Single methylation of 23S rRNA triggers late steps of 50S ribosomal subunit assembly

Taiga Arai, Kensuke Ishiguro, Satoshi Kimura, Yuriko Sakaguchi, Takeo Suzuki, and Tsutomu Suzuki

Department of Chemistry and Biotechnology, Graduate School of Engineering, University of Tokyo, Tokyo 113-8656, Japan

Edited by Harry F. Noller, University of California, Santa Cruz, CA, and approved July 20, 2015 (received for review April 6, 2015)

Ribosome biogenesis requires multiple assembly factors. In Escherichia coli, deletion of RlmE, the methyltransferase responsible for the 2′-O-methyluridinyl modification at position 2552 (Um2552) in helix 92 of the 23S rRNA, results in slow growth and accumulation of the 45S particle. We demonstrate that the 45S particle that accumulates in ΔrlmE is a genuine precursor that can be assembled into the 50S subunit. Indeed, 50S formation from the 45S precursor could be promoted by RlmE-mediated Um2552 formation in vitro. Ribosomal protein L36 (encoded by rplJ) was completely absent from the 45S precursor in ΔrlmE, and we observed a strong genetic interaction between rlmE and rplJ. Structural probing of 23S rRNA and high-salt stripping of 45S components revealed that RlmE-mediated methylation promotes interdomain interactions via the association between helices 92 and 71, stabilized by the single 2′-O-methylation of Um2552, in concert with the incorporation of L36, triggering late steps of 50S subunit assembly.

Ribosome assembly | post-transcriptional modification | rRNA methyltransferase | RlmE | L36

Ribosomes are essential ribonucleoprotein complexes that translate the genetic information encoded by mRNA into protein. Intact bacterial ribosomes, which have a sedimentation coefficient of 70S, consist of large (50S) and small (30S) subunits. Each subunit can be reconstituted in vitro from its individual component ribosomal RNAs and proteins (1–5), indicating that these components contain all of the information necessary to automatically assemble into functional subunits. Because the intermediate particles observed in an in vitro reconstitution of the subunits are similar to those observed in vivo, assembly maps that describe the order of ribosomal protein binding in vitro are useful tools for understanding ribosome biogenesis in the cell (6). However, assembly is slower in vitro than in vivo, and nonphysiological conditions, such as high-salt concentration and high temperature, are required for in vitro assembly.

In the cell, ribosome assembly is coordinated with the transcription and processing of large precursor rRNAs encoded by the rrn operon (3, 7). Once transcription starts, nascent transcripts rapidly form local secondary and tertiary structures that serve as binding sites for the primary ribosomal proteins, thereby initiating ribosome assembly. However, hierarchical assembly starting at the 5′-terminal domains of the 16S and 23S rRNAs is not essential for ribosome formation in the cell (8) because nonribosomal factors, termed “assembly factors,” facilitate rapid and efficient assembly of ribosomal particles. Ribosome biogenesis is an energy-consuming process in which a number of assembly factors, including RNA helicases, GTPases, protein chaperones, and rRNA-modifying enzymes, support efficient assembly of each subunit (6, 9, 10). Individual mutation or deletion of several assembly factors causes accumulation of immature 50S subunits that sediment at 40S or 45S. For example, the RNA helicase SrmB participates in the early stage of 50S assembly (11, 12); deletion of smrB results in accumulation of a 40S particle lacking L13, one of the primary binders that initiates 50S assembly in vitro (13). Deletion of the RNA helicase DeaD also results in accumulation of a 40S particle; however, none of the primary binders are missing from this particle, indicating that DeaD acts later than SrmB during 50S assembly (9). The 40S particle that accumulates in ΔdeaD is a genuine precursor of 50S, rather than a dead-end particle (14). DbpA, another RNA helicase, is also involved in 50S assembly. Overexpression of the R331A active-site mutant of DbpA in Escherichia coli results in accumulation of a particle that sediments at 45S at low Mg2⁺ concentration, but increases in density and comigrates with the 50S subunit at high Mg2⁺ concentration (15), suggesting that the 45S particle is structurally flexible and can undergo conformational changes upon binding of Mg2⁺. A similar 45S particle that undergoes Mg2⁺-dependent alteration in density appears in cells expressing the G80E/D85N mutant of ObgE (CgtAΔ) (16), an essential GTPase involved in the late steps of 50S assembly (10).

Some rRNA-modifying enzymes play critical roles in ribosome assembly. RlmE (RrmJ, FtsJ) is an S-adenosylmethionine (AdoMet)-dependent methyltransferase that catalyzes 2′-O-methylation of uridine at position 2552 (Um2552) in the A-loop (Helix 92) of 23S rRNA (Fig. 1 A and B) (17). RlmE is highly conserved in all domains of life, as well as in organelles, indicating the functional importance of this enzyme. Um2552 is situated at 5′ adjacent to G2553 (Fig. 1 B), which is an essential base that anchors the CCA terminus of A-site tRNA in the 50S subunit (18). The absence of a 2′-O-methyl group at Um2552 in the rlmE deletion strain of E. coli (ΔrlmE) decreases the rate of programmed frameshifting and read-through of stop codons (19), indicating that Um2552 is required for optimization of translational accuracy and thus allows natural recoding events. Deletion of rlmE resulted in a growth defect and accumulation of a 45S particle, especially at low temperature (Fig. 1 C) (20). The particle was similar to that observed in E. coli cells

