Atomic Structures of the 30S Subunit and Its Complexes with Ligands and Antibiotics

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The two subunits that make up the ribosome have both distinct and cooperative functions. The 30S ribosomal subunit binds messenger RNA (mRNA) and is involved in the selection of cognate transfer RNA (tRNA) by monitoring codon–anticodon base-pairing during the decoding process. Both subunits work in concert to move tRNAs and mRNAs relative to the ribosome in translation, and both are the target of a large number of naturally occurring antibiotics. Thus, useful information about the mechanism of translation can be gleaned from structures of both individual subunits and the intact ribosome. In this paper, we describe our work on the determination of the atomic structure of the 30S ribosomal subunit and its complexes with RNA ligands, antibiotics, and initiation factor IF1. The results provide structural insights into how the ribosome recognizes cognate tRNA and discriminates against near-cognate tRNA. They also provide a structural basis for understanding the action of various antibiotics that target the 30S subunit.

Knowledge of the atomic structure of the 30S is the culmination of decades of work on its structure and function. The state of the ribosome field two years ago has been summarized in a recent book (Garrett et al. 2000). The initial breakthrough in the goal toward a crystal structure of the ribosome was the crystallization of 50S subunits suitable for diffraction studies by Yonath and coworkers (Yonath et al. 1980). Although these original crystals did not diffract very well, over the course of several years, improved crystals of the 50S subunit were obtained that eventually resulted in diffraction to 3 Å resolution (von Böhlen et al. 1991). Another important advance was the introduction of cryoelectron microscopy that the 30S subunit crystals in MPD diffracted to about 10–12 Å resolution initially (Yonath et al. 1988; Yusupov et al. 1988), and by 1995, their diffraction limit had only improved to about 7–8 Å resolution (Schluenzen et al. 1995). However, by that time, advances in single-particle reconstruction methods had shown that the 30S subunit was conformationally different in the active, inactive, and 50S-bound states (Frank et al. 1995; Lata et al. 1996).

CRYSTALLIZATION OF THE 30S SUBUNIT

Crystals of both the 30S subunit and 70S ribosomes from Thermus thermophilus that were suitable for structural studies were first reported by the Puschino group (Trakhanov et al. 1987; Yusupov et al. 1987). These crystals used 2-methyl-2,4-pentanediol (MPD) as the precipitant, and, shortly afterward, crystals of the 30S in a mixture of ethanol and ethylbutanol were reported by the Yonath group (Glotz et al. 1987). The 30S subunit crystals in MPD diffracted to about 10–12 Å resolution initially (Yonath et al. 1988; Yusupov et al. 1988), and by 1995, their diffraction limit had only improved to about 7–8 Å resolution (Schluenzen et al. 1995). However, by that time, advances in single-particle reconstruction methods in cryo-electron microscopy had shown that the 30S subunit was conformationally different in the active, inactive, and 50S-bound states (Frank et al. 1995; Lata et al. 1996). This suggested to us that conformational variability of the 30S subunit could be the reason for the poor diffraction reported for its crystals, and we reasoned that, by analogy with smaller enzymes, binding of substrates or cofactors might result in complexes that were conformationally homogeneous and therefore yield better crystals.

As a starting point, we used the original crystallization conditions reported by the Puschino group. To our surprise, a straightforward optimization of the original conditions resulted in crystals that diffracted to beyond 3 Å at third-generation synchrotron sources (Clemons et al. 1999; Wimberly et al. 2000). Unlike improved crystals of the 30S subunit produced by the Yonath group (Yonath et al. 1998; Tocilj et al. 1999), our crystals do not require soaking in tungsten clusters and heat treatment to result in diffraction to high resolution, and the reason they diffract so well will become apparent below.

STRUCTURE DETERMINATION

Just three years ago, it was not clear that the crystallographic phase problem could be solved for asymmetric units as large as a ribosomal subunit. It has long been known that heavy atom clusters of tungsten or tantalum could be used for obtaining phase information for large molecules (Blundell and Johnson 1976). They were used in the structure determination of large complexes such as...
ribulose biphosphate carboxylase and the nucleosome core particle (O’Halloran et al. 1987; Andersson et al. 1989; Knablein et al. 1997). The use of clusters was also suggested for ribosomes (Thygesen et al. 1996). The first concrete demonstration that such clusters could yield phase information for ribosome-sized problems came when they could be visualized directly in difference-Patterson maps of the 50S subunit and resulted in electron density maps that showed unambiguous right-handed helices corresponding to A-form RNA duplexes (Ban et al. 1998).

