Exploration of the conserved A+C wobble pair within the ribosomal peptidyl transferase center using affinity purified mutant ribosomes

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ABSTRACT

Protein synthesis in the ribosome’s large subunit occurs within an active site comprised exclusively of RNA. Mutational studies of rRNA active site residues could provide valuable insight into the mechanism of peptide bond formation, but many of these mutations cause a dominant lethal phenotype, which prevents production of the homogeneous mutant ribosomes needed for analysis. We report a general method to affinity purify in vivo assembled 50S ribosomal subunits containing lethal active site mutations via a U1A protein-binding tag inserted onto the 23S rRNA. The expected pH-dependent formation of the A2450+C2063 wobble pair has made it a potential candidate for the pH-dependent conformational change that occurs within the ribosomal active site. Using this approach, the active site A2450+C2063 pair was mutated to the isosteric, but pH-independent, G2450/U2063 wobble pair, and 50S subunits containing the mutations were affinity purified. The G/U mutation caused the adjacent A2451 to become hyper-reactive to dimethylsulfate (DMS) modification in a pH-independent manner. Furthermore, the G/U mutation decreased both the rate of peptide bond formation and the affinity of the post-translocation complex for puromycin. The reaction rate (k_pep) was reduced ~200-fold for both puromycin and the natural aminoacyl-tRNA A-site substrate. The mutations also substantially altered the pH dependence of the reaction. Mutation of this base pair has significant deleterious effects upon peptidyl transferase activity, but because G/U mutation disrupts several tertiary contacts with the wobble pair, the assignment of A2450 as the active site residue with the neutral pK_a important for the peptidyl transferase reaction cannot be fully supported or excluded based upon these data.

INTRODUCTION

All cells synthesize polypeptides within the ribosome, a large ribonucleoprotein particle. The 50S large ribosomal subunit catalyzes peptide bond formation within the peptidyl transferase center (PTC), an active site comprised exclusively of RNA (1,2). The reaction substrates are an aminoacyl-tRNA and a peptidyl-tRNA, which bind to the A site and P site of the ribosome, respectively (3). The PT reaction involves aminolysis of the ester linkage in the peptidyl-tRNA by the aminoacyl-tRNA α-amino group.

Within the past few years, extensive structural information about the ribosome has been generated, providing a wealth of new insight regarding the chemical details of this biologically ancient enzyme (1,2,4–6). These structural studies confirm years of biochemical data, while demonstrating that the PT reaction is catalyzed within the 23S rRNA. The observation that the ribosome catalyzes peptide bond formation using an RNA active site led to the obvious question: how does the rRNA contribute to peptide bond catalysis? It has already been established that nucleotides within the rRNA are responsible for correctly binding and positioning the 3′-CCA ends of the A-site and P-site tRNA for substrate reactivity (7–11), but is there further catalytic involvement by rRNA nucleotides in the PT reaction?

The pH dependence of the PT reaction potentially provided useful information about ribosomal catalysis. Early kinetic experiments demonstrated that the reaction rate increases with increasing pH, revealing an apparent reaction pK_a of 7.2–7.6 (12–14). This was originally thought to be titration of an active site histidine residue, but the absence of amino acid residues in the active site have forced this interpretation to be reconsidered. Recent measurements involving a rapid factor dependent PT assay demonstrated that there are two reaction pK_a’s, and thus titration of two groups contributes to the
reaction (15). One pKₐ of 6.9 was attributed to deprotonation of the small molecule A-site substrate puromycin (Pmn), whereas the second pKₐ of ~7.5 was attributed to deprotonation of a ribosomal residue.

The chemical identity of the ribosomal group with a 7.5 pKₐ remains unknown, though two possibilities have been suggested previously (15). (i) The titratable group is the N3 position of A2451, which acts as a general base to deprotonate the α-amino group of the A-site tRNA. To serve this role, the pKₐ of A2451 N3 would have to be perturbed by as much as 6 pH units. Although this possibility cannot be ruled out, the charge relay mechanism that was originally proposed to achieve such a perturbation is not consistent with subsequent experimental data (1,16). (ii) The titratable group is the N1 of A2450, which is protonated within the pH dependent and highly conserved non-canonical A2450+2063 wobble pair adjacent to the site of peptide bond formation (15,17). N1 protonation allows it to serve as a hydrogen bond donor to the O2 of cytosine (Figure 1A).

A2450 N1 protonation is consistent with the orientation of A2450 and C2063 in the 50S crystal structure solved at acidic pH (pH 5.8) (2). The pKₐ of the adenosine N1 in a model A+C pair is perturbed toward neutrality (pKₐ = 6.0–6.5) as determined by using NMR (18). This N1 perturbation corresponds to a pKₐ shift of ~2–3 pH units, but the local environment of the PT active site could perturb the pKₐ still higher. In this scenario, the protonated A+C wobble conformation would be the less active form of the ribosome, whereas the deprotonated form would more efficiently catalyze the reaction (15). At pHs below neutrality, A2450 would be protonated and the base pair would be intact; however, at elevated pHs the A2450 would be deprotonated, and the base pair would be destabilized. The A+C wobble could act as a pH-dependent molecular switch between the fully active and the less active forms of PTC.

In order to test the role of the A2450-C2063 wobble pair in the PT reaction, it would be useful to site specifically mutate these nucleotides. The G•U wobble pair is isosteric with an A+ C pair, but pH independent (Figure 1A). However, because the ribosome is essential for cellular viability, mutations in critical active site residues often cause lethal or dominant lethal phenotypes, which make conventional mutagenesis difficult (19).

