the 3' end of the aminoacylated tRNA, whereas the efficiency of peptide bond formation depends on the tRNA remote interactions. The rotatory motion guides the A-site tRNA to land at the P-site in an orientation appropriate for the creation of the two basepairs. This double basepair seem to stabilize the orientation of P-site tRNA at the conformation essential for the P-site tRNA catalytic role in peptide bond formation (Dorner *et al.*, 2002; Weinger *et al.*, 2004). Hence, the rotatory motion not only leads to a configuration suitable for peptide bond formation (Agmon *et al.*, 2003; Bashan *et al.*, 2003a), it also places the reactants at a distance reachable by the O2' of the P-site tRNA A76.

Remote placement of the A-site 3' end of the tRNA seems to be designed to tolerate variability in PTC binding, as it is required to comply with the ability of the ribosome to accommodate all of the amino acids, and to allow for the rotatory motion. It appears therefore that the tRNA

size and shape and the overall ribosome architecture determines the position of the tRNA molecules and the universal basepairs, described above, establish the approximate inclination of the A-site tRNA 3' end, and facilitates P-site mediated catalysis. Accurate A-site tRNA alignment, however, is governed by its remote interactions, and since such placement is the prerequisite for the processivity of protein biosynthesis, it appears that the role played by the remote interactions supersedes all others. This conclusion is supported by the finding that in the absence of these interactions, similar, albeit distinctly different, binding modes are formed (Moore and Steitz, 2003; Yonath, 2003a; 2003b) which contrary to substrate orientation dictated by remote interactions, leads to optimal stereochemistry for the formation of a peptide bond. Hence, binding independent of remote directionality leads to various orientations, each requiring conformational rearrangements to participate in formation of a peptide bond (Moore and Steitz, 2003).

In short, by identifying the linkage between the universal ribosomal symmetry and the substrate binding mode, the integrated ribosomal machinery for peptide-bond formation, amino acid polymerization and translocation within the PTC, was revealed (Agmon *et al.*, 2003; Bashan *et al.*, 2003a). This machinery is consistent with results of biochemical and kinetic studies (Gregory and Dahlberg, 2004; Nierhaus *et al.*, 1980; Sievers *et al.*, 2004; Youngman *et al.*, 2004), proposing that positioning of the reactive groups is the critical factor for catalysis of intact tRNA substrates, and does not exclude assistance from ribosomal or substrate moieties. Hence, by offering the frame for correct substrate positioning, as well as for catalytic contribution of the P-site tRNA 2'-hydroxyl group, as suggested previously (Dorner *et al.*, 2002; Weinger *et al.*, 2004), the ribosomal architectural-frame governs the positional requirements, and provides the means for substrate mediated chemical catalysis.

PTC mobility and antibiotics synergism

The two universally conserved nucleotides A2602 and U2585 that bulge towards the PTC center (Figs. 3B and 3C) and do not obey the symmetry, are extremely flexible. In D50S A2602 is placed beneath A73 of A-site tRNA, within contact distance throughout the course of the rotation. Similarly, U2585, situated under A2602 and closer to the tunnel entrance, is located within a contact distance to bound amino acid throughout the A- to P-site motion.

Nucleotide A2602 exhibits a large variety of conformations in different complexes of the large subunit (Agmon *et al.*, 2003; Bashan *et al.*, 2003a). A2602 is involved in several tasks other than peptide bond formation, such as nascent peptide release (Polacek *et al.*, 2003) and anchoring tRNA A- to P-site passage (Agmon *et al.*, 2003; 2005;

Bashan *et al.*, 2003a; 2003b; Zarivach *et al.*, 2004). Sparsomycin, which targets A2602 (Bashan *et al.*, 2003a; Hansen *et al.*, 2003; Porse *et al.*, 1999), is a potent universal antibiotics agent, hence less useful as anti-infective drug. Comparisons between sparsomycin binding sites in D50S (Bashan *et al.*, 2003a) and H50S (Hansen *et al.*, 2003) indicated the correlation between antibiotics binding mode and the ribosomal functional-state. By binding to non-occupied large ribosomal subunits, sparsomycin stacks to A2602 and causes striking conformational alterations in the entire PTC, which should influence the positioning of the tRNA in the A-site, thus explaining why sparsomycin was considered to be an A-site inhibitor, although it does not interfere with A-site substrates (Goldberg and Mitsugi, 1966; Monro *et al.*, 1969; Porse *et al.*, 1999). In its position in D50S, sparsomycin faces the P-site. Hence, it can also enhance non-productive tRNA-binding (Monro *et al.*, 1969). Conversely, when sparsomycin enters the large subunit simultaneously with a P-site substrate or substrate-analog, it can cause only a modest conformational alteration of A2602, and because the P-site is occupied by the P-site substrate, sparsomycin stacking to A2602 appears to face the A-site (Hansen *et al.*, 2003).