At the same time, anomalous scattering has seen increasing use in structure determination, especially through the use of the multi-wavelength anomalous diffraction (MAD) technique on crystals of proteins labeled with selenomethionine (Hendrickson et al. 1990; Hendrickson 1991). It was shown by us that treatment of MAD as a special case of standard multiple isomorphous replacement (MIR) resulted in excellent phases despite the nominally very small signal from anomalous scattering (Ramakrishnan et al. 1993; Ramakrishnan and Biou 1997). A test calculation shows that a fully selenomethionylated ribosome would have too little signal to be useful (Clemons et al. 2001). However, the LII edges of many elements typically have 3–4 times the anomalous scattering of the K-edge of selenium, and were previously used for both protein (Kahn et al. 1985; Weis et al. 1991) and RNA (Cate et al. 1996) structure determination. Calculations based on the original equation by Crick and Magdoff (1956) showed that even 10–20 sites per 30S subunit would result in a measurable signal (Clemons et al. 2001), offering an alternative route to the phasing problem. This conclusion appears to have been reached independently by the various groups working on ribosome crystallography. Our atomic structure of the 30S, as well as that of the 50S (Ban et al. 2000) and the 70S (Cate et al. 1999), were all solved primarily using LII edges of compounds such as osmium or iridium hexammine, although in each case, heavy atom clusters were very useful in the determination of low-resolution phases.

The structure of the 30S in our laboratory was determined in two steps: Initially, a 5.5 Å resolution structure was determined, which implicitly described the correct molecular packing in the crystal (Clemons et al. 1999). Even at this resolution, we were able to interpret about a third of the structure because of the large amount of prior structural and biochemical data on the 30S subunit. We were able to place proteins whose structures had been determined in isolation, deduce the fold and location of the previously unknown protein S20, and determine the fold of the entire central domain of 16S RNA and the relative location of the functionally important helix 44 along the interface of the subunit.

Last year, we published the complete atomic structure of the 30S subunit at 3 Å resolution contains 1513 out of 1521 nucleotides of 16S RNA and all 20 polypeptide chains present in the crystal (S2-S20, THX), with 98% of the sequence built into the molecule (Wimberly et al. 2000). The missing regions occur exclusively at the termini of the RNA and proteins.

The secondary structure of Thermus 16S RNA is very close to that predicted on the basis of phylogenetic comparisons (Gutell 1996), which is shown in Figure 1 along with the standard Brimacombe numbering for the various double helical elements (Glotz and Brimacombe 1980; Mueller et al. 1997). Figure 1 also indicates the insertions and deletions relative to the Escherichia coli 16S RNA sequence, and both sets of numberings are shown to aid biochemical analysis. The secondary structure domains of 16S RNA form distinct domains in three dimensions (Fig. 2a), so that the 5’ domain forms the body, the central domain forms the platform, the 3’ major domain forms the head, and the 3’ minor domain consists largely of two helices that lie on the intersubunit interface. This is different from the structure of 23S RNA, where the division of secondary structure domains appears to be more arbitrary and where the secondary structure domains are closely intertwined in three dimensions. The three-dimensional structure of 16S RNA has been described briefly (Wimberly et al. 2000), and a detailed analysis is in progress.

The shape of the 30S is very similar to that seen by electron microscopy. In addition to the classic features of the head, shoulder, and platform, one can also see the “beak” and “spur” that were observed using cryoelectron microscopy (Frank et al. 1995; Stark et al. 1995). From a comparison of the various electron microscopic images of the 30S subunit (Frank et al. 1995; Lata et al. 1996), it appears that the crystal structure is closer to the 50S-bound form than to either the active or inactive conformations of the isolated subunit.

The overall shape and gross morphological features of the 30S subunit are largely determined by the RNA component (cf. Figs. 2a and 2c). The RNA contains more than 50 regular helices. Many of the regions that were considered to be single-stranded loops in the secondary structure are in fact irregular extensions of regular double helices, so the molecule can be thought of in three dimensions as largely helical. Neighboring helical elements are often coaxially stacked; e.g., helices 16 and 17 in the 5’ domain or helices 21, 22, and 23 in the central domain. These coaxially stacked helices are packed using primarily three types of interactions, all of which use the minor groove as the packing interface (Wimberly et al. 2000). A notable feature is the use of highly conserved adenines to make tertiary interactions by binding in the minor groove of an adjacent double-helical section of RNA. As described below, this feature is also used in decoding.

The proteins mainly decorate the back and periphery of the 30S subunit (Fig. 2b), whereas the interface surface consists largely of ribosomal RNA (Fig. 2c). A glaring exception to this rule is S12, which is located near the decoding site at the interface. Proteins S13 and S19 also lie mainly on the 50S side of the head.

Many of the proteins have highly elongated extensions or internal loops (Fig. 3). A particularly dramatic example is S12, which is the only protein that lies directly at the intersubunit interface, but has a long amino-terminal

**GLOBAL ARCHITECTURE**

The final refined atomic model of the 30S subunit at 3 Å resolution contains 1513 out of 1521 nucleotides of 16S RNA and all 20 polypeptide chains present in the crystal (S2-S20, THX), with 98% of the sequence built into the molecule (Wimberly et al. 2000). The missing regions occur exclusively at the termini of the RNA and proteins.
extension that threads through the 30S subunit and makes contact with S8 and S17 on the back side. In general, these extensions are highly basic, make intimate contacts with ribosomal RNA, and help with the assembly and folding of the 30S subunit. Similar extensions have also been observed in the 50S structure (Ban et al. 2000).