Several approaches have been developed to biochemically analyze dominant lethal ribosomal mutants. One approach is to express the rRNA mutant in the background of a second mutation that confers antibiotic resistance. The mixed ribosomal population is incubated with the antibiotic, which inactivates the wild-type (wt) ribosomes in the population, but leaves the mutant ribosomes unaffected (20,21). However, antibiotics typically reduce activity only ~100-fold (22), which leaves a relatively small signal-to-noise ratio for mutant analysis. Any mutation that slows reactivity by an amount approaching 100-fold would be uninformative in such an assay. A second approach is to analyze the kinetics of peptide bond formation in a mixed ribosome population, and then deconvolute the rates of the two populations (15). Although this approach

![Figure 1](https://example.com/figure1.png)

**Figure 1.** (A) Schematic representation of an A+C (upper panel) and a G•U wobble pair (lower panel). In this study the A2450+C2063 pair is mutated to a G•U pair. (B) Diagram of the U3X-tag inserted into the 23S rRNA for affinity purification. The three U1A-binding sites are delineated in gray.
was successfully performed on an A2451U active site point mutant, separation of the two rates was extremely laborious. Ribosomal reconstitution using in vitro transcribed 23S rRNA and total 50S ribosomal proteins provides a third means to generate mutant ribosomes (23–25). However, the reconstitution is notoriously inefficient and the resultant ribosomes are significantly less active than the in vivo assembled particles.

Here, we report an alternative approach to generate homogeneous ribosomes with dominant lethal point mutations in the 23S rRNA. In this system mutated rRNA is inducibly expressed and assembled into 50S subunits in vivo, and the mutant ribosomes are purified from wt subunits using a protein affinity tag on the RNA (Figure 1B). The RNA tag is a modified U1A-binding site that is recognized by His6-tagged U1A protein which is co-expressed with the mutant rRNA. Affinity purification produced 50S particles in sufficient yields and purity for use in chemical footprinting and PT reaction analysis. We utilized this purification system to test the effect of the pH-independent A2450G→C2063U mutant wobble base pair on peptide bond formation.

MATERIALS AND METHODS

Expression and purification of His6U1A

His6U1A was expressed from a pET15 vector (Qiagen) and purified on Ni2+–nitrilotriacetate (Ni-NTA) agarose (Qiagen) according to the manufacturer’s protocol. Following column purification, the eluant was dialyzed twice against 1 litre of 20 mM Tris–HCl (pH 7.5), 100 mM NaCl, 1 mM DTT and 20% glycerol. The protein was loaded onto a hydroxyapatite column using an Akta FPLC (Pharmacia), eluted with 0.5 M ammonium sulfate and dialyzed against 1 litre of 20 mM Tris–HCl (pH 7.5), 100 mM KCl, 1 mM DTT, 0.1 mM EDTA and 10% glycerol, and stored at −20°C.

Construction of the U3X-tagged rRNA plasmid

A multi-step procedure was used to insert the U3X-tag at various positions in the rmb operon of plasmid pLK35. A new set of AscI/XhoI/BsiWI restriction sites were inserted at four target sites (see Results) in 23S rRNA using Quick-Change(tm) (Stratagene). None of these restriction enzymes cleaves pLK35. The modified pLK35 plasmids were digested with AscI and BsiWI, gel purified and CIP treated. The annealed and phosphorylated oligonucleotides corresponding to the U3X insert were ligated to the cut plasmid. Transformed colonies were screened for loss of the XhoI site and gain of an EcoRV site. The plasmids were confirmed by sequence analysis (Yale University, Keck facility). The final sequence of the U3X-tag is shown in Figure 1B. A single U1A-binding site (U1X) of sequence CCTCGACCCGAATTCCATGCA-CTCCGAGATTGGTCTCTGAA was inserted using the QuickChange protocol.

Purification of 50S ribosomal subunits

Escherichia coli 50S ribosomal subunits were purified from pop2136 cells as described in (26) with the following modifications. Spin times were adjusted for a Beckman Ti70 rotor. Following the first wash step, the ribosomes were resuspended in 20 mM Tris–HCl (pH 7.6), 0.5 mM NH4Cl, 1.5 mM magnesium acetate, 0.5 mM EDTA and 7 mM 2-mercaptoethanol, and incubated at 4°C for 4 h to dissociate the subunits. The subsequent wash step was done with this low magnesium buffer. Mixed 30S and 50S particles were stored at 4°C until the Ni-NTA purification step. We found that ribosomes purified with the buffer and salt conditions reported here were more active in peptidyl transfer than ribosomes prepared according to (27).

Homogeneous U3X-tagged 50S particles for use in small-scale affinity purifications were isolated from MC250 cells by zonal centrifugation in a Beckman Ti 14 rotor on a 10–40% sucrose gradient (450 ml) in 20 mM Tris–HCl (pH 7.6), 60 mM NH4Cl, 5.2 mM magnesium acetate, 0.25 mM EDTA and 3.0 mM 2-mercaptoethanol (45 000 r.p.m. for 4 h 15 min at 4°C). 50S subunits were stored at −80°C in 70 mM NH4Cl, 30 mM KCl, 7 mM MgCl2, 1 mM DTT, 0.5 mM EDTA and 50 mM Tris–HCl (pH 7.6).