Significance

Ribosome biogenesis requires a number of assembly factors. However, the exact mechanism of this process remains largely unknown. We could successfully reconstitute the 50S subunit from the 45S precursor by a 2′-O-methylation of U2552 (Um2552) mediated by rRNA methyltransferase RlmE, in the presence of the wash fraction from crude ribosomes. To our knowledge, this experiment is the first demonstration of enzymatic formation of a ribosomal subunit in vitro from its precursor via the action of an assembly factor. Mechanistic studies revealed that RlmE-mediated Um2552 formation promotes interdomain interactions of 23S rRNA, in concert with the incorporation of L36, triggering late steps of 50S subunit assembly. RlmE and Um2552 are conserved in other organisms, including human, indicating the functional importance of this process in ribosome biogenesis.

Author contributions: Tsutomu Suzuki designed research; T.A., K.I., S.K., Y.S., and Takeo Suzuki performed research; Y.S. contributed new reagents/analytic tools; T.A., K.I., and Tsutomu Suzuki analyzed data; and T.A. and Tsutomu Suzuki wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

Author contributions: Tsutomu Suzuki designed research; T.A., K.I., S.K., Y.S., and Takeo Suzuki performed research; Y.S. contributed new reagents/analytic tools; T.A., K.I., and Tsutomu Suzuki analyzed data; and T.A. and Tsutomu Suzuki wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

1To whom correspondence should be addressed. Email: ts@chembio.t.u-tokyo.ac.jp.

This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1506749112/-/DCSupplemental.
overexpressing mutants of DbpA (Fig. 1C) (15) or ObgE (16). It remains unknown, however, whether this 45S particle is able to form an active 50S subunit (21). The cold-sensitive phenotype, along with accumulation of the 45S particle, was alleviated by overexpression of the obgE or engA GTPase (22), indicating that RlmE is an assembly factor that acts at a late step of 50S formation. However, the mechanism by which RlmE-mediated Um2552 formation facilitates late 50S assembly remains unknown.

In this study, we demonstrate that the 45S particle that accumulates in ΔrlmE is a genuine precursor of the 50S subunit. Furthermore, we partially recapitulate the formation of 50S from 45S in the presence of recombinant RlmE and AdoMet in vitro. Genetic and biochemical studies suggested that the Um2552 formation by RlmE facilitates interdomain interactions between H92 and H71, in concert with the incorporation of L36 and other ribosomal proteins, leading to correct placement of the central protuberance and other domains during late steps of 50S assembly.

Results

The 45S Particle Is an Assembly Intermediate of the 50S Subunit. In sucrose density gradient (SDG) centrifugation analysis (Fig. 1C), the 45S particle in ΔrlmE cultured at low temperature (22 °C) sedimented between 30S and 50S at a low Mg2+ concentration (0.5 mM), as previously reported (20), but increased its sedimentation coefficient to comigrate with the 50S subunit at a high Mg2+ concentration (10 mM); thus, the 45S particle is unable to associate with the 30S subunit to form the 70S ribosome, even at 10 mM Mg2+. As noted above, a similar particle that also exhibits Mg2+-dependent alteration of density appears in cells overexpressing the R331A mutant of DbpA (Fig. 1C) (15). To characterize the physical properties of the 45S particle of ΔrlmE, we used SDG centrifugation to analyze the 45S particle together with the 50S subunit at various Mg2+ concentrations (Fig. 1D).

The particle gradually increased its density in response to the increasing Mg2+ concentration and almost overlapped with the 50S subunit at 2.0 mM Mg2+, indicating that the 45S particle is structurally flexible and can alter its conformation consecutively in response to the Mg2+ concentration. The result clearly ruled out the possibility that the precursor has two distinct structural conformers bearing 45S and 50S.

The 45S particle of the ΔrlmE strain has been speculated to be a dead end or breakdown product, rather than a bona fide precursor, of the 50S subunit (21). To investigate whether the 45S particle in ΔrlmE is an authentic precursor of the 50S subunit, we cultured the E. coli ΔrlmE strain in the presence of [13C] adenosine (Ado), to label rRNAs with 13C-Ado, and then cultured the cells for several hours in LB medium with nonlabeled Ado. We then used mass spectrometry to monitor the 13C-to-12C ratio of Ado in rRNAs from 30S, 45S, and 50S (Figs. S1 and S2). As shown in Fig. 1E, the proportion of 13C-labeled 16S rRNA in 30S decreased moderately over time at a constant speed in normal medium because newly synthesized 16S RNA composed of 12C-Ado constantly replaced 13C-labeled 16S RNA. On the other hand, 13C-labeled rRNAs in 50S clearly increased during the first 40 min after cultivation in normal medium, followed by a moderate decrease, as newly synthesized RNA replaced 13C-labeled RNA. Consequently, the proportion of 13C-labeled rRNAs in 45S plunged during the first 80 min. If the 45S particle is not a precursor for 50S, 13C-RNA in 50S should not increase in normal medium in this experiment. Moreover, we monitored ribosome profiles in ΔrlmE over a time course after blocking transcription with rifampicin (Fig. 1F). In SDG analysis performed at 0.5 mM Mg2+, the level of 45S particle gradually decreased, and about half of the particle was converted to 50S within 120 min. In SDG analysis performed at 10 mM Mg2+, the 70S peaks increased over time whereas the 50S peak decreased proportionally (Fig. S3). Taken together, these results clearly demonstrate that the 45S particle is not a dead-end product; instead, at least half of this 45S particle is a genuine precursor that can be assembled into a 50S subunit, which, in turn, can productively associate with the 30S subunit to form the 70S ribosome.