The base of U2585 undergoes a substantial conformational alteration in a complex of D50S with Synercid[®] - a synergistic antibiotic agent, of which one part binds to the PTC and the other blocks the protein exit tunnel (Agmon *et al.*, 2004; Harms *et al.*, 2004). This recently approved injectable drug with excellent synergistic activity, is a member of the streptogramins antimicrobial drug family in which each drug consists of two synergistic components (S_A and S_B), capable of cooperative converting weak bacteriostatic effects into lethal bactericidal activity. In crystals of D50S-Synercid[®] complex, obtained at clinically relevant concentrations, the S_A component, dalfopristin, binds to the PTC and induces remarkable conformational alterations, including a flip of 180° of U2585 base hence paralyze its ability to anchor the rotatory motion and to direct the nascent protein into the exit tunnel (Agmon *et al.*, 2004).

As the motions of U2585 are of utmost importance to cell vitality, it is likely that the pressure for maintaining the processivity of protein biosynthesis will attempt recovering the correct positioning of U2585, by expelling or relocating dalfopristin, consistent with dalfopristin low antibacterial effect. The S_B component of Synercid[®], quinupristin, is a macrolide that binds to the common macrolide binding pocket (Auerbach *et al.*, 2004; Schluenzen *et al.*, 2001). Due to its bulkiness, quinupristin is slightly inclined within the tunnel, and consequently does not block it efficiently (Agmon *et al.*, 2004; Harms *et al.*, 2004), thus rationalizing its reduced antibacterial effects compared to erythromycin. Since within the large ribosomal subunit both Synercid[®] components interact with each other, the non-productive flipped positioning of U2585 is stabilized, and the way out

of dalbopristin is blocked. Hence, the antimicrobial activity of Synercid[®] is greatly enhanced.

Thus, the two components of this synergetic drug act in two radically different fashions. Quinupristin, the S_B component, takes a passive role in blocking the tunnel, whereas dalbopristin, the S_A component, plays a more dynamic role by hindering the motion of a vital nucleotide at the active site, U2585. It is conceivable that such mode of action consumes higher amounts of material, compared to the static tunnel blockage, explaining the peculiar composition of 7:3 dalbopristin:quinupristin in the optimized commercial Synercid[®], although the crystal structure of the complex D50S-Synercid[®] indicates binding of stoichiometric amounts of both components.

The mild streptogramins reaction on eukaryotes may be linked to the disparity between the 180° flip of U2585 in D50S (Harms *et al.*, 2004) and the mild conformational alterations of U2585 imposed by the S_A compounds on eukaryotic or archaeal ribosome, as seen in the complex of H50S with Virginiamycin-M, a streptogramin_A component (Hansen *et al.*, 2002b). This significant difference in binding modes to eubacterial vs. archaeal ribosomes appears to reflect the structural diversity of PTC conformations (Harms *et al.*, 2001; Yonath, 2002; Yusupov *et al.*, 2001), consistent with the inability of H50S PTC to bind the peptide bond formation blocker clindamycin, as well as the A-site tRNA competitor chloramphenicol (Mankin and Garrett, 1991).

On ribosome evolution and conservation

The entire symmetrical region is highly conserved, consistent with its vital function. Sampling 930 different species from three phylogenetic domains (Cannone *et al.*, 2002) shows that 36% of all of *E. coli* 23S RNA nucleotides, excluding the symmetrical region, are “frequent” (namely, found in > 95% of the sequences), whereas 98% of the symmetrical region nucleotides are categorized as such. The level of conservation increases in the inner most shell of the symmetry related region. Thus, among the 27 nucleotides lying within 10 Å distance from the symmetry axis, 75% are highly conserved, among these seven are absolutely conserved.

The universality of the symmetrical region hints that the ribosomal active site evolved by gene fusion of two separate domains of similar structures, each hosting half of the catalytic activity. Importantly, whereas the ribosomal internal symmetry relates nucleotide orientations and RNA backbone fold (Fig. 2A), there is no sequence identity between related nucleotides in the A- and the P-regions. The preservation of the three-dimensional structure of the two halves of the ribosomal frame regardless of the sequence demonstrates the rigorous requirements of accurate substrate positioning in stereochemistry supporting

peptide bond formation. Similarly, protein L16, the only ribosomal protein contributing to tRNA positioning (Agmon *et al.*, 2003; Bashan *et al.*, 2003a), displays conserved tertiary structure alongside diverged primary sequence.

Consistently, results of recent experiments addressing the functional conservations of the ribosome, show that the translational factor function and subunit-subunit interactions are conserved in two phylogenetically distant species, *E. coli* and *T. thermophilus*, despite the extreme and highly divergent environments to which these species have adapted (Thompson and Dahlberg, 2004). Similarly, mutations in *T. thermophilus* 16S and 23S rRNAs, within the decoding site and the PTC, produced phenotypes that are largely identical to their mates in mesophilic organisms (Gregory *et al.*, 2005). The contribution of protein L2 to the ribosomal polymerase activity may also shed some light on ribosome evolution. Protein L2 is the only protein interacting with both the A- and the P-regions (Agmon *et al.*, 2005). Although the entire protein is not required for the formation of a single peptide bond (Nitta *et al.*, 1998), among its two residues involved in these interactions, one (229) was shown to be essential for the elongation of the nascent chain (Cooperman *et al.*, 1995). It appears, therefore, that the main function of L2 is to provide stabilization to the PTC while elongation takes place. Stabilization of the ribosomal frame is mandatory for maintaining accurate substrate positioning, which, in turn, is required for enabling the rotatory motion, but is irrelevant to single peptide bond formation (Yonath, 2003b). This finding is consistent with the assumption that the ancient ribosome was made only from RNA and that the proteins were added later, in order to increase its fidelity and efficiency.