Small-scale affinity purification of U3X-tagged ribosomes

His6U1A protein (500 pmol) was mixed with pre-equilibrated Ni-NTA agarose (from 40 μl of 50% slurry) in 100 μl 1× PD for 30 min at 4°C. The beads were centrifuged gently for 2 min at 4°C, excess U1A was removed, and the beads were washed twice in 1× PD buffer (50 mM Tris–HCl (pH 7.6) 70 mM NH4Cl, 30 mM KCl, 10 mM MgCl2, 5 mM imidazole, 10 μg/ml tRNA and 10 μg/ml heparin). Ribosomes were added (167 pmol) in a 100 μl volume. After 1 h of rocking at 4°C, unbound ribosomes were removed, the beads were washed twice with 1× PD, and the His6U1A-bound ribosomes were eluted in 1× EB (same as 1× PD except it also contained 200 mM imidazole and lacked tRNA and heparin).

Construction of a His6U1A plasmid for co-expression with rRNA

Plasmid PPU used to co-express the His6U1A protein was constructed as follows. The ~3.9 kb pACYC184 (New England Biolabs) BamHI/HindIII fragment, containing the chloramphenicol resistance marker and the ColE1 compatibility group origin of replication, was ligated to the 1.7 kb BamHI/HindIII digestion product of pAS1 (Clontech), which contained the lambda P1 promoter. The His6U1A coding sequence was PCR amplified with primers containing a BamHI site at both ends and cloned into the BamHI site in the pACYC184/pAS1 plasmid. The orientation and sequence of the final product was confirmed by sequence analysis (Yale University, Keck facility). P1 is controlled by the lambda phage cI857 repressor, which is integrated into the genome of pop2136 cells. Incubation at 42°C inactivates the repressor, leading to gene expression. The ColE1 origin is compatible with the pLK35-derived plasmids used to express the rRNA.

Large-scale induction and affinity purification of U3X-tagged ribosomes

rRNA transcription was induced by growing pop2136 cells containing plasmid pLK35-U3X-B at 30°C in ampicillin (Amp) until they reached an OD600 of 0.1. The cells were grown for an additional 2–3 h at 42°C until they reached a final OD600 between 0.8 and 1.0. Cells co-induced for rRNA and His6U1A expression were grown using the same approach
except both Amp and chloramphenicol were included in the media. 50S ribosomal subunits were purified from these cells as described above.

The heterogeneous ribosomal subunits were mixed with Ni-NTA agarose in a 50 ml conical tube. An aliquot of 1 ml of Ni-NTA agarose (50:50 slurry) was added per 40 nmol of crude ribosomes, and 7.5× vol of 1× PD per ml of bead. The mixture was rocked in the cold room for 20 min. The slurry was poured over an empty column, and the flow-through was collected and re-poured over the column. The column was washed in 20× column volumes of 1× PD, and eluted in 25 ml of 1× EB. The eluted ribosomes were pelleted for 3.5 h at 60,000 r.p.m. in a Ti70 ultracentrifuge rotor (Beckman). Purified ribosomes were resuspended in ribosome storage buffer (see above) and stored at −80°C.

**Primer extension assay to measure purification efficiency**

23S rRNA was isolated from ribosomes by phenol/chloroform extraction. A 32P-radiolabeled primer complementary to the 23S rRNA sequence was annealed downstream of the U3X insertion site in 100 mM KCl, 50 mM Tris–HCl, pH 8.5. The primer was extended with AMV Reverse Transcriptase (AMV-RT Roche) for 30 min at 42°C in manufacturer supplied buffer supplemented with 1 mM each dCTP, dATP, dTTP and ddGTP. Products were resolved on a 20% 19:1 acrylamide gel. For the U3X-tagged rRNA, the first incorporated G is four nucleotides from the end of the primer, whereas the first G is seven nucleotides from the primer in the untagged 23S rRNA. Relative band intensities of the primer, whereas the first G is seven nucleotides from the primer in the untagged 23S rRNA. Relative band intensities for the +4 and the +7 products were quantified on a STORM PhosphorImager (Molecular Dynamics) and ImageQuant software. rRNA purity was calculated by comparing the ratio of the +4 band intensity (tagged) to the total intensity of the +4 and +7 bands.

**70S PT kinetic analysis and post-translocation complex formation**

Preparation of the 30S subunits, tRNAs, initiation and elongation factors was performed as described previously (15,16,26). 30S subunits were reactivated by heating for 1 h at 37°C in buffer 50 mM Tris–HCl, 20 mM Bis–Tris (pH 7.2) 30 mM KCl, 70 mM NH4Cl and 20 mM MgCl2. The 70S initiation complex was prepared in buffer A (50 mM Tris–HCl, 20 mM Bis–Tris (pH 7.2) 30 mM KCl, 70 mM NH4Cl and 7 mM MgCl2) by incubating wt 30S ribosomes with 50S subunits (1 μM each), MFT-mRNA (4 μM; coding for fMet-PheThr) and f[3H]Met-tRNAfMet (1.5 μM) in the presence of initiation factors 1, 2 and 3 (1.5 μM each, MFT-mRNA (4 μM; coding for fMet-PheThr) and f[3H]Met-tRNAfMet (1.5 μM) in the presence of initiation factors 1, 2 and 3 (1.5 μM each). FMR-mRNA (4 μM; coding for fMet-PheThr) and f[3H]Met-tRNAfMet (1.5 μM) was added. The rate of fMetPhe formation was measured upon mixing equal volumes (14 μl) of initiation (0.2 μM) and tRNA complexes (1.2 μM) in a quench-flow KinTek apparatus at 37°C. The reaction products released from f[3H]MetPhe-tRNAfMet by alkaline hydrolysis in 0.5 M KOH (for 30 min at 37°C), separated by reverse phase high-performance liquid chromatography (RP-HPLC), and quantified using double-label radioactivity counting.