Figure 1. Secondary structure diagram of 16S RNA from *Thermus thermophilus*, colored by domain (5’ domain red, central domain green, 3’ major domain orange, and 3’ minor domain blue). Insertions in the *Thermus* sequence relative to *E. coli* occur in the 5’ and 3’ major domains and are shown in green. Deletions are indicated with black tick marks and a “Δn” to show the deletion of n nucleotides. The *T. thermophilus* numbering is shown by tick marks for every 10th nucleotide and labels for every 50 nucleotides, colored as for each domain. Corresponding *E. coli* numbering and tick marks are shown in black throughout. (Modified, with permission, from http://www.rna.icmb.utexas.edu; see Gutell 1996.)
Figure 2. (a) Tertiary structure of 16S RNA in the 30S subunit, colored according to domains as in Fig. 1. (b, c) Overviews of the back and interface sides of the 30S subunit, respectively. (Panel a reprinted, with permission, from Wimberly et al. 2000. Copyright Nature.)
Interestingly, proteins characterized as primary binders in the 30S assembly map are all globular and lack the long extensions. These proteins generally bind at multihelix junctions, such as S4 at the 5-way junction of helices 3, 4, 16, 17, and 18; S7 at the junction of helices 29, 30, 41, and 42; and S15 at the junction of helices 20, 21, and 22. The primary binders are very likely to be involved in condensing 16S RNA around multihelix junctions in agreement with a great deal of prior data (Nowotny and Nierhaus 1988; Agalarov and Williamson 2000). Finally, proteins with similar or even identical folds, such as S6, S10, and S11, bind RNA in different ways, suggesting that it is difficult to make inferences about RNA binding from topology.

**COMPARISON WITH AN INDEPENDENT 30S STRUCTURE**

The same month that our structure was published, an independent 3.3 Å structure of the 30S subunit from the Max Planck/Weizmann group was reported (Schluenzen et al. 2000). A comparison reveals a very similar overall conformation of the 30S subunit in the two structures. Thus, there appears to be little evidence to support the claim that heating in the presence of tungsten clusters leads to a functionally more active form of the particle (Tocilj et al. 1999; Schluenzen et al. 2000). Although the global path of the RNA backbone is in agreement with ours, there are many significant differences in the details. In the Max Planck/Weizmann structure, regions of RNA such as the 560 and 967 loops are missing, irregular conformations of RNA are often modeled differently, and the RNA registry differs from ours in several regions, with these regional differences accounting for about 300 out of 1500 nucleotides. The root mean square deviation (rmsd) between equivalent phosphorus atoms in these regions of discrepancy is 6.1 Å compared to 3.6 Å for the whole RNA. In the Max Planck/Weizmann structure, the previously unsolved proteins are modeled as Ca traces with unidentified residues and are often noncontiguous and incomplete. Major differences in proteins fall into the categories of missing domains (e.g., S2, S4), missing extensions or loops (e.g., S7, S9, S13), or incompatible topologies (e.g., S11, S12, S17). Part of their S3 corresponds to our S14, and S20 has completely reversed polypeptide chain direction. We attribute most of these discrepancies to differences in interpretation of the electron density rather than genuine differences in conformation. Support for this view comes from the observation that recent structures of the 30S subunit published by the Max Planck/Weizmann group are in very close agreement with the structure we originally published about 8 months earlier, with an rmsd between phosphorus atoms of about 1 Å (Pioletti et al. 2001). Any differences that
remain could be the result of slightly different crystallization conditions as well as the fact that tungsten clusters are part of the Max Planck/Weizmann structure, but not of ours.

**BINDING SITES FOR tRNA AND mRNA**

A precise analysis of the tRNA- and mRNA-binding sites was made easier by the occurrence of two fortuitous interactions. The spur (helix 6) from a neighboring molecule in the crystal is inserted into the P site of the 30S subunit and mimics P-site tRNA (Carter et al. 2000). Interestingly, the 3’ end of 16S RNA is folded back into the mRNA-binding cleft, and nucleotides are visible in both the P- and the E-site codons. The terminal three bases of this “pseudo-message” form noncanonical base pairs with the “anticodon” from the spur. Thus, the form of the 30S we crystallized appears to have mimics for tRNA in the P site and mRNA in the P and E sites. In hindsight, it is clear why this crystal form diffracts well even in the absence of bound ligands: It effectively mimics a ligand-bound form through crystal contacts. It is also interesting that in the absence of added mRNA, the 3’ end of 16S RNA is folded back into the 30S cleft in crystals of 70S ribosomes also (see Noller et al., this volume). This suggests that the binding of the 3’ end of 16S RNA into its own mRNA-binding cleft in the 30S subunit is not just a crystal-packing artifact. Rather, the intramolecular burial of the 3’ end of 16S RNA into the message-binding cleft may be energetically more favorable than binding mRNA for entropic reasons, except when extensive additional compensating interactions are made with mRNA; e.g., in base-pairing of the 3’ end with the Shine-Dalgarno sequence. This feature may therefore have the benefit of making initiation more specific.