Involvement in maintaining the symmetry region architecture, and consequently in peptidyl transferase activity can also be attributed to protein L36. This small Zn containing protein is situated in the middle of four parallel helices and seems to stabilize their overall conformation. Two of these helices are part of the symmetry related region and two are the non-symmetrical extensions of the PTC main components. Furthermore, at its location, L36 interactions can also connect these helices with the elongation factors binding sites. Hence, in addition to stabilizing the conformation of the symmetry related region, it may also be involved in transmitting information about factor binding. The possible availability of alternative route for signaling and/or alternative means for conformation preservation, may account for the absence of L36 in some species, such as *H. marismortui*.

The ribosomal tunnel

Elongation arrest and tunnel mobility Nascent proteins emerge out of the ribosome through an exit tunnel, a uni-

versal feature of the large ribosomal subunit first seen in the mid eighties (Milligan and Unwin, 1986; Yonath *et al.*, 1987). This tunnel is adjacent to the PTC and its opening is located at the other end of the subunit (Fig. 4). Lined primarily by ribosomal RNA, this tunnel is rather kinked, has a non-uniform diameter, and contains grooves and cavities (Ban *et al.*, 2000; Harms *et al.*, 2001). Among the few r-proteins reaching its wall, the tips of extended loops of proteins L4 and L22 create an internal constriction. Five years ago, when first observed at high resolution in H50S crystal structure, this tunnel was assumed to be a firmly built passive and inert conduit for nascent chains (Ban *et al.*, 2000; Nissen *et al.*, 2000). However, biochemical results, accumulated during last decade, indicate that the tunnel plays an active role in sequence-specific gating of nascent chains and in responding to cellular signals (Etchells and Hartl, 2004; Gong and Yanofsky, 2002; Johnson, 2005; Nakatogawa and Ito, 2002; Stroud and Walter, 1999; Tenson and Ehrenberg, 2002; Walter and Johnson, 1994; White and von Heijne, 2004; Woolhead *et al.*, 2004). Furthermore, co-translational folding of nascent polypeptides into secondary structures while still within the ribosomal tunnel has been detected in several cases (e.g. Eisenstein *et al.*, 1994; Hardesty *et al.*, 1995; Woolhead *et al.*, 2004). Such initial folding events within the ribosomal tunnel seem to serve signaling between the cell and the protein-biosynthetic machinery (Johnson, 2005) rather than as segments of the correct fold of the mature protein.

Consistently, the crystal structures of complexes of the large ribosomal subunit from the eubacterium *D. radiodurans*, D50S, revealed a crevice adjacent to the tunnel that can be exploited for initial folding (Fig. 4B and Amit *et al.*, 2005) and indicated that the tunnel has the capability to oscillate between conformations (Fig. 4C), and that these alterations could be correlated with nascent protein sequence discrimination and gating (Bashan *et al.*, 2003b; Berisio *et al.*, 2003a), as well as with its trafficking into its chaperone-folding cradle (Baram and Yonath, 2005; Baram *et al.*, 2005). Analysis of these structures also shows that at its entrance, the tunnel diameter may limit the passage of highly folded polypeptides. Furthermore, in specific cases, likely to be connected with nascent-chain-tunnel interactions, the tunnel entrance properties accompanied by the incorporation of rigid residues, such as proline, may hamper the progression of protein sequences known to arrest elongation (Gong and Yanofsky, 2002; Nakatogawa and Ito, 2002).

So far most of the tunnel functional roles have been attributed to mobile extended loops of ribosomal proteins that penetrate its walls, which are primarily made of ribosomal RNA. Examples are the tips of extended loops of proteins L22 and L23 that seem to provide communication routes for signaling between the ribosome and the cell, as their other ends are located on the solvent side of