The RlmH methyltransferase, which is responsible for 3-methylpyrrolidinyluridine (m3Ψ) formation at position 1915 (m3Ψ) in 23S rRNA, employs 70S ribosome as a substrate (23). Therefore, if the...
45S particle of $\Delta rlmE$ were a breakdown product of mature 50S subunits, $m^{\Psi}1915$ would be present in 23S rRNA of the 45S particle. We analyzed the posttranscriptional modification of 23S rRNAs obtained from the 45S particle and 50S subunit of $\Delta rlmE$ cells. The extracted 23S rRNA was digested by RNase $T_1$ and analyzed by capillary liquid chromatography (LC)/nanoelectrospray ionization mass spectrometry (MS). In 50S subunits from both the WT and $\Delta rlmE$, only the $m^{\Psi}1915$-containing fragment (5AAC m$^{\Psi}$AP4AACGp) was detected, and very little unmethylated fragment was observed (Fig. 1G). By contrast, in the 45S particle of $\Delta rlmE$, a large fraction of the corresponding fragment was unmethylated (Fig. 1G). Because $m^{\Psi}1915$ is the sole methylation site in this fragment, we can conclude that position 1915 is unmethylated in the 45S particle. It is reasonable to attribute the small fraction of methylated fragments to contaminating 50S subunits. Thus, the 45S particle is a genuine precursor, rather than a breakdown product, of the mature 50S subunit. Therefore, we will refer to this particle as the “45S precursor.”

Enzymatic Formation of the 50S Subunit from the 45S Precursor by RlmE-Mediated Um2552 Methylation. To investigate whether RlmE-mediated 2'-O-methylation of U2552 facilitates 50S assembly, we tried to generate the 50S subunit from the 45S precursor in vitro in the presence of recombinant RlmE and AdoMet. Initially, we used crude ribosomes obtained from the 45S precursor in vitro in the presence of recombinant RlmE and AdoMet. In total, about 20% of the 45S precursor was converted to 50S subunits. Under these conditions, we also observed an increase in the 70S peak relative to the 50S and 30S peaks in SDG analysis at 10 mM Mg$^{2+}$ (Fig. S4A). Thus, we proved that the newly synthesized 50S could productively associate with the 30S subunit to form the 70S ribosome. We next determined the 2'-O-methylation status of U2552 in 23S rRNA from both 45S and 50S particles after reconstitution. LC/MS analysis detected the RNA trinucleotide C2551–G2552 in 23S rRNA from 45S and 50S particles (Fig. S4B). Without incubation with RlmE and AdoMet, only unmethylated trimer (CUGp) was detected in both 45S and 50S. After incubation with RlmE and AdoMet, however, the methylated trimer (CUmGp) was clearly detected along with the unmethylated trimer (CUGp) (Fig. S4B). Approximately 50% of 23S rRNA in the 45S was methylated whereas 70% of 23S rRNA from the 50S was methylated.

Furthermore, we attempted to enzymatically reconstitute the 50S subunit in vitro from the isolated 45S particle. To this end, the isolated 45S particle was incubated with RlmE and AdoMet in the presence or absence of the wash fraction from the crude ribosomes. We could detect small amounts of 50S, which were increased by RlmE-mediated Um2552 formation in the presence of the wash fraction (Fig. 2B and Fig. S4C). About 25% of the 45S precursor was converted to 50S subunits. Moreover, the reconstituted 50S was able to associate with the 30S subunit to form the 70S ribosome (Fig. 2C). LC/MS analysis of the RNA trinucleotide C2551–G2552 in 23S rRNA clearly showed that Um2552 was concentrated in the reconstituted 50S relative to the 45S particle (Fig. 2D). Um2552 was found in ~50% and 20% of 23S rRNAs in the 50S and 45S particles, respectively. These data clearly rule out the possibility that RlmE methylates the 50S subunit to facilitate 70S formation. Taken together, these observations demonstrate that Um2552 is indeed formed in the 45S precursor and that methylated 45S is subsequently converted into the 50S subunit.