The 7.8 Å structure of the 70S with mRNA and tRNA identified two elements of the 30S subunit, helix 27 and helix 44 (Cate et al. 1999). Using these two elements, we were able to superimpose the 70S structure onto our atomic model for the 30S subunit to arrive at a model for the interaction of tRNA with the 30S subunit (Fig. 4a). The superposition provided additional evidence that the spur and 3’ end of 16S RNA were good mimics of P-site tRNA and mRNA, respectively, as discussed above. In addition, it allowed us to deduce details of the interactions of tRNA and mRNA with the 30S subunit (Carter et al. 2000). As an example, details of the interaction of the P-site tRNA mimic and codon with the 30S subunit are shown in Figure 4b.

As had been revealed by previous biochemical data, the P-site tRNA makes extensive interactions with the 30S subunit. On the other hand, the A site is shallow and wide, which is consistent with both its lower affinity for tRNA and the need to allow tRNA to rotate significantly from its initial conformation as part of the ternary complex with EF-Tu (Stark et al. 1997) to its eventual conformation after GTP hydrolysis and accommodation, in which the acceptor arm of tRNA swings into the peptidyl transferase site. Additionally, the structure shows that helix 44, helix 34, protein S12, and the 530 loop are all part

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**Figure 4.** tRNA-binding sites of the small subunit. (a) Overview of the three tRNA molecules modeled onto the 30S structure using the 7.8 Å structure of 70S in complex with tRNA and mRNA. The two 16S elements used for the superposition (helices 44 and 27) are shown in green. (b) Details of the interactions in the P site made by the spur (helix 6) from a neighboring subunit in the crystal that mimics tRNA. The 3’ end of 16S RNA is folded into the cleft of the 30S and mimics mRNA (dark blue). The spur is shown in red, and the elements of 16S RNA and ribosomal proteins are labeled and colored independently for clarity. Dotted yellow lines indicate base-pairing between the spur and the codon mimic (dark blue), and dotted black lines indicate hydrogen bonds. Bases with black filling have previously been implicated in P-site tRNA or mRNA binding. (Reprinted, with permission, from Carter et al. 2000 [copyright Nature] http://www.nature.com.)
of the A site where decoding occurs. This confirms the suggestion by Brimacombe and coworkers based on cross-linking experiments with mRNA that these elements are close to the decoding site (Dontsova et al. 1992).

The tRNA-binding sites have several striking features that were previously either unsuspected or only hinted at. Whereas the A and P sites consist mainly of RNA elements, the E-site tRNA includes extensive interactions with proteins S7 and S11. A second unexpected finding was that long tendrils of extended polypetide chains from S9 and S13 make their way into the tRNA-binding sites. In particular, the lengths of the S9 tail and its terminal lysine residue are highly conserved, suggesting that it makes a crucial interaction with tRNA.

The tRNA interactions deduced from the 30S structure are consistent with a large body of biochemical data and have subsequently been confirmed by the 5.5 Å resolution structure of the 70S ribosome with mRNA and tRNA (Yusupov et al. 2001), as well as our complex of mRNA and an anticodon stem-loop of tRNA with the 30S subunit (Ogle et al. 2001).

ROLE OF THE 30S SUBUNIT IN DECODING

A central role of the 30S is decoding, or selection of the correct aminoacylated tRNA for participation in peptidyl transferase. The discrimination inherent in codon–anticodon base-pairing leads to a free-energy difference of only 2–3 kcal/mole between cognate tRNA and near-cognate tRNA (which has a single mismatch). This difference would predict an error rate of about 10−2, which is 1–2 orders of magnitude higher than the observed error rate of translation (Kurland 1992).

Ever since the discovery that antibiotics can increase the error rate of translation, it has been proposed that the ribosome can modulate the fidelity of translation by directly monitoring codon–anticodon base-pairing (Davies et al. 1964). The site where this monitoring occurs is referred to as the decoding site, and by definition, it must be part of the environment of A-site tRNA. Two possible mechanisms have been proposed to account for the improved accuracy. The first involves recognition by the ribosome of the geometry of correct codon–anticodon base-pairing in the same way that enzymes recognize the geometry of the correct substrate and discriminate against incorrect ones (Eigem and De Maeyer 1966; Potapov 1982; Thompson and Dix 1982). In an alternative mechanism, selection of the correct tRNA would be based on a kinetic proofreading scheme, in which the energy discrimination inherent in codon–anticodon pairing is used twice, by having an initial selection and a “proofreading” step that are separated by an irreversible step (Hopfield 1974; Ninio 1975). In the case of the ribosome, the irreversible step would be the hydrolysis of GTP by EF-Tu. Thus, an error rate of 10−2 at each step could result in a total error rate as low as 10−4 if each step was allowed to reach equilibrium. Because of the need for speed, the overall accuracy is reduced from the maximum possible, but it is still sufficient to account for the error rate of protein synthesis. Therefore, in the context of proofreading schemes, it is not clear that recognition of codon–anticodon pairing geometry by the ribosome is even necessary. However, when a slowly hydrolyzable GTP analog was used, it was shown that even a single selection step could theoretically have a discrimination sufficient to account for the accuracy of protein synthesis, provided equilibrium could be reached, which gave evidence for the ability of the ribosome to recognize the geometry of codon–anticodon base-pairing (Thompson and Karim 1982).