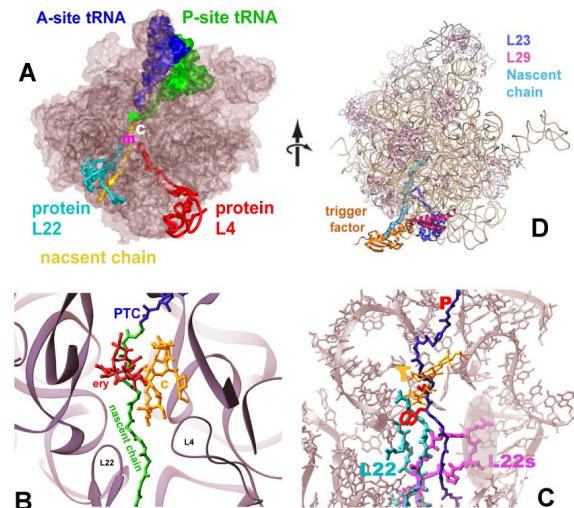


Fig. 4. The ribosome tunnel. **A.** A section through D50S (in light purple) with docked A- and P-sites tRNA and modeled polyaniline nascent chain (yellow). The approximate positions of the PTC (P), the hydrophobic crevice (C), and the macrolide pocket (m) are marked. The main chains of proteins L4 and L22 are shown in red and cyan, respectively. **B.** The hydrophobic crevice (C), in relation to the PTC, the macrolide-binding site, represented by erythromycin (ERY), to the tunnel constriction composed of the tips of the elongated loops of proteins L4 and L22, and to the possible path of the nascent chain (modeled as polyaniline and shown in green), Rapamycin binding mode is shown in gold. **C.** A view parallel to the tunnel long axis (rRNA in olive green) with a modeled nascent chain (blue). The tip of the ribosomal protein L22 beta-hairpin at its native and swung (L22_s) conformations, the latter induced by troleandomycin (T, in gold) binding, are shown in cyan and magenta, respectively. The modeled polypeptide chain (blue) represents a nascent protein with the sequence motif known to cause SecM (secretion monitor) elongation arrest. This motif is located about 150 residues from the N-terminus and has the sequence XXXXXW-XXXXXXXXXXP, where X is any amino acid and P (proline) is the last amino acid to be incorporated into the nascent chain (based on Nakatogawa and Ito, 2002). The positions of two key residues for nascent protein arrest, proline and tryptophane, are highlighted in red, to indicate the stunning correlation between its position and that of troleandomycin (T, in gold). The specific proline of SecM that is required for the arrest when incorporated into the protein at the PTC is the top amino acid of the modeled nascent chain is designated by P. The shaded area designates the region where mutants bypassing the arrest were depicted (Nakatogawa and Ito, 2002). **D.** A side view of the structure of trigger factor in complex with D50S (represented by purple-brown RNA backbone and purple-pink ribosomal proteins). The bound trigger factor binding domain is shown in orange, and a modeled polypeptide chain in cyan. Ribosomal proteins L29 and L23 are highlighted in magenta and blue, respectively. Note the elongated loop of L23, a unique eubacterial feature, which reaches the interior of the tunnel, to a location allowing its interaction with the emerging nascent chain.

the ribosome, in the proximity of the tunnel opening (Baram and Yonath, 2005; Berisio *et al.*, 2003a; Harms *et al.*, 2001). Furthermore, the beta-hairpin tip of L22 can swing across the tunnel around its accurately placed hinge (Fig. 4C), and gate the tunnel. This motion appears to provide a general mechanism for elongation arrests triggered by specific cellular conditions, since the interacting nucleotides with the swung L22 hairpin tip are identical to those identified in mutations bypassing tunnel arrest (Agmon *et al.*, 2003; Bashan *et al.*, 2003b; Berisio *et al.*, 2003a). Thus, this elongated ribosomal protein (Harms *et al.*, 2001; Unge *et al.*, 1998) that stretches along the large subunit may not only contribute to the dynamics associated with tunnel arrest, but also participate in signal transmission between the cell and the ribosomal interior.

Intra-ribosome chaperon activity? The crystal structure of a complex of the large ribosomal subunit from *D. radiodurans* co-crystallized with rapamycin, a polyketide with no inhibitory activity, revealed that rapamycin binds to a crevice located at the boundaries of the nascent protein exit tunnel, opposite to the macrolide pocket (see below and Figs. 4A and 4B). Being adjacent the ribosome tunnel, but not obstructing the path of nascent chains at extended conformation, this crevice may provide the site for local cotranslational folding of nascent chains (Amit *et al.*, 2005). The size of this crevice is suitable for accommodating small secondary structural elements, and therefore may provide nascent chains a site for adopting a particular fold at the early stage of tunnel passage, consistent with a large range of biochemical evidence, obtained mainly for transmembrane proteins, implicating cotranslational folding (Etchells and Hartl, 2004; Johnson, 2005; Stroud and Walter, 1999; Tenson and Ehrenberg, 2002; Walter and Johnson, 1994; White and von Heijne, 2004; Woolhead *et al.*, 2004).

Similar to rapamycin, transmembrane protein segments are highly hydrophobic, and therefore may be accommodated within the crevice. Hence, this crevice may provide the space as well as the hydrophobic patch that might act as an inner-tunnel chaperone, consistent with findings interpreted as nascent chain folding near the PTC, which was proposed to correlate with sequential closing and opening of the translocon at the ER membrane (Woolhead *et al.*, 2004). Hence the detection of the crevice confirms that the tunnel possesses specific binding properties, and suggests that this crevice plays a role in regulating nascent protein progression, thus acting as an intra-ribosome chaperon.

The cotranslational folding may be only transient, until messages are transmitted to other cell components [e.g. the translocon pore (Etchells and Hartl, 2004; Woolhead *et al.*, 2004). Alternatively, it is conceivable that once small nucleation centers are formed, they may progress through the tunnel by temporary expansions of the tunnel-

diameter, as observed recently for translation-arrested ribosomes (Gilbert *et al.*, 2004). Cotranslational folding is frequently observed for eukaryotic membrane proteins. These may possess a comparable crevice, as a similar feature could be identified also in the archaeal H50S. However, although existence of a crevice is postulated in ribosomes from all kingdoms of life, this does not imply structural identity, since phylogenetic diversity should play a considerable role in its detailed structure, as found at the macrolide-binding pocket (Auerbach *et al.*, 2004; Baram and Yonath, 2005; Pfister *et al.*, 2004; Yonath, 2005; Yonath and Bashan, 2004). Hence, the binding affinities of this crevice should vary, explaining why rapamycin is not known to strongly inhibit membrane proteins translation.