Post-translocation complexes were prepared by incubating EF-Tu-GTP[14C]Phe-tRNAfMet (1 μM; 1000 d.p.m./pmol) with initiation complexes (1 μM) for 10 min at 20°C. Peptide bond formation resulted in ribosomes carrying deacylated tRNAfMet in the P-site and [3H]MetMet[tRNAfMet in the A-site. Upon incubation (5 min) with EF-G (0.02 μM) and GTP (1 mM), [3H]MetMet[tRNAfMet was translocated to the P-site. Complexes were purified by ultracentrifugation as described above. Quench-flow assays were performed at 37°C by mixing equal volumes of ribosome (0.2 μM) and Pmnn (as indicated) solutions and analyzed as described above.

**A2451 reactivity to dimethylsulfate**

Ribosomes were reacted with dimethylsulfate (DMS) as described in (17) except the reaction was carried out in 50 mM buffer (MES, pH 6.0, HEPEs, pH 6.5–8.5), 150 mM KCl and 15 mM MgCl2. All ribosomes were activated at 42°C for 10 min in reaction buffer prior to incubation with DMS. The primer used for this analysis was 2493 (28).

**RESULTS**

**Design and placement of an RNA affinity tag**

Affinity purification of 50S ribosomes containing 23S rRNA mutations requires that the tag be placed on the rRNA. We selected an affinity tag recognized by the eukaryotic protein splicing factor U1A for the following reasons: (i) U1A is a low molecular weight protein (14 kDa), so its presence should be minimally disruptive to the ribosome provided that the RNA tag is properly positioned; (ii) U1A binds the 11 nt consensus RNA sequence with high specificity and affinity (29,30); (iii) U1A is a eukaryotic protein, so there should be low background binding to E.coli RNAs; and (iv) the recombinant U1A is efficiently expressed in E.coli with good yields and high purity (29,31). Based upon these considerations, we designed a ~100 nt RNA tag (U3X) containing three tandem copies of the U1A consensus-binding site (Figure 1B). The ability of the U3X affinity tag to bind U1A was confirmed by gel-shift analysis with recombinant His6-U1A protein (data not shown).

The site of U3X-tag insertion within the 23S rRNA was selected based upon phylogenetic and in vivo complementation analysis. To function in affinity purification, the RNA tag must be solvent-accessible, it must not disrupt rRNA secondary and tertiary structures, and it must not disrupt major functions of the ribosome, such as subunit association, factor binding or ribosome assembly. Three potential tag insertion sites were selected in helix 9 (termed A), helix 46 (termed B) and helix 54 (termed C) by examining where 23S rRNA sequence expansions had been tolerated during evolution (32). All three sites showed length variation >100 nt. Thus, all of these locations could theoretically accommodate an insertion of the size of ~100 nt U3X-tag. A fourth tag insertion site in helix 98 (termed D) was chosen based upon experimental precedence of extending helix 98 (33). We inserted a single U1A-binding site, U1X, at this position because the
naturally occurring sequence variation ranged only 0–19 nt. The theoretical compatibility of each site was visually confirmed by inspection of the *Haloarcula marismortui* 50S subunit structure (2).

The compatibility of the U3X insertions was tested in an *E. coli* strain (MC250) where the plasmid encoded U3X-tagged rRNA was the only rRNA operon available to the cell (34). The strain lacks all seven chromosomal rRNA operons and thus needs to be supplemented with rRNA from a plasmid. We transformed the MC250 cells with a plasmid containing the rrnB operon (pLK35) with the U3X-tag sequence inserted at positions A, B or C, or the U1X-tag sequence inserted at position D. Cell growth was well supported by rRNA production from the wt plasmid, but the U3X insertion at positions A and C, and U1X insertion at position D substantially interfered with cell growth (data not shown). The effect upon position D insertion was surprising given that this site was tolerant of a tRNA insertion in a previous report (33). Insertion at site B was the only tagged rRNA construct that permitted ribosomal function at levels similar to wt. This construct (pLK35-U3X-B) was used for all subsequent analysis.

**Inducible expression of rRNA mutants in *E. coli***

The goal in developing this system was the expression and purification of dominant lethal 23S rRNA mutants. In addition to the tag, this required an inducible expression system in which mutated 23S rRNA could be produced in the background of chromosomal 23S rRNA transcription. We selected the inducible expression system under the control of a chromosomal encoded temperature-sensitive lambda repressor in *E. coli* strain pop2136 (35). Cells were initially grown at 30°C, where the mutant rRNA was not expressed, and then shifted to 42°C to induce rRNA expression prior to lysing the cells and purifying the 50S ribosomal subunits.

The U3X-tagged rRNA served as a marker of mutant ribosome expression in this mixed population of wt and mutant ribosomes. The U3X-tag sequence on the rRNA was confirmed by primer extension. After 2–3 h of induction at 42°C, ~50% of total cellular rRNA and 20% of the fully assembled 50S subunits contained the U3X-tagged 23S rRNA (data not shown). These values were comparable to previously reported expression levels of rRNA from the Pt promoter (20,34,36). Thus, this brief induction period was sufficient to achieve reasonably high levels of tagged ribosome expression.