LOCATION OF THE DECODING SITE

Where is the decoding site in the 30S subunit? Data from footprinting (Moazed and Noller 1990), cross-linking (Dontsova et al. 1992), and genetics (O’Connor et al. 1995, 1997) have implicated the region that includes helix 44, the 530 loop, and helix 34 in the ribosomal A site. These regions are all close together in the structure. More recent work on decoding has focused on two universally conserved residues, A1492 and A1493, which are part of an internal loop in helix 44. These bases are essential for both viability and A-site tRNA binding (Yoshizawa et al. 1999). This internal loop is also the binding site for the antibiotic paromomycin and related aminoglycosides, which increase the error rate of protein synthesis. In a nuclear magnetic resonance (NMR) structure of paromomycin bound to a fragment of helix 44, it was shown that the drug displaces A1492 and A1493 toward the minor groove of helix 44 (Fourmy et al. 1996). On the basis of biochemical experiments, it was concluded that the N1 of the two bases directly interacted with the 2′ OH of mRNA (Yoshizawa et al. 1999), and that the bases were involved in recognition of the shape of the codon–anticodon helix via minor-groove interactions. However, the same data could be used to arrive at a completely different model that involved interaction of the codon–anticodon helix with the major groove of helix 44 (VanLoock et al. 1999). In any case, when the 7.8 Å structure of the 70S ribosome with tRNA was determined, modeling the NMR structure of the internal loop with or without paromomycin into the 70S electron density suggested that the adenines were too far away from the codon–anticodon helix to have a direct role in decoding (Cate et al. 1999).

These discrepancies began to be resolved when the structure of the 30S subunit complexed with spectinomycin, streptomycin, and paromomycin showed that rather than displace them into the minor groove in a way that would preserve the A1493–A1408 base pair reported in the NMR structure (Fourmy et al. 1996), paromomycin in fact flipped A1492 and A1493 completely out of the internal loop of the helix so that they pointed into the A site (Carter et al. 2000). Modeling of A-site tRNA and mRNA showed that these bases would be in a position to bind directly in the minor groove of the codon–anticodon helix. However, they would do so in a way that would place the N1 of the adenines pointing toward tRNA rather than interacting with the 2′ OH of mRNA, as had previously been proposed (Yoshizawa et al. 1999). There are
several examples of highly conserved adenines in 16S RNA that form tertiary interactions with the minor groove of A-form helices. These interactions involve hydrogen bonding of the adenines with the 2′ OH of both strands of the minor groove in a manner that would be sensitive to the shape and width of the groove. On the basis of this analysis, it was proposed that the binding of cognate tRNA would induce a similar conformational change in A1492 and A1493 to the one induced by paromomycin. The energetic cost of the conformational change would be paid for by compensating interactions between the adenines and the codon–anticodon helix. In the case of near-cognate tRNA, the fit would not be perfect, so that the induced changes would not be energetically favorable. However, paromomycin, by partially paying the energetic cost of the induced changes, would promote the binding of near-cognate tRNA and thus lead to an increased error rate.

**RECOGNITION OF CODON–ANTICODON BASE-PAIRING BY THE RIBOSOME**

The interactions by which the ribosome senses codon–anticodon pairing at the A site were determined experimentally by directly soaking mRNA in the form of a uridine hexanucleotide, as well as the anticodon stem-loop of its cognate phenylalanyl-tRNA, into crystals of the 30S subunit (Ogle et al. 2001). This work directly revealed the codon and anticodon stem-loop in the A site. In doing so, it showed that our earlier modeling of the A-site tRNA was reasonably accurate, and also that a sharp kink exists in the RNA between the A- and P-site codons, as could be expected if each were to be part of an A-form helix with its corresponding anticodon.