The first encounter with ribosome associated chaperone The complex process of folding newly synthesized proteins into their native three-dimensional structure is vital in all kingdoms of life. Although, in principle, protein can fold with no assistance of additional factors, since their sequences entail their unique folds, under cellular conditions nascent polypeptides emerging out of the ribosomal tunnel are prone to aggregation and degradation, and thus require assistance. The cellular strategy to promote correct folding and prevent misfolding involves a large arsenal of molecular chaperones (Bukau *et al.*, 2000; Frydman, 2001; Gottesman and Hendrickson, 2000; Hartl and Hayer-Hartl, 2002; Rospert, 2004; Thirumalai and Lorimer, 2001). These proteins are found in all kingdoms and the existence of ribosome-associated chaperones is a highly conserved principle in eukaryotes and prokaryotes, although the involved components differ between species.

In eubacteria, the folding of cytosol proteins is coordinated by three chaperone systems: the ribosome-associated Trigger Factor, DnaK, and GroEL. Trigger factor (TF), a unique feature of eubacteria, is the first chaperone encountering the emerging nascent chain. This 48 kDa modular protein is composed of three domains, among which the TF N-terminal domain (TFa) that contains a conserved "signature motif", mediates the association with the ribosome (Maier *et al.*, 2005). It cooperates with the DnaK system, and their combined depletion causes a massive aggregation of newly synthesized polypeptides as well as cell death above 30°C (Deuerling *et al.*, 1999). Biochemical studies showed that TF binds to the large ribosomal subunit at 1:1 stoichiometry by interacting with ribosomal proteins L23 and L29 (Blaha *et al.*, 2003; Kramer *et al.*, 2002).

Protein L23 exists in ribosomes from all kingdoms of life, but belongs to the small group of ribosomal proteins that display a significant divergence from conservation. Thus, in all species, it is built of an almost identical globular domain. In eubacteria, however, it possesses a unique feature, a sizeable elongated loop, which in of *D. radiodurans* extends from the vicinity of the tunnel opening all the way into the

tunnel interior (Fig. 4D) and in (Harms *et al.*, 2001), and can actively interact with the nascent protein passing through it (Baram and Yonath, 2005; Baram *et al.*, 2005), implying a possible dynamic control.

The high resolution crystals structure of D50S in complex with the TFA domain from the same source showed that the “signature motif” and its few aminoacids extension (called the “extended signature motif”), interact with the large ribosomal subunit near the tunnel opening (Fig. 4D) at a triple junction between ribosomal proteins L23 and L29 and the 23S rRNA (Baram *et al.*, 2005), consistent with a previous suggestion (Kristensen and Gajhede, 2003). Despite the similarity between the overall structures of the ribosome-bound and the unbound TFA (Ferbitz *et al.*, 2004; Ludlam *et al.*, 2004), significant difference were detected between their conformations and that of the bound TFA, indicating a substantial conformational rearrangement of TFA upon binding to the ribosome. These alterations result in the exposure of a sizable hydrophobic patch facing the interior of the ribosomal exit tunnel, which should increase the tunnel’s affinity for hydrophobic segments of the emerging nascent polypeptide. Thus, the trigger factor prevents aggregation of the emerging nascent chains by providing a competing hydrophobic environment (Baram *et al.*, 2005).

In D50S, protein L23 exposes a sticky hydrophobic patch, located in the wall of the ribosomal tunnel and available for interactions with hydrophobic regions of the progressing nascent chain. These interactions may be involved in co-translational folding of nascent polypeptides into secondary structures while still within the ribosomal tunnel, and such events may trigger signaling to the cell, for recruiting TF and initiating its binding. Thus, the subjection of L23 elongated loop may affect, in turn, its interaction with TF. Similar to the undetected conformational changes in the chimeric complex (Ferbitz *et al.*, 2004), mainly due to the disorder of the corresponding TFA region, the possible involvement of L23 loop in initial folding and/or FT attraction could not be seen in the chimeric complex, since like in eukaryotes, L23 of H50S lacks the elongate loop that penetrates the tunnel.

It seems, therefore, that protein L23 plays multiple roles in eubacteria. It is essential for the association of TF with the ribosome, and since the tip of its internal loop can undergo allosteric conformational changes thus modulating the shape and the size of the tunnel (Baram and Yonath, 2005; Baram *et al.*, 2005), it may control the pace of the entrance of the nascent chain into its shelter and serve as a channel for cellular communication with the nascent chain while progressing in the tunnel.