**Affinity purification of tagged ribosomes**

Small-scale trial experiments demonstrated that U3X-tagged ribosomes could be retained and eluted on a Ni-NTA column. Ribosomal proteins were retained on the column only when the 23S rRNA was U3X-tagged, as evidenced by SDS–PAGE analysis (Figure 2A, compare lanes 7 and 11). Bound ribosomes were eluted by adding imidazole, which disrupts the His<sub>S</sub>U1A interaction with Ni-NTA.

Although the trial experiments with purified His<sub>S</sub>U1A were successful, generating a sufficient quantity of His<sub>S</sub>U1A protein for large-scale ribosomal affinity purification would immediately become a limiting factor. To circumvent the His<sub>S</sub>U1A purification steps, we designed a two-plasmid system that utilized temperature-sensitive co-expression of both the U3X-tagged rRNA and the His<sub>S</sub>U1A protein. Plasmid PPU was constructed with the His<sub>S</sub>U1A protein expressed under the control of the temperature-sensitive lambda repressor and a ColE1 compatible origin of replication (for details see Materials and methods). Co-expression of rRNA from the pLK35-U3X-B plasmid and His<sub>S</sub>U1A protein from the PPU plasmid did not slow pop2136 cell growth significantly (data not shown). We anticipated that the His<sub>S</sub>U1A would bind the U3X-tag in *vivo*, and remain bound during the subsequent purification. This would obviate the need for independent purification of the U1A protein. In addition, U1A binding *in vivo* could help protect the U3X-tag from ribonuclease degradation damage that may occur during the expression and purification process. Improved longevity of the RNA tag could lead to improved yields of mutant 50S ribosomes.

A large-scale purification of U3X-tagged ribosomes from a mixed population was performed to test the yield and purity of the affinity purification process. Separation of the purified ribosomes by ultracentrifugation in a sucrose gradient showed that the eluted fraction contained exclusively 50S subunits (data not shown). The yield of purified U3X-tagged ribosomes was ~4% of the total ribosomes in the cell at the time of lysis. The eluted fraction was tested for purity by primer extension analysis. Greater than 95% of the 50S ribosomal subunits in the purified fraction contained a U3X-tag (Figure 2B). This is likely to underestimate the purity because the analysis gives an estimate of 3% tagged even when the ribosomes were 100% untagged (Figure 2B, lane 2).

**Characterization of the affinity purified ribosomes**

Several assays were used to determine whether the rate of peptide bond formation was affected by the U3X-tag or the affinity purification process. The first activity assay measured how well affinity purified tagged ribosomes initiated translation. Untagged 50S subunits or purified U3X-tagged 50S subunits were incubated with f<sup>1</sup>Met-tRNA<sup>Fmet</sup>, 30S subunits, mRNA, initiation factors and GTP. Binding of f<sup>1</sup>Met-tRNA<sup>Fmet</sup> to the 70S particle was measured by nitrocellulose filtration. Within the 15 min incubation, 73% of the wt ribosomes and 61% of the tagged ribosomes bound to the fMet-tRNA<sup>Fmet</sup>. This suggests that initiation activity was not significantly affected by the tag or the purification protocol.

The ability of affinity purified ribosomes to form a dipeptide was tested by mixing initiation complexes containing a P-site f<sup>1</sup>Met-tRNA<sup>Fmet</sup> with [<sup>14</sup>C]Phe-tRNA<sup>Phe</sup>, which had been pre-incubated with EF-Tu, GTP, phosphoenolpyruvate and pyruvate kinase (16). HPLC analysis of the product measured the extent of dipeptide formed by unpurified or affinity purified ribosomes in this single-turnover reaction. The untagged ribosomes reacted to an end point of 60%. Thus, the U3X-tag and the affinity purification process had only minimal effects on the extent of dipeptide formation.

Next, we characterized the U3X ribosomes for tripeptide formation. In this reaction the fMet-tRNA<sup>Phe</sup> is translocated to the P-site by EF-G. This post-translocation complex is incubated with Pmn, a small molecule mimic of the
aminoacyl-tRNA. Pmn reaction with the fMetPhe-tRNA results in the tripeptide fMetPhePmn (16). The reaction end point for the untagged ribosomes was 59%, whereas 33% of the U3X-tagged ribosomes reacted. This suggests that the inactive fraction in the tagged ribosomes was slightly larger than the untagged, possibly resulting from the purification process. In both cases, the majority of the reaction was completed by the first time point (10 s).

The rate of the PT reaction was measured under conditions in which the chemical step is expected to be rate limiting (15) (Figure 3A). The untagged ribosomes reacted at a rate of 12 s⁻¹, which was 2-fold faster than the rate of the affinity purified ribosomes (6 s⁻¹) (pH 7.5, 37°C). Thus, there is a slight deleterious effect of tagging and affinity purifying the ribosomes, but it is a modest effect that can be readily controlled for. Thus, in all of the cellular viability and ribosomal reactivity assays, tag insertion and affinity purification has no more than 2-fold effect.