The binding of cognate tRNA induces global domain movements in the 30S subunit. It also causes local changes around the decoding site. In the absence of tRNA or paromomycin, A1492 and A1493 are stacked inside helix 44, and G530 is in the syn conformation (Fig. 5a). The binding of paromomycin causes A1492 and A1493 to flip out of the internal loop of helix 44, while leaving G530 in the syn conformation (Fig. 5b). On tRNA binding, A1492 and A1493 flip out of helix 44 as we had predicted, but also cause G530 on the other side of the codon–anticodon helix to switch from a syn to an anti conformation (Fig. 5c). A comparison of Figures 5b and 5c shows that, apart from having no effect on G530, paromomycin alone does not induce precisely the same conformational changes in A1492 and A1493. This suggests that the antibiotic facilitates incorporation of near-cognate tRNA by contributing in part toward the energetic cost of changes required for recognition, rather than by creating a fully preformed binding site. Since the adenines are largely disordered in the native structure as judged by a very high B factor, but well ordered in the complexes with paromomycin or tRNA, there must be some additional entropic cost in the changes induced by binding.

A1493 makes intimate contacts with the first base pair of the codon–anticodon helix (Fig. 6a), whereas A1492 and G530 together span the minor groove of the second base pair and interact with each other via their N1 positions (Fig. 6c). The third base pair is not as closely monitored, with only the codon nucleotide making interactions with the ribosome (Fig. 6d). This is consistent with the requirement of the genetic code that the third position be able to accommodate noncanonical base pairs such as a GU wobble (Crick 1966; Yokoyama and Nishimura 1995). Moreover, the observation that the ribosome makes extensive interactions with the codon base but not the anticodon base is consistent with previous analysis suggesting that the codon base conformation is more restricted than that of the anticodon (Yokoyama and Nishimura 1995). These experiments show how the ribosome recognizes the geometry of codon–anticodon base-pairing, consistent with the slower requirement at the

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**Figure 5.** Overview of A-site tRNA binding. Discrete states of the 30S A site deduced from three different crystal structures of (a) native, unbound 30S, (b) the 30S in complex with paromomycin, and (c) the 30S in complex with mRNA and cognate tRNA. In all panels, tRNA ASL is gold, A-site mRNA codon purple, P-site mRNA codon green, protein S12 brown, and important bases involved in conformational changes red. Magnesium ions are shown as magenta spheres. (a) The A site of the native 30S subunit (Wimberly et al. 2000). A1492/3 (red, right) are stacked in the center of helix 44, and G530 is in its syn conformation. C1054 is shown in the upper left corner. (b) In the paromomycin-bound 30S (Carter et al. 2000), the bases of A1492 and A1493 have been pushed out into the A site by the antibiotic binding to the center of helix 44 (yellow sticks, lower right). (c) In the 30S:ASL:U6 complex, A1492/3 flip out to monitor the codon–anticodon interaction and G530 has switched to its anti conformation to stabilize the interaction of the second position. Two Mg ions are visible in the place where A1492/3 are located in the native structure. (Reprinted, with permission, from Ogle et al. 2001 [copyright AAAS] http://science.org.)
wobble position. The structure explains previous chemical modification data and also rationalizes the requirements for adenines at 1492 and 1493 and a guanine at 530. Only purines would be able to span a minor groove, but a guanine at 1492 or 1493 could not have the same conformation as either adenine because of a steric clash with the minor groove from its NH2 at the N2 position. Guanines also would not be able to make the analogous hydrogen bonds through their N1 positions. Similarly, only guanine at G530 would be able to form the observed bond with A1492 through the N1 position. Finally, because DNA has no 2’ OH group and, also, DNA–RNA hybrids have a different groove width and shape, the structure explains the classic observation that DNA is a poor template for translation but is less so in the presence of the aminoglycoside neomycin (McCarthy and Holland 1965). It also explains the inability of the ribosome to bind tRNA in the A site when the codon consists of DNA rather than RNA (Potapov et al. 1995).

Recently, we have also determined the structure of the anticodon stem-loop of near-cognate tRNA in complex with the 30S subunit. This near-cognate tRNA, which codes for leucine, has a GAG anticodon, and hence has a GU wobble in both the first and third positions using the poly-U mRNA. In the absence of paromomycin, no density is visible for tRNA in the A site, whereas weak density is visible for mRNA. In the presence of paromomycin, the structure is very similar to that of cognate tRNA. However, because the first base pair of the codon–anticodon helix is a GU wobble pair rather than a Watson-Crick base pair, the fit of A1493 with the minor groove of this base pair is distorted and hence energetically less favorable. Interestingly, in comparison with the cognate case, it is the codon base rather than the anticodon that has moved to accommodate the wobble pair at the first position (Fig. 6b).

These experiments show that the ribosome recognizes cognate tRNA through an induced-fit mechanism that monitors the geometry of the minor groove of codon–anticodon base-pairing. In doing so, the ribosome uses principles found in other systems. The interactions of A1493 and A1492 with the minor groove are identical to those classified as type I and type II A minor interactions (Nissen et al. 2001), which are found in tertiary interactions in the group I intron, and both 23S RNA and 16S RNA. Much of the free energy of the interaction ap-
pears to arise from the shape complementarity of the adenosines to the minor groove (Doherty et al. 2001). The principles of minor groove recognition by surface complementarity and sequence-independent hydrogen bonding to ensure fidelity of base-pairing are also employed by DNA and RNA polymerases (Doublie et al. 1998; Kiefert et al. 1998; Cheetham and Steitz 1999).