Antibiotics targeting the ribosomal tunnel

Antibiotics selectivity: the key for therapeutic effectiveness Ribosomes show a high level of universality in

sequence and almost complete identity in function, therefore the imperative distinction between pathogens and human, the key for antibiotics usefulness, is achieved by subtle structural difference within the antibiotics binding pockets of the prokaryotic and eukaryotic ribosomes (Auerbach *et al.*, 2004; Yonath and Bashan, 2004). Both *D. radiodurans* and *H. marismortui* are non-pathogenic organisms. Nevertheless, there are major differences between the suitability of their ribosomes to serve as pathogen models. Thus, although *D. radiodurans* is an extremely robust gram-positive eubacterium that can survive in harsh environments, it is best grown under conditions almost identical to those allowing for optimal biological activity of *E. coli* (Harms *et al.*, 2001) and shows striking sequence similarity to it. Moreover, contrary to archaeal and halophilic ribosomes, which possess typical eukaryotic elements at the principal antibiotics targets and are not inhibited by antibiotics at the clinically useful concentrations (Mankin and Garrett, 1991; Sanz *et al.*, 1993), *D. radiodurans* ribosomes are targeted by the common ribosomal antibiotics at clinically relevant concentrations in a fashion similar to most pathogens (Auerbach *et al.*, 2004; Schluenzen *et al.*, 2001). Thus, the availability of structures of antibiotics complexed with ribosomes from both species provides unique tools for investigating the structural basis for antibiotics selectivity.

A striking example is the immense influence of the minute difference between adenine and guanine in position 2058, which was found to dictate the affinity of macrolides binding. Macrolides are natural and semi-synthetic compounds, which rank highest in clinical usage. They are characterized by a macrolactone ring to which at least one sugar moiety is attached (Fig. 5). The first widely used macrolide drug is erythromycin, a 14-member lactone ring, decorated by a desosamine and cladinose sugars. Ketolides belong to a novel class of the macrolide family, characterized by a keto group at position 3 of the macrolactone ring, a single amino-sugar moiety and an extended hydrophobic arm (Fig. 5). This recently developed drug family was designed to act against several macrolide resistant bacterial strains. Both macrolides and ketolides were shown, crystallographically, to bind to a specific pocket in the eubacterial tunnel, called below the “macrolide binding-pocket”. Both act by producing a steric blockage of the ribosome exit tunnel, hence hampering the progression of nascent chains (Auerbach *et al.*, 2004; Berisio *et al.*, 2003a; 2003b; Hansen *et al.*, 2002b; Pfister *et al.*, 2004; 2005; Schluenzen *et al.*, 2001; 2003; Tu *et al.*, 2005; Yonath, 2005; Yonath and Bashan, 2004).

This high affinity pocket is composed of nucleotides belonging to the 23S RNA (Fig. 5A) and is located at the tunnel upper side, below the PTC and above the tunnel constriction (Fig. 4A). All currently available crystal structures of complexes of 14-membered ring macrolides with large subunits (Berisio *et al.*, 2003a; Schluenzen *et al.*, 2001; Tu *et al.*, 2005) show that the interactions of the

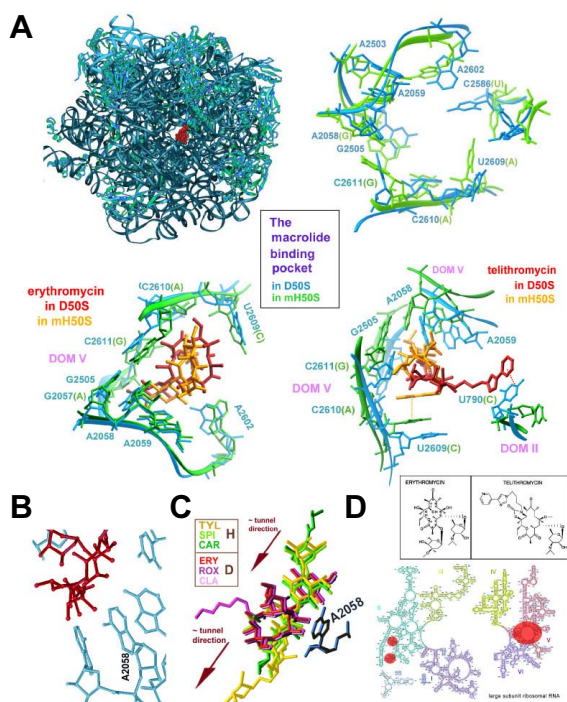


Fig. 5. The macrolide binding pocket. **A.** Top left: a view into D50S ribosome tunnel, with bound erythromycin (red). The ribosomal RNA and ribosomal proteins are shown in dark and light blue, respectively. Top right and bottom: the free, erythromycin and telithromycin bound pockets, in D50S (cyan) and H50S (green), highlighting the differences in sequence and orientation (green letters in parenthesis refer to the type of the nucleotide in *H. marismortui* if different from that of *D. radiodurans*). In bottom right, the stacking interactions between telithromycin and the binding pocket in both D50S and H50S, are shown by dotted lines. Note the superiority of tunnel (pocket) blocking in D50S, compared to mH50S. **B.** Zoom into D50S macrolide pocket (cyan), showing the close proximity between 2058 and the bound erythromycin (red). **C.** Superposition of the locations of three 16-membered macrolide tylosin (TYL), carbomycin (CAR) and spiramycin (SPI) bound to H50S, on the locations of three 14-membered macrolides, erythromycin (ERY), clarithromycin (CLA) and roxithromycin (ROX) bound to D50S, showing that the 16-membered macrolides should not severely hamper nascent protein passage. The location of A2058 and the approximate tunnel direction are also shown. Note the larger distance between the nucleotide at position 2058 and the desosamine sugars of the three 16-member macrolides, compared to the 14-member compounds. **D.** The two-dimensional diagram of the 23S RNA. The different RNA domains are encircled. The areas highlighted in light red are those involved in macrolide (only domain V) and ketolidase (both domains V and II). (Insert) The chemical compositions of erythromycin and telithromycin.