**Mutation and kinetic characterization of the active site A2450G•C2063U wobble pair mutation**

The U1A tag affinity purification system was used to investigate the active site A2450G•C2063U wobble pair mutation in an effort to test if the A2450 N1 is the functional group in the PTC with a pKₐ of 7.5. As discussed above, the A2450 N1 is likely to have a perturbed pKₐ based upon the wobble conformation of the equivalent A2485•C2104 base pair in the *H. marismortui* 50S crystal structure (A2450•C2063 in *H. marismortui*) and the NMR studies show that the adenosine N1 pKₐ is perturbed in A+•C wobble pairs (18). A G•C pair is isosteric with an A+•C wobble pair, but it is pH independent (Figure 1A). Therefore, we set out to test how mutation of the A+•C pair to a G•U would affect the pH dependence of the active site.

The A2450G•C2063U mutation, like many active site mutations, has a dominant lethal phenotype. Ribosomes with the G•U mutation could not support growth when provided as the

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**Figure 2. Affinity purification of U3X-tagged 50S ribosomes.** (A) Selective retention of U3X-tagged 50S ribosomes by His₆U1A protein. Ribosomes or ribosomes containing a U3X-tag were affinity purified by incubation with Ni-NTA and His₆U1A protein. Aliquots of the batch purification process were analyzed by SDS-PAGE, as indicated, and stained with Coomassie Blue. (B) Analysis of ribosome purity by primer extension. Lane 1, no template. Lane 2, untagged rRNA control. Lane 3, homogeneous U3X-tagged rRNA control. Lanes 4–6, affinity purified 23S rRNA from U3X-tagged wt or G•U wobble mutant (GU) ribosomes. Lanes 5 and 6 are from two independent large-scale purifications of GU ribosomes. Lane 7, GU mutant 23S rRNA from ribosomes in the translocation complex.
only source of ribosomes in MC250 cells (data not shown). Furthermore, expression of the mutant interfered with growth when co-expressed with wt ribosomes in pop2136 cells (data not shown). Therefore, in order to biochemically assay ribosomes with this mutation, it was necessary that they should be affinity purified.

The A2450G/C2063U point mutations were introduced into the U3X-tagged rRNA construct pLK35-U3X-B. The resulting plasmid was co-transformed with the U1A protein expression plasmid into pop2136 cells. Tagged mutant ribosomes and the His6U1A protein were co-induced at 42°C and the ribosomes affinity purified to a level >95% pure (Figure 2B, lanes 5 and 6). The yield after purification was 4–7% of all ribosomes for wt, and 4–5% of all ribosomes for G/CU mutants. Mutation of the active site did not cause significant reduction in cell growth during the 2–3 h induction.

The mutant ribosomes were analyzed using the same PT assays outlined above. Initiation was not adversely affected by the mutation. A total of 55% of the mutant ribosomes bound fMet-tRNA compared to 61% for purified wt ribosomes. The end point of dipeptide formation was largely unaffected by the mutant, in that there was 50% product formation after 10 min at 20°C compared to 60% for purified wt ribosomes, therefore the dipeptide formation occurred with near quantitative yields. The fMetPhe-tRNAs were translocated to the P-site, and the amount of U3X-tagged rRNA in the post-translocation complex measured by primer extension. The percentage purity of the rRNA was unchanged (95%; Figure 2B, lane 7) indicating that there was no selection against the G2450/C2063 mutant in the post-translocation complex. This complex was then assayed for peptide bond formation upon Pmn addition. The reaction end points were unaffected by the G/CU mutant (33% for purified G/CU wobble ribosomes and 27% for purified wt ribosomes), but the reaction rates were substantially different. The wt affinity purified ribosomes reacted at a rate of 6 s⁻¹ (10 mM Pmn, 37°C, pH 7.2; Figure 3A).

The observed decrease in the rate of Pmn reaction on the mutant ribosomes may be due to a slower rate of catalysis or to a weaker binding of Pmn. To distinguish between the two possibilities, we measured the dependence of $k_{\text{pep}}$ on Pmn.

Figure 3. Kinetic analysis of G/CU mutant ribosomes. (A) Rate of tripeptide formation in the reaction between fMetPhe-tRNA$^{\text{Phe}}$ and Pmn. The extent of reaction is plotted as a function of time for wt 50S ribosomes prepared using standard protocols from pop2136 (closed circles) or MRE600 (open circles) cells, U3X-tagged wt 50S ribosomes (closed squares) and U3X-tagged G/CU mutant ribosomes (open squares) prepared by affinity purification. The reaction was normalized to the final amplitudes of the extent of tripeptide formation. (B) Concentration dependence of Pmn reaction (37°C, pH 7.2). Further increase in concentration was not feasible due to limited solubility of Pmn. (C) pH dependence of the G2450/C2063 affinity purified mutant ribosomes at 10 mM Pmn (37°C). The reaction rate is plotted versus the pH in the range 6.2–8.3. (D) Rate of dipeptide formation in the reaction between fMet-tRNA$^{\text{Met}}$ and Phe-tRNA$^{\text{Phe}}$. The extent of reaction is plotted as a function of time for wt 50S ribosomes (closed circles) and U3X-tagged G/CU mutant ribosomes (open circles) prepared by affinity purification. The reaction is normalized to the final amplitudes of the extent of dipeptide formation.
concentration (Figure 3B). The affinity of Pmn to the ribosomes was strongly decreased by the G•U wobble mutation to a $K_a$ of $\sim$80 mM compared to 3 mM measured with the wt ribosomes (15). The rate constant of peptide bond formation extrapolated to Pmn saturation was 0.1 s$^{-1}$ (pH 7.2), 180-fold slower than on the wt ribosomes at the same pH.