The U2552C Mutation Causes Accumulation of the 45S Precursor. Next, we sought to clarify whether RlmE, acting as an assembly factor, transiently interacts with the 45S precursor to promote 50S assembly, or whether instead the Um2552 modification is itself required for 50S assembly. To address this issue, we generated the U2552C mutant of 23S rRNA, which is not predicted to be a substrate of RlmE. In the crystal structure of the 50S subunit, Um2552 forms a base-triple with C2556 and U1955, which is inverted in the 45S subunit (Um2552, the U-to-C replacement at position 2552 does not affect this interaction because a cytosine base also has a C2 carbonyl group. In addition, the U2552C mutation does not affect peptidyl-transferase activity in vitro (18). To express the mutant rRNA, a plasmid harboring the mB operon with the U2552C mutation was introduced into an E. coli Δ7 prrn strain (8, 27) in which all seven chromosomal m operons were disrupted; consequently, all ribosomes in this strain were replaced with the U2552C mutant ribosomes. As in $\Delta rlmE$, the U2552C mutant exhibited a cold-sensitive phenotype, but much slower growth even at 30 °C (Fig. 3A). The
ribosome profile revealed that the 45S precursor accumulated in the U2552C mutant (Fig. 3B), and we confirmed that the accumulated 45S precursor increased in density and overlapped with the 50S subunit in SDG analysis performed in 10 mM Mg$_{2+}$ (Fig. S5). The level of 45S accumulation in the U2552C strain is not so different from that in ∆rlmE (Fig. 3B); nevertheless, the U2552C strain grows much slower than ∆rlmE (Fig. 3F). This observation might be explained by low translational fidelity caused by the U2552C mutation that prevents 2'-O-methylation (19), in addition to the retarded 50S assembly by this mutation. We also made a double-mutant strain, ∆rlmE/U2552C, by deleting rlmE in the U2552C strain; the resultant strain was also cold-sensitive (Fig. 3A) and accumulated the 45S precursor (Fig. 3B). However, for unknown reasons, this strain grew better than the U2552C mutant (Fig. 3A). It is possible that the U2552C mutation modulates the configuration of RlmE on the ribosome, causing some adverse impact on ribosome biogenesis.

We next analyzed 2'-O-methylation at position 2552 in the U2552C mutant. In the WT strain (KT103) (8), ∼75% of 23S rRNA was 2'-O-methylated at position 2552 (Fig. 3C). No methylation was observed in ∆rlmE or ∆rlmE/U2552C, which were used as negative controls. In the U2552C mutant, only 15% of 23S rRNA was 2'-O-methylated (Fig. 3C). Because RlmE had a weak ability to methylate C2552, the U2552C mutation efficiently abolished 2'-O-methylation at position 2552 without disruption of rlmE. It remained possible, however, that the U2552C mutation hindered RlmE binding to H92 in the 23S rRNA. Because RlmE can bind to the 50S subunit (20), we measured the relative abundance of RlmE attached to the mutant and WT 50S subunits by stable isotope labeling by amino acids in cell culture (SILAC) (28). The tryptic digests were analyzed by LC/MS to detect specific peptides containing C-terminal Lys residues from RlmE and L3 (Table S1). The unlabeled peptide from mutant 50S and the fully labeled corresponding peptide from WT 50S (with 8-Da higher mass due to the presence of [15N] Lys) were eluted simultaneously from LC and detected in the same mass spectra (Fig. 3D). The intensity ratio of these peptides was normalized against that of L3-derived peptides. The results revealed that the same amounts of RlmE were attached to the mutant and WT 50S subunits by stable isotope labeling by amino acids in cell culture (SILAC) (28). The tryptic digests were analyzed by LC/MS to detect specific peptides containing C-terminal Lys residues from RlmE and L3 (Table S1). The unlabeled peptide from mutant 50S and the fully labeled corresponding peptide from WT 50S (with 8-Da higher mass due to the presence of [15N] Lys) were eluted simultaneously from LC and detected in the same mass spectra (Fig. 3D). The intensity ratio of these peptides was normalized against that of L3-derived peptides. The results revealed that the same amounts of RlmE were attached to the mutant and WT 50S subunits (Fig. 3E); thus, accumulation of 45S precursor in the U2552C mutant is not due to the loss of ability to bind RlmE, but instead to the lack of 2'-O-methylation at position 2552, indicating that the Um2552 modification is directly involved in 50S assembly.

**RlmE Contributes to 50S Assembly in Concert with the Incorporation of L36.** We next analyzed the protein components of the 45S precursor that accumulated in ∆rlmE cells. Comparison of the components to those in the 50S subunit revealed several additional bands in the 45S precursor, which we analyzed by peptide mass fingerprinting (Fig. 4A). The additional proteins were factors responsible for 50S assembly: three RNA helicases, Dcl, SmrB, and RhfE; three rRNA-modifying enzymes, RlmKL, RluB, and RluC; and one assembly factor, YhbY. This result also supports the idea that the 45S precursor of ∆rlmE is an authentic assembly intermediate of the 50S subunit. Possible roles of these assembly factors are discussed in SI Discussion.