Since our experiments involve just the anticodon stem-loop of tRNA in the context of the 30S subunit, a natural question concerns what state these structures represent along the reaction pathway. The conformation of the anticodon stem-loop is virtually identical to that of the A-site tRNA in the 70S structure (Yusupov et al. 2001), suggesting that it represents a post-accommodation state of tRNA. Our structure therefore has no direct bearing on the initial recognition step in which a ternary complex of EF-Tu with tRNA and GTP binds to the ribosome. However, we note that the protections observed on the binding of the ternary complex of EF-Tu with tRNA and GTP binds to the ribosome. However, we note that the protections observed on the binding of the ternary complex of EF-Tu with tRNA and GTP to the ribosome are essentially the same as those observed on nonenzymatic tRNA binding (Powers and Noller 1994), and both are consistent with our structure. This suggests that despite a large change in the orientation of the tRNA, many of the interactions of the ribosome with the codon-anticodon helix observed here probably occur in the initial recognition step and persist through GTP hydrolysis to accommodation. It is clear that A1492 and A1493 are in a conformationally variable region of the 16S RNA (Lodmell and Dahlberg 1997). Chemical probing studies show that ram and restrictive ribosomes have altered chemical reactivity in portions of helices 27 and 44 of 16S RNA (Allen and Noller 1989). More recent chemical probing (Lodmell and Dahlberg 1997) and cryo-electron microscopy (Gabashvili et al. 1999) provide evidence that the ram and restrictive ribosomes may have altered conformations throughout the 30S subunit.

A model has been proposed in which the ribosome switches between ram and restrictive states even during normal translation (Lodmell and Dahlberg 1997). After hydrolysis of GTP by EF-Tu, the ribosome enters a restrictive conformation that favors dissociation of tRNAs. This is followed by a transition to another state in which the tRNA is stably bound and can become available for peptide-bond formation. Near-cognate tRNAs are more likely to dissociate from the restrictive conformation, whereas cognate tRNAs are more likely to progress to peptide-bond formation. Restrictive mutations alter the energetics of the ribosome so that the lifetime of the restrictive state is prolonged, and hence incorrect tRNAs are more likely to be rejected. In contrast, ram mutations “destabilize” this restrictive state, so that more tRNAs (even near-cognate ones) are likely to stay bound to the ribosome.

The structure of the 30S subunit shows the locations of the streptomycin-binding site as well as mutations that affect translational accuracy. The environment of helix 27 in the 30S crystal structure is shown in Figure 7a. The bases with altered chemical reactivity in the ram and restrictive states (Allen and Noller 1989) lie at the interface between helix 27 and helix 44. In our structure, bases 888–890 of helix 27 are present in the S-turn conformation thought to be characteristic of the ram state (Lodmell and Dahlberg 1997). As shown in Figure 7b, the ram mutations that occur in S4 and S5 lie mainly at the interface between the two proteins (Clemons et al. 1999; Wimberly
et al. 2000). This interface is contiguous with an RNA–RNA interface between helix 1 and the 530 loop (helix 18). Together, they represent a hybrid protein–RNA interface that could be disrupted during conformational changes in the 30S subunit. Streptomycin binds tightly in a pocket where it interacts with the phosphate backbone from four regions of 16S RNA (helices 1, 18, 27/28, and 44) as well as the highly conserved Lys-45 (E. coli Lys-42) of S12 (Fig. 7c) (Carter et al. 2000). All of these regions lie at the interface between the shoulder and platform domains of the 30S subunit (Fig. 7d).

Together, these observations implicate movement at this interface in the conformational changes associated with proofreading. At one end of the interface, mutations in S4 and S5 that disrupt their interaction lead to the \textit{ram} phenotype, probably by destabilizing the restrictive state and thus accelerating the transition to a \textit{ram} state. In contrast, mutations in S12 that disrupt interactions at the other end of the interface probably prolong the lifetime of the restrictive state, whereas streptomycin, which stabilizes interactions in the S12 region, suppresses the effect of these mutations (Kurland et al. 1996). In the extreme case of streptomycin-dependence mutations in S12, the transition to the \textit{ram} state is so unfavorable that streptomycin is actually required for function. Interestingly, the single known mutation in S12 that leads to streptomycin resistance without a hyperaccurate phenotype is K45R (E. coli K42). This lysine makes direct hydrogen bonds with two OH groups on streptomycin as well as a salt bridge to A913. Its mutation to an arginine would have the effect of disrupting streptomycin binding but would not disrupt its interactions with the ribosome, since the arginine could make the same salt bridge to A913.