desosamine sugar and the lactone ring play a key role in macrolide binding. These contacts involve predominantly

the main constituents of the macrolide binding-pocket, namely nucleotides A2058-A2059 of the 23S RNA Domain V (Figs. 5A–5D). The second macrolide sugar, namely the cladinose, interacts directly with the ribosome only in a few cases (Berisio *et al.*, 2003a; Schluzzen *et al.*, 2001). Three closely related 14-membered macrolides, namely erythromycin and its semi-synthetic derivatives, clarithromycin and roxithromycin, exhibit exceptional consistency in their binding modes to the macrolide-binding pocket (Fig. 5C) (Schluzzen *et al.*, 2001). The high binding affinity of these macrolides was found to originate mainly from their interactions with nucleotide 2058. In all eubacteria this nucleotide is an adenine, and this adenine provides the means for prominent macrolide interactions. In eukaryotes, as well as in the archaeon *H. marismortui*, it is a guanine.

Consistently, these structures indicated that owing to increased bulkiness, a guanine in position 2058 should impose spatial constraints and hamper macrolide binding, in accord with the resistance mechanisms that are modifying the chemical identity of this nucleotide either by A → G mutation, or by *erm* methylation (Blondeau *et al.*, 2002; Courvalin *et al.*, 1985; Katz and Ashley, 2005; Poehlsgaard and Douthwaite, 2003; Sigmund *et al.*, 1984; Vester and Douthwaite, 2001; Weisblum, 1995). For over three decades it has been known that mutations in proteins L22 and/or L4 can also induce resistance to the 14-membered antibiotics (Davydova *et al.*, 2002; Pereyre *et al.*, 2002; Poehlsgaard and Douthwaite, 2003; Wittmann *et al.*, 1973). Although these proteins are rather close to the macrolide binding pocket, the structures of the macrolide complexes do not indicate a direct contact with these proteins. Nevertheless, the increase in A2508 size accompanied with the alterations in the tunnel conformation at its constriction, similar or identical to those seen crystallographically (Berisio *et al.*, 2003a) or by electron microscopy (Gabashvili *et al.*, 2001), could be correlated with this antibiotic-resistant mechanism. Thus, at its swung conformation, the tip of protein L22 hairpin loop, reached protein L4.

To circumvent the acute problems associated with macrolide resistance by modification of A2058, several new compounds have been designed. These include macrolide derivatives, in which the core macrolactone ring has been modified, to 15- (e.g. azithromycin) or 16- (tylosin, carbomycin A, spiramycin and josamycin) membered rings, all exhibiting activity against some MLS_B resistance strains (Bryskier *et al.*, 1993; Poulsen *et al.*, 2000). Ketolides present a yet another chemical approach, based on the addition of rather long extensions, such as alkyl-aryl or quinollyallyl, to the core macro lactone ring, expected to provide additional interactions, thus minimizing the contribution of 2058-9 region.

Drug binding to ribosomes with guanine at position 2058 may superficially indicate low level of selectivity, hence

should reduce its clinical relevance. This intriguing question triggered a through comparison between antibiotics binding modes to eubacteria, represented by D50S and to eukaryotes, represented by H50S. This comparison revealed a prominent differences in the effectiveness of tunnel blockage (Figs. 5A and 5C), which could be linked to the specific architecture of the two macrolide binding pockets.

Indeed, in H50S there are seven nucleotides that differ from the typical eubacteria, among them three present purine/pyrimidine exchange, and most of the conserved nucleotides have different conformations (Fig. 5A). Accordingly, the binding modes and consequently the therapeutical usefulness of macrolides that bind to H50S, namely the 16-membered ring compounds (Hansen *et al.*, 2002b) are considerably different from those found in D50S. Thus, in D50S the macrolides occupy most of the tunnel space, whereas in H50S the 16-membered ring macrolides lie almost parallel to the tunnel wall and consume a smaller part of it (Fig. 5C). These differences are likely to result from the sequence and conformational divergence of the macrolide binding pocket, in accordance with the low drug affinity to H50S, which forced the usage of immense excess (several orders of magnitude above the clinical levels) of antibiotics for obtaining measurable binding to H50S (Hansen *et al.*, 2002b; 2003), contrary to the usages of clinically relevant drug concentrations in the complexes of D50S (Auerbach *et al.*, 2004; Bashan and Yonath, 2005; Berisio *et al.*, 2003a; 2003b; Harms *et al.*, 2004; Schlunzen *et al.*, 2001; 2003; Yonath, 2005). Hence, the crystallographically observed differences in the antibiotics binding modes demonstrate the interplay between structure and clinical implications and illuminate the distinction between medically meaningful and less relevant binding.