Previous work demonstrated that the 45S particle of ∆rlmE lacks some ribosomal proteins, including L5, L16, L18, L25, L27, L28, L30, L33, and L35 (15, 29). We used the SILAC method to measure the abundance of ribosomal proteins in the 45S precursor relative to their abundance in the 50S subunit. Because the stoichiometries of L1, L2, L3, and L4 were unchanged (15, 29), we focused on ribosomal proteins smaller than L5. We subjected 45S and 50S to tryptic digestion and analyzed the digests by LC/MS to detect and quantitate specific peptides (Table S1) from 23 species of ribosomal proteins. We confirmed that L5 was significantly less abundant in the 45S precursor (Fig. 4B), as reported (15). In addition, the levels of L5, L6, L16, L18, L19, L25, and L27 were reduced in the 45S precursor (Fig. 4B). Although this result was basically consistent with previous reports (15, 29), we did not observe any reduction in L28. Because L28 was easily stripped off under the high-salt condition (Fig. 4B), indicating weak association of this protein, this discrepancy might be explained by differences in sample preparation. In addition, we analyzed small proteins by urea/Tris-tricine PAGE (Fig. 4C). L36, which is the smallest protein in the 50S subunit and can barely be detected by MS, was not detectable in the 45S precursor. As a
control, we analyzed the 50S subunit from ΔrpmJ and confirmed that L36 was not detectable (Fig. 4C). No L36 was present in the 45S precursors accumulated in the U2552C mutant or in a strain overexpressing the DbpA R331A mutant (Fig. 4C), indicating that 45S precursors from these mutants shared similar characteristics.

Because both L35 and L36 were absent from the 45S precursor in ΔrlmE, we investigated whether the 45S precursor also accumulated in knockout strains lacking rpmJ (L35) or rpmJ (L36). In ΔrpmJ, we observed a shoulder on the left side of the 50S peak at low Mg2+ concentration (Fig. 4D) that disappeared at high Mg2+ concentration (Fig. 4D), indicating marginal accumulation of the 45S precursor in ΔrpmJ. Deletion of rpmJ causes accumulation of the 50S particle that is unable to productively associate with the 50S subunit (30), leading us to hypothesize that the 45S precursor would also accumulate in ΔrpmJ. Consistent with this prediction, in ΔrpmJ, we detected a 45S precursor exhibiting Mg2+-dependent alteration of its density (Fig. 4D). The level of 45S accumulation in ΔrpmJ, however, was lower than that in ΔrlmE (Fig. 1C), mirroring the more robust growth phenotype of ΔrpmJ relative to ΔrlmE (Fig. 4E). Because rpmJ is the last gene in the spc operon, we suspected that deletion of rpmJ might affect the expression of secY and other genes. To explore this possibility, we complemented ΔrpmJ with plasmid-encoded rpmJ (Fig. S6), demonstrating that 45S accumulation in ΔrpmJ is not due to a polar effect caused by rpmJ deletion; rather, L36 is required for a late assembly of the 50S subunit.

To further investigate the genetic interaction between L36 and RlmE, we generated a double deletion lacking both rpmJ and rpmI (ΔrlmE/ΔrpmJ). This strain exhibited a severe synthetic growth defect at 37 °C and was completely unable to grow at low temperature (Fig. 4E). Furthermore, strong accumulation of 45S precursor was also observed in the double-deletion strain (Fig. S7), indicating that incorporation of L36 acts cooperatively with modification of Um2552 by RlmE to promote late steps of 50S assembly.

We next analyzed the status of the Um2552 modification in the 45S precursor of ΔrpmJ. In the 50S subunit from WT cells, 85% of 23S rRNA was 2′-O-methylated (Fig. 4F). In ΔrpmJ, 40% of 23S rRNA was 2′-O-methylated in the 45S precursor whereas 85% was 2′-O-methylated in the 50S subunit (Fig. 4F). These data suggest that L36 incorporation is not necessary for the formation of Um2552 mediated by RlmE.

Stripping Ribosomal Components from the 45S Precursor. The Mg2+-dependent alteration of its sedimentation coefficient suggests that the 45S precursor has a flexible structure in which intramolecular interactions have not yet been fully established and stabilized. We investigated this physical property by stripping ribosomal components using high concentrations of salt. Specifically, we incubated the 45S precursor from ΔrlmE and the WT 50S subunit in a solution containing 500 mM ammonium chloride in the presence of 0.5 mM or 10 mM Mg2+, and then subjected the samples to SDG to separate components [upper fraction (U) in Fig. 4G] that dissociated from the particles [lower fraction (L) in Fig. 4G]. From each fraction, rRNAs and protein components were prepared separately and then analyzed by denaturing PAGE for rRNAs (Fig. 4G, Upper) and SDS/PAGE for proteins (Fig. 4G, Lower). No components dissociated from the WT 50S under these conditions. By contrast, 5S rRNA and several protein components dissociated from the 45S precursor, especially at low Mg2+ concentration. By contrast, at high Mg2+ concentration, dissociation of these components was clearly suppressed, indicating that binding of numerous Mg2+ ions stiffens the internal structure of the 45S precursor, thereby preventing dissociation of its components. Mass-spectrometric analysis revealed the dissociated proteins in the upper fraction to be L5, L6, L9, L14, L16, L25, L27, and L28. To determine the levels of these dissociated proteins, we treated the particle with high-salt solution at low Mg2+ concentration, isolated it from the lower fraction of...
the SDG, and then quantitated its residual components by SILAC (Fig. 4B). We confirmed that the levels of the dissociated proteins were clearly reduced in the particle after high-salt treatment. Because L5, L18, and L25 associate with 5S rRNA, we assume that these proteins dissociate from the 45S precursor along with 5S rRNA as a result of high-salt treatment. L6, L14, and L16 are in close proximity to the region surrounding H92 and the L36-binding site, implying that the structure of this region is immature.