Without knowing the structure of the restrictive and \textit{ram} states, it is difficult to predict how streptomycin and accuracy mutations would affect the transition between
them. Moreover, it remains unclear how the changes in conformation observed in the mutants relate to conformational changes in the ribosome during the elongation cycle. In addition to structural work, the ram and restrictive states need to be placed rigorously in a kinetic pathway from initial selection to peptidyl transferase. Although the details of proofreading remain to be elucidated, the movement between the platform and shoulder domains of the 30S subunit that has been observed recently on cognate tRNA binding or IF1 binding (Carter et al. 2001; Ogle et al. 2001) supports the proposal that movement between these domains is involved.

**COMPLEXES WITH ANTIBIOTICS**

Many antibiotics act by binding to the large or small ribosomal subunit, and an enormous amount of work has gone into elucidating the biochemical basis for their action (Spahn and Prescott 1996). Most antibiotics appear to bind to the RNA component of the ribosome, although many resistance mutations map to ribosomal proteins.

Despite decades of biochemical work, it has not been possible until recently to obtain direct structural information on the binding of antibiotics to ribosomes because of the lack of a high-resolution structure of either subunit. A notable exception was the site in helix 44 that contains an internal loop implicated in decoding. This region was shown to bind antibiotics as an isolated fragment (Purohit and Stern 1994) and was used to determine the structures of paromomycin and gentamicin in complex with RNA (Fourmy et al. 1996; Yoshizawa et al. 1998).

With the determination of atomic structures for ribosomal subunits, the situation has changed dramatically. It is now possible to determine the structures of complexes of antibiotics with either ribosomal subunit by co-crystallization or by soaking the native crystals in antibiotic. Refinement of the coordinates of the native structure against diffraction data from the antibiotic complex directly reveals density corresponding to the antibiotic molecules in difference Fourier maps. Using this approach, we have determined the structure of the 30S subunit in complex simultaneously with streptomycin, spectinomycin, and paromomycin (Carter et al. 2000) and also separately with tetracycline, hygromycin B, and pactamycin (Brodersen et al. 2000). The locations of these antibiotics in the 30S structure are shown in Figure 8. These structures shed light on the mechanism of action of the antibiotics, as discussed above for paromomycin and streptomycin in the context of decoding and proofreading. We briefly mention below the results for the other antibiotics.

**Tetracycline**

Tetracycline is known to have a single strong binding site in the entire ribosome, located in the 30S subunit, in addition to several weaker binding sites on both subunits (Epe et al. 1987; Kolesnikov et al. 1996), but only the strong binding site is responsible for its inhibitory action during translation (Buck and Cooperman 1990). We observe two tetracycline sites in the 30S subunit. The strong or primary binding site makes contacts with an irregular minor groove of helix 34 and peripherally with bases 964–967 in helix 31. This position places it in the A site of the 30S subunit, in a position that would sterically clash with A-site tRNA after accommodation. The structure thus explains the observation that tetracycline bind-

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**Figure 8.** Antibiotics bound to the 30S subunit. (a) Overview of the 30S subunit outlining the central region in which the antibiotics we have studied bind. (b) The central region expanded to show the locations of streptomycin (blue), spectinomycin (green), paromomycin (red), tetracycline (orange), pactamycin (cyan), and hygromycin B (magenta).
During initiation, the 30S subunit binds three initiation factors, IF1, IF2, and IF3, along with initiator tRNA. As a first step in understanding the structure of the initiation complex, we have determined the structure of IF1 bound to the 30S subunit. The structure shows that IF1 interacts directly with A1492 and A1493 in the A site, and its binding would sterically prevent the binding of A-site tRNA. Both of these are consistent with earlier biochemical data (Moazed et al. 1995; Dahlquist and Puglisi 2000). The binding of IF1 causes a change in the conformation of helix 44 that is propagated to result in global domain movements, and since IF1 is known to affect the rate but not extent of dissociation of 30S and 50S (Godefroy-Colburn et al. 1975), it is likely that the IF1 complex with the 30S represents a transition state that has a lowered activation energy barrier for subunit association or dissociation.

CONCLUSIONS AND FUTURE PROSPECTS

These are exciting times to be working on ribosome structure. The recent atomic structures of the 30S and 50S, and the subsequent lower resolution but complete molecular model of the 70S with tRNA, have changed the way that ribosome biology will be studied. The structure of the 30S has given us insights into tRNA and mRNA recognition, decoding, and antibiotic function, as well as interactions with initiation factor IF1. The atomic structures of both subunits have been invaluable in the molecular interpretation of the 5.5 Å resolution structure of the 70S ribosome. Nevertheless, a large number of important questions remain, even in the context of the bacterial ribosome, let alone its more complicated eukaryotic counterpart. These range from the sequence of events during initiation and termination to an understanding of how GTP hydrolysis by the various factors is triggered by the ribosome and how this energy is used in various steps such as proofreading or translocation. We also need to learn much more about the molecular details of the conformational changes in the ribosome during translation.

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