**pH dependence of G•U mutant ribosomes**

In wt ribosomes, the slope of the reaction rate as a function of pH is 1.5, which has been interpreted to mean that the reaction involves at least two ionizable groups (15). One of these is the $\alpha$-amino group of Pmn, while the second is an ionizable group in the ribosome. The PT reaction rates of the G•U wobble ribosomes were measured as a function of pH and plotted (Figure 3C). The measurements were carried out at limiting concentration of Pmn (10 mM), because saturating concentrations cannot be achieved due to limited solubility. The slope of the pH dependence of rate is 0.95, a value that would be consistent with the removal of an ionizing A+C base pair in the active site; however, there is no inflection in the pH dependence between pH 5 and 8, which is unexpected. Although several explanations are possible, it appears that the $pK_a$ of Pmn is no longer a limiting factor in the assay. Most likely, the loss of the Pmn $pK_a$ appears to result from the inability to saturate the A-site with Pmn, as the pH dependence under the $k_{cat}/K_M$ conditions may reveal ionizing groups involved in Pmn binding, rather than in catalysis.

**Peptide bond formation with the natural A-site substrate**

Given that the mutant ribosomes appeared capable of forming dipeptide but showed a large loss of reactivity with Pmn, we set out to determine if the inactivity resulted from the use of a small molecule tRNA mimic rather than full tRNAs. The rate of dipeptide formation was measured between fMet-tRNA$^{fM}$ and Phe-tRNA$^{Phe}$ for the wt and mutant ribosomes (Figure 3B). The wt ribosomes produced fMetPhe-tRNA$^{Phe}$ at a rate of 7.5 ± 0.5 s$^{-1}$, while the rate of the mutant ribosomes was 0.04 ± 0.003 s$^{-1}$, a 200-fold rate reduction at a saturating concentration of the ternary complex (1.2 μM). Thus, the reduced activity of these ribosomes does not appear to be an artifact of the Pmn assay. The rate of peptide bond formation is substantially reduced even with full tRNA substrates in the A-site and P-site.

**Chemical footprinting of the rRNA active site in G•U wobble mutants**

Given the substantial rate reduction and change in reaction pH dependence, we next asked if the mutant caused a conformational change in the PTC that could be detected by chemical footprinting. We assayed for DMS alkylation at A2451, a residue whose DMS reactivity in *E. coli* has been argued to reflect an inactive PTC conformation (37). There is a striking DMS-dependent reverse transcriptase stop at A2451 in the G•U mutant ribosomes (Figure 4). The intensity of the A2451 reactivity was substantially greater than that has been observed previously in the wt ribosomes (37,38), and the reactivity was pH independent (Figure 4). A2451 alkylation was independent of any of the standard methods used to alter the PTC conformational state (37). The extent of methylation remained constant between 50S subunits tested directly after purification and subunits that had been ‘activated’ by incubating at 42°C for 10 min. Furthermore, there was no change in subunits ‘inactivated’ by dialysis to remove potassium and sodium ions, and subunits that were ‘inactivated’ and subsequently ‘reactivated’ by adding 150 mM KCl and incubating at 42°C. These data suggest that the G•U mutant ribosomes are locked in an inactive and pH-independent conformation.

**DISCUSSION**

Detailed knowledge of how RNA catalyzes peptide bond formation would provide valuable insight into how this ancient enzyme functions. Biochemical analysis of ribosomal mutants is impaired by the fact that many active site mutations have a dominant lethal phenotype. We set out to develop a robust system for producing in vivo assembled mutant ribosomes in order to facilitate biochemical characterization of dominant lethal mutations. The approach involved introducing a U1A affinity tag into the 23S rRNA and co-expressing the mutant with a His6-tagged form of the U1A protein. A His6-tagged U1A protein binds both to Ni-NTA agarose beads and the tagged ribosomes, making it possible to purify
tagged ribosomes away from cellular expressed wt ribosomes. The tag insertion at A1205 and affinity purification were minimally disruptive to the PT reaction. Mutant ribosomes are greater than 95% pure after one affinity-purification step. Ribosomal yields following purification were sufficient for footprinting experiments and kinetic analysis of peptide bond formation, and were successfully used to examine ribosome reaction rates.

In the course of this study two other groups reported an affinity purification approach to isolate large ribosomal subunits with dominant lethal 23S rRNA mutations (39,40). Leonov et al. (39) introduced a streptavidin aptamer sequence into helix 25 and used this tag to affinity purify ribosomes with a G2655C mutation that affects translocation. Youngman et al. (40) inserted an MS2 protein-binding site into helix 98. They used this system to extensively characterize mutations at four active site positions for their effect on the PT and peptide release reaction rates. They did not explore mutations at A2450+C2063, and they did not investigate the pH dependence of the mutant ribosome reactivity.

The PTC undergoes a conformational change with a pK_a of \(~7.5\) and the PT reaction is promoted by deprotonation of an active site group with the same pK_a (15,17). The two most commonly suggested candidates for the titratable site group are the N3 of A2451 and the protonated A2450+C2063 wobble pair (2,15,17,38). A2451 was localized to the reaction center repeatedly by using biochemical footprinting studies (9,41,42). It was reported to have a DMS profile in E.coli that correlates with the pK_a of PT, though subsequent studies showed the apparent pH-dependent reactivity must correspond to a conformational change (17,38). Mutation of A2451 to U altered the reaction pK_a as assayed in a mixed ribosomal population (15), though the possibility that this was an indirect effect resulting from an active site conformational change cannot be excluded. The biochemical data on A2451 leaves open the possibility that a different residue contains the ionizable group in the active site. For this reason, we used the affinity purification method to characterize the A2450G\(\cdot\)C2063U wobble pair mutant.