We also carried out the same experiments on 45S precursors isolated from ΔrpmE (Fig. S8) or from a strain that overexpressed the DbpA R331A mutant (Fig. S8); in both cases, the results were similar to those obtained with the 45S precursor from ΔlmE.

**Structural Probing of 23S rRNA in the 45S Precursor Using Hydroxyl Radicals.** To investigate the structural differences in 23S rRNA between the 45S precursor and the 50S subunit, we performed hydroxyl-radical probing to analyze the solvent accessibility of the rRNA sugar-phosphate backbone (31). The probing was conducted at both low and high Mg2+ to identify the regions sensitive to Mg2+ concentration. Cleaved positions were determined by primer extension covering the entire sequence of the 23S rRNA (Fig. 5 A and B and Fig. S9). Compared with the 23S rRNA cleavage pattern in the 50S subunit, we found that a number of sites specifically cleaved in the 23S rRNA of the 45S precursor, which could be categorized into two types of regions. Regions of the first type were exposed to solvent specifically in the 45S precursor, regardless of Mg2+ concentration (Fig. 5A), and we designated these sites as “45S-specific cleavage” (color-coded in blue in the figures). Regions of the second type were sensitive to hydroxyl-radical cleavage only at low Mg2+ concentration (Fig. 5B), and we designated these sites as “low Mg2+ cleavage” (color-coded in red in the figures). We mapped both types of regions onto the crystal structure of the E. coli 50S subunit (Fig. 5C).

In the 45S precursor, the loop sequence of H92, including positions 2552, 2554, and 2555, was more sensitive to hydroxyl radicals, regardless of the Mg2+ concentration (Fig. 5A and D). In the crystal structure, H92 interacts with H71 in domain IV via a base-triple composed of Um2552, C2556, and U1955 (Figs. 5D and 6A). We also observed 45S-specific cleavages at positions 1950–1956 in H71 (Fig. S9G). These results strongly suggest that 2′-O-methylation of Um2552 plays an important role in stabilizing the Um2552–C2556–U1955 base-triple, thereby facilitating the interdomain interaction between H92 and H71 (Figs. 1B, 5D, and 6A); therefore, the absence of 2′-O-methylation at position 2552 destabilizes this region. The weak binding of L14 in the 45S precursor (Fig. 4B) could be explained by instability of the interaction between H92 and H71 (Fig. 5D). Furthermore, we observed 45S-specific cleavages in H89, H91, and H97 (Fig. 5A, C, and D and Fig. S9L), which, together with H42, form a binding pocket for L36 (Fig. 5D). The complete absence of L36 in the 45S precursor (Fig. 4C) is consistent with this observation. We also detected higher levels of 45S-specific cleavage in H38 (Figs. S9C and D and S104), which serves as a binding site for L16 and 5S rRNA. This observation explains why 5S rRNA, along with three bound proteins, dissociated from the 45S precursor after high-salt treatment (Fig. 4B and G) and also explains the low stoichiometry and weak association of L16 in the 45S precursor (Fig. 4B). Likewise, 45S-specific cleavages were present in helices 11, 13, and 31 (Figs. S9A and B and S10B), which forms a binding site for L35, explaining the low level of L35 in the 45S precursor (Fig. 4B). The low Mg2+-specific cleavages were apparently concentrated in domain III, including helices 52 and 55–58 (Fig. 5 B and C and Fig. S9 E and F), indicating that Mg2+-dependent alteration of the sedimentation coefficient of the 45S precursor originates from a dynamic conformational change in domain III that is mediated by Mg2+ binding. H52 in domain III and H66 in domain IV engaged in an interdomain interaction (Fig. 5E), and their contact site contained low Mg2+-specific cleavages (Fig. 5E and Fig. S9 E and H), indicating that the H52–H66 interaction in the 45S precursor is stabilized at high Mg2+ concentration.

**Discussion**

RlmE preferentially methylates the 50S subunit rather than the 45S particle that accumulates in ΔlmE (20). On the basis of this observation, the 45S particle has been regarded as a dead-end or


breakdown product of the 50S subunit (21). However, in this study, we demonstrated that the 45S particle could actually be converted to the 50S subunit in ΔrlmE (Fig. 1 E and F), demonstrating that 45S is a genuine precursor of 50S. Furthermore, we could partially reconstitute the 50S subunit from the 45S precursor in vitro by RlmE-mediated formation of Um2552 (Fig. 2B). This experiment is the first demonstration of the in vitro formation of 50S from its precursor, mediated by the enzymatic activity of an assembly factor. Given that most 50S subunits in cells are associated with 30S subunits at sufficient concentrations of Mg\(^2+\), it is reasonable to speculate that the physiological substrate for RlmE is not the 50S subunit, but rather the 45S precursor, because Um2552 in H92 of 23S rRNA is located on the interface side of the 50S subunit (24–26). Quantitative analyses of rRNA modifications by HPLC/MS have shown that the level of Um2552 surges during late steps of 50S assembly (32, 33), and these results are consistent with our observations.