A sole G → A mutation enabled macrolides binding

Further comparison, supporting the above conclusions, became possible as G2058 in *H. marismortui* 23S rRNA has recently been mutated to an adenosine (Tu *et al.*, 2005). This mutation (called below mH50S) increases macrolide binding affinity by 10000-fold, but does not significantly improve the effectiveness of the binding mode, as the magnitude of tunnel blockage in mH50S remains lower than that achieved by the same drug in the eubacterial D50S (Figs. 5A and 5C). Furthermore, based on azithromycin binding mode to mH50S (Tu *et al.*, 2005), the impressive gain in drug affinity, achieved by the G2058A mutation, is not accompanied by a comparable alteration in its binding mode compared to H50S wild-type, where 2058 is a guanosine. This seemingly surprising finding indicates that although 2058 identity determines whether binding occurs, the conformations and the chemical identities of the other nucleotide in the macrolide-pocket govern the antibiotics binding-modes and,

subsequently, the drug effectiveness.

Interestingly, all mH50S bound macrolides/ketolides share a similar macrolactone conformation, which is almost identical to that suggested by NMR studies to be of the lowest free energy at ribosome-free environments, therefore more likely to occur in vacuum or dilute solutions. These ribosome-free experiments ignored the ribosome, which by providing a significant interaction network, alters radically the drug environment. Hence, preservation of conformation of the drug in isolation is inconsistent with the high binding affinities between ribosomes and macrolides/ketolides. The preservation of the experimental ribosome-free macrolactone conformation in telithromycin-mH50S complex (Fig. 5A) supports the separation between binding and effectiveness. Thus, in mH50S telithromycin does not create the prominent interactions of ketolides with domains II (Fig. 5D), which are consistent with resistance data, and independently identified by footprinting, mutagenesis (e.g. Hansen *et al.*, 1999; Vester and Douthwaite, 2001; Xiong *et al.*, 1999), and crystallographic experiments, using the eubacterium *D. radiodurans* (Berisio *et al.*, 2003a; Schlunzen *et al.*, 2003). Likewise, high level of similarity between the binding modes of telithromycin and erythromycin is inconsistent with the profound differences detected between the susceptibility of A2058G ribosomes to ketolides, as compared with no influence on the susceptibility to macrolides (Pfister *et al.*, 2005).

The rationale behind the strange properties of the macrolides/ketolides binding modes to mH50S may be linked to the high salinity (> 2.5 M KCl) essential for *H. marismortui* optimal growth and for maintaining its integrity (Shevack *et al.*, 1985; Yonath, 2002). High salinity is also maintained within H50S crystals, although significantly lower from the optimal value. It is conceivable that in addition to the phylogenetic and conformational variability between archaea and eubacteria, which leads to dissimilarities between antibiotics conformations in D50S vs. H50S-mH50S (Baram and Yonath, 2005; Pfister *et al.*, 2005), the high salinity within the H50S (and mH50S) crystals masks potential ribosomal entities that could have interact with the drug. The similarity between the conformation of the macrolactone ring of unbound telithromycin, and the resemblance between the binding modes of erythromycin, azithromycin and telithromycin to H50S and/or mH50S (Tu *et al.*, 2005) support the notion that the high salinity in H50S crystals provides a semi ribosome-free environment to the bound drug, allowing it to maintain its conformation in ribosome-free environment.

To conclude, the A → G Mutation of 2058 in *H. marismortui* ribosome was found to be most beneficial for ribosomal-antibiotics research. It confirmed that 2058 is the key player in macrolide binding; it clarified the distinction between mere binding and antibiotics' inhibitory effectiveness; and it provided structural insight into the intriguing question,

which is also of utmost importance for drug development: What is the relevance of “minimum free-energy conformation” determined in ribosome-free environment to antibiotics binding and their therapeutic effectiveness? In other words: is there a correlation between the “minimum free-energy conformation” of a drug, determined in ribosome-free environment and its therapeutic effectiveness?

Future expectations Combating resistance to antibiotics has been a major concern in recent years. However, although pathogens resistance to antibiotics is believed to be the most severe problem in antibiotics usage, in attempts at combating it by novel design and/or by the improvement of existing antibiotics, the selectivity issue must play a key role. Impressive progress has been made by developing chemically improved (e.g. ketolides) as well as synergistic drugs. These open the gates for the introduction of further species specific anchors, thus increasing selectivity, and for providing alternative interactions, thus reducing the rate of the appearance of resistance. However, the battle is far from its end and that additional major effort is necessary.

The conclusions drawn from the crystallographic structures of the antibiotics complexes with bacterial ribosome provide indispensable tools for enhancement of the antibiotic efficiency. These structures show that the drugs' chemical properties govern its exact interactions, and that variations in drug properties appear to dominate the exact nature of seemingly identical mechanisms of drug resistance. Hence, the elucidation of common principles, combined with the variability in binding modes, including the discovery of a non-inactivating specific binding to the ribosome, justify expectations for the design of improved antibiotics properties by chemical modifications of existing compounds as well as by the design of novel drugs, based on the structural information.

Acknowledgments Thanks are due to Rita Berisio and to all members of the ribosome group at the Weizmann Institute for the constant assistance. X-ray diffraction data were collected at ID19/SBC/APS/ANL and ID14/ESRF-EMBL. The US National Inst. of Health (GM34360), the Human Frontier Science Program Organization (HFSP: RGP 76/2003), and the Kimmelman Center for Macromolecular Assemblies, provided support. AY holds the Martin and Helen Kimmel Professorial Chair.

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