In the H.marismortui 50S crystal structure determined at pH 5.8, the A2450+C2063 (A2485+C2104 in H.marismortui) pair is in a wobble configuration (2) (Figure 5). Although protons are not visible in this structure, the arrangement of the two nucleotides and the pH of the solution is consistent with A2450 (A2485) N1 protonation. One model we have suggested is that deprotonation of this group at near neutral pH could induce a conformational change in the PTC that promotes reactivity. The G\(\cdot\)U wobble pair is isosteric with the A\(\cdot\)C wobble, but its formation is pH independent in the neutral pH range (Figure 1A). Assuming there are no other changes in the active site resulting from the mutations, the G\(\cdot\)U would stabilize the wobble pair, which would keep the active site in the less active conformation and reduce the pH dependence of the reaction. This hypothesis was explored with affinity purified A2450G\(\cdot\)C2063U mutant 50S ribosomes using both chemical reactivity and enzyme kinetics assays.

The results of the A2451 DMS reactivity experiments on the G\(\cdot\)U mutant ribosomes are somewhat unexpected. In the H.marismortui 50S crystal structure, the N1 of A2451 (A2486) is hydrogen bonded to G2061 (G2102) and is not solvent accessible (Figure 5). 50S E.coli ribosomes in an ‘activated’ conformation (induced by heating to 42°C) are notably unreactive to DMS at A2451 (37). Therefore, DMS reactivity at A2451, which occurs at the N1 position, indicates that the residue is solvent exposed and in a conformation different than that observed in the crystal structure. In the context of the G2450\(\cdot\)U2063 mutation, A2451 becomes substantially more reactive to DMS modification, which is in contrast to the expectation that the G\(\cdot\)U mutation would stabilize the conformation observed in the crystal structure. However, as expected from the experimental design, the reactivity is pH independent. Furthermore, A2451 is insensitive to any efforts.

![Figure 5](image-url) Structure of conserved residues within the 23S rRNA active site from the H.marismortui 50S subunit crystal structure (2). The A2450+C2063 wobble pair (E.coli numbering) is shown in orange, A2451 is shown in blue, G2061 and G2447 are shown in green and C2501 is in yellow. The N1 of A2451 that reacts with DMS is shown as an enlarged sphere. The hydrogen bonds between A2450+C2063 that are likely to be disrupted by mutation to a G\(\cdot\)U pair are shown in red. This includes interactions within the A2450+C2063-C2501 triple, as well as the G2061-A2451-G2447 triple stacked above it.
to ‘activate’ or ‘inactivate’ the ribosomes. Thus, these data suggest that mutation to G•U results in a pH-insensitive active site conformation, but the conformation is not the one observed in the low pH 50S crystal structure.

The affinity purified G•U mutant ribosomes were also analyzed for effects on peptide bond formation. Mutation had no significant effect prior to formation of the post-translocation complex, including initiation and translocation, but the G•U mutation reduced the rate of peptide bond formation with both Pmnm and aminoacyl-tRNA by ~200-fold. The magnitude of the Pnm effect was similar to that of the A2451U mutation, as assayed in a mixed ribosome population (15). Although the pH dependence of the residual reactivity changed substantially, it did not follow a pattern consistent with any previously defined model of the reaction mechanism. The substantial reduction in the rate of peptide bond formation is consistent with the hypothesis that A2451 DMS reactivity is indicative of an inactive conformation.

The contradictory nature of the data suggests that the G2450•U2063 mutation is not isomorphous with the A+G pair within the confines of the tightly folded PTC. The A2450+2063 pair is part of a base triple that includes C2501 (E.coli numbering) (Figure 5). This triple stacks with a second triple (G2447•A2451•G2061) that includes the active site A2451. Although the conformation of the isolated G•U wobble pair is predicted to be isoceric with an A+G pair, this is unlikely to be true of the entire C2501• A2450•2063 base triple. At least two of the three hydrogen bonds to C2501 are lost in the G•U mutation (A2450 N6-C2501 N3 and C2063 N4-C2501 O2). For both hydrogen bonds the G•U mutation would cause lone pairs of electrons to be juxtaposed. Unfortunately, there is no combination of the naturally occurring nucleosides that would produce a pH-independent base triple that is isoceric with C2501• A2450•2063. C2501 also makes a tertiary contact to the 2′-OH of G2061 (Figure 5), a hydrogen bond that appears to connect the two base triples. It is possible that displacement of U2063 away from C2501 would in turn displace G2061 to connect the two base triples. It is possible that displacement of U2063 away from C2501 would in turn displace G2061 away from A2451. This would explain the hypersensitivity of A2451 to DMS modification. In addition to these effects, the G•U mutation introduces an amino group into the active site at position 2450, which may explain why the effect on peptide bond formation is so large. As a result of these considerations, the assignment of A2450 as the active site residue with the neutral pKₐ important for the PT reaction cannot be supported or completely excluded based upon these data on the G2450•U2063 mutation. However, it is clear that mutation of this base pair has a significant deleterious effect upon PT activity.

We have successfully developed a gentle non-disruptive affinity purification method for isolation of in vivo assembled lethal mutant 50S ribosomes. The relative simplicity of the co-expression and purification method will make it possible to biochemically explore other mechanistic hypotheses regarding the role of rRNA in peptide bond synthesis.

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