To obtain insights into the function of an RNA modification responsible for a specific biological process, it is necessary to determine whether the RNA-modifying enzyme or the RNA modification itself plays a pivotal role in the process. In this study, we resolved this issue by examining the U2552C mutant. This mutation reduced 2′-O-methylation at position 2552 (Fig. 3C) and accumulated the 45S precursor (Fig. 3B) without hindering RlmE binding to the 50S subunit (Fig. 3 D and E). These data strongly suggest that RlmE binding to the 45S precursor does not contribute to 50S assembly; instead, Um2552 modification itself plays a critical role in triggering late steps of 50S assembly. In general, a 2′-O-methylated residue favors the C3′ endo ribose pucker conformation (34). The solution structure of H92, determined by NMR, revealed that 2′-O-methylation of Um2552 restrains the conformational dynamics of the A-loop structure by influencing Um2552–C2556 pairing (35) (Fig. 1B). In the crystal structure of the 50S subunit, Um2552 forms a base-triple along with C2556 and U1955 in H71 (Fig. 6A). If Um2552 were to adopt the C2′ endo ribose pucker, the base-triple would not be stabilized. Therefore, it is likely that the Um2552 modification stabilizes the base-triple Um2552–C2556–U1955 by adopting the C3′ endo conformation, facilitating the H92–H71 interaction and thereby promoting Um2552 incorporation into the 50S subunit (Fig. 6B).

In the crystal structure of the 50S subunit, Um2552 forms a base-triple along with C2556 and U1955 in H71 (Fig. 6A). If Um2552 were to adopt the C2′ endo ribose pucker, the base-triple would not be stabilized. Therefore, it is likely that the Um2552 modification stabilizes the base-triple Um2552–C2556–U1955 by adopting the C3′ endo conformation, facilitating the H92–H71 interaction and thereby promoting Um2552 incorporation into the 50S subunit (Fig. 6B).

The unique physicochemical feature of the 45S precursor is the consecutive alteration of its sedimentation coefficient in response to Mg\(^2+\) concentration (Fig. 1D). This structural property of the 45S precursor can be observed only in vitro. In the cell, where the Mg\(^2+\) concentration is sufficiently high, this structural change does not occur. However, this unique characteristic provides us with useful information about the structural properties of the 45S precursor, including flexible regions and solvent-accessible regions, during ribosome biogenesis in the cell. Our structural probing of 23S RNA from the 45S precursor revealed that helices 52, 56, 57, and 58 in domain III were more sensitive to hydroxyl radical cleavages at low Mg\(^2+\) concentration than at high Mg\(^2+\) concentration (Fig. 5 B and C and Fig. S9), indicating that these regions undergo Mg\(^2+\)-dependent structural changes. Intriguingly, H52 in domain III associates with H66 in domain IV via a “kissing-loop” interaction (Fig. 5E). Based on the observation that the interacting loops in both helices exhibited low Mg\(^2+\)-specific cleavage, it is likely that the H52–H66 interaction in the 45S precursor is stabilized at high Mg\(^2+\) concentration. In the crystal structure of the 50S subunit, four Mg\(^2+\) ions are present in H52 (Fig. 5E) whereas no Mg\(^2+\) is present in H66 (26). Mg\(^2+\) ions are likely to stabilize H52 to facilitate the kissing-loop interaction with H66, allowing the association between domains III and IV (Fig. 5 C and E). Moreover, H52 interacts with L28 (Fig. 5E and Fig. S11), which dissociates from the 45S precursor after high-salt stripping (Fig. 4B). Destabilization of the H52–H66 interaction might affect the stable association of L28. Because L28 bridges domains I, III, and V of the 23S rRNA (Fig. S11), correct positioning of L28 seems to stabilize domain III by strengthening its interactions with other domains. L28 also associates with H11 (domain V), which contained 45S-specific cleavage sites (Figs. S10B and S11). H11 is situated in close proximity to H13, where L35 binds (Fig. S10B); therefore, L35 incorporation might contribute to the stable binding of L28. Taken together, these observations indicate that, within the 45S precursor, interdomain interactions within the 23S rRNA are not yet stably established. Instead, domain III has a flexible structure that can be stabilized by Mg\(^2+\) ions. The alteration of the sedimentation coefficient of the 45S precursor seems to be primarily due to stabilization of interdomain interactions.
interactions and condensation of domain III in response to increasing Mg\(^{2+}\) concentration.

In this study, we clearly demonstrate that \textit{rlmE} and \textit{rpmJ} interact genetically, strongly suggesting that Um5252 and L36 play pivotal roles in late steps of the 50S subunit assembly. \textit{rlmE} was much more cold-sensitive (Fig. 4E), and accumulated much higher levels of the 45S precursor, than \textit{rpmJ} (Figs. 1C and 4D). In addition, L36 was completely absent from the 45S precursor in \textit{ΔrlmE} (Fig. 4C) whereas \sim 40\% of 45S precursor in \textit{ΔrpmJ} contained Um5252 (Fig. 4F), indicating that L36 incorporation is not required for Um5252 formation. These observations suggest that structural rearrangements of the 45S precursor induced by Um5252 are involved in incorporation of L36.

On the basis of the genetic and biochemical observations in this study, we propose a mechanistic model for late steps of the 50S subunit assembly triggered by the RlmE-mediated Um5252 modification (Fig. 6 B and C). Detailed discussion of this model is provided in SI Discussion. Um5252 stabilizes the A-loop structure and the Um2552–C structure of late assembly of the 50S subunit differ between \textit{E. coli} and \textit{B. subtilis} (39 as a platform for 5S rRNA and the other components of the central protuberance. Stable binding of L16 and L27 plays a critical role in spanning domains V and II and properly positions H38–39. A platform for 5S rRNA and the other components of the central protuberance (Fig. 6C). Association of domains V and II, as well as L27 binding to H86, affects L35 incorporation. L35 might affect stable binding of L28, which bridges domains I, III, and V. Taken together, these data indicate that all of the events in late steps of the 50S subunit assembly triggered by Um5252 formation take place on the interface side of the 45S precursor, including the central protuberance (Fig. 6B). The interface is not formed until a late step of 50S assembly; therefore, immature 50S cannot associate with the 30S subunit to produce unproductive 70S ribosomes incapable of participating in translation.

The 45S particle is also found in \textit{Bacillus subtilis}. Ribosomal (B. subtilis) 40S particles in the gradient were fractionated on a Piston Gradient Fractionator (BIOCOMP), and the A\textsubscript{520} was measured using a UV monitor and plotted as a function of time (10 min at 4 °C after addition of aluminia) for each fraction. The cell lysates were digested into nucleosides at 37 °C for 1 h in 60 μL of reaction mixture consisting of 25 mM ammonium acetate (pH 7.2), 1.33 mM nucleotides (μL/μL), and subjected to Q Exactive Hybrid Quadrupole-Orbitrap Mass Spectrometer (Thermo Fisher). One-tenth of the mixture was dried in vacuo, dissolved in 20 μL of 0.1% formic acid (FA), and analyzed on an Xcalibur 1200 elite (Thermo Fisher) apparatus. The mass range was narrowed to 1000–3000 Da, and the resolving power was set to 60,000 at 200 Da.

### Materials and Methods

#### Strains and Plasmid Constructions.

The \textit{E. coli} deletion strains \textit{ΔrlmE}:Km\(^{r}\) (\textit{JW3146}) and \textit{ΔrpmJ}:Km\(^{r}\) (\textit{JW3261}) and their parental strain \textit{BW25113} were provided by the National Institute of Genetics, Japan. We disrupted \textit{rrnB} (ribonuclease I) and \textit{rlmE} in \textit{BW25113} with the cat cassette by one-step gene inactivation (43) to yield strains \textit{ΔrrnB}:Cm\(^{r}\) and \textit{ΔrlmE}:Cm\(^{r}\), respectively. Single-colony isolation was repeated twice for all strains constructed in this study. Knockout was confirmed by colony PCR. \textit{ΔrlmE}:Cm\(^{r}\) was used as a reagent for construction of strains \textit{ΔrlmE}:Km\(^{r}\), \textit{ΔrrnB}:Km\(^{r}\), and \textit{ΔrlmE}:Cm\(^{r}\) and \textit{ΔrrnB}:Cm\(^{r}\), \textit{ΔrlmE}:Cm\(^{r}\) by P1 transduction. These strains were used for preparation of 45S precursors for a series of biochemical analyses. To construct the double-deletion strain, \textit{ΔrrnB}/ΔrpmJ, \textit{ΔrrnB}:Cm\(^{m}\) was introduced into the \textit{ΔrrnB}:Km\(^{m}\) strain by P1 transduction. \textit{dbpA} and \textit{rpmJ} were PCR-amplified from the \textit{E. coli} and \textit{B. subtilis} genomes by using the primer pair dbpA and rpmJ, respectively.

The amplified products \textit{dbpA} and \textit{rpmJ} were inserted into the XhoI/PstI and Ncol/PstI sites of pBAD-CMYC-HisA (Invitrogen) to yield pBAD-dbpa and pBAD-rpmJ, respectively. In pBAD-dbpa, the \textit{R331A} mutation was introduced using the QuikChange site-directed mutagenesis kit (Agilent Technologies). The \textit{E. coli} \textit{Δr} C\textsubscript{rrnB} strain \textit{KT101} (8), which harbors \textit{pbR101} (44) encoding the \textit{rrnB} operon and \textit{rnb} as a counter selectable marker, was used to examine the U2552C mutation in the 23S rRNA. Disruption of \textit{rlmE} in \textit{KT101} was performed by P1 transduction to generate \textit{KT101/ΔrlmE}. \textit{pbR101} in WT \textit{KT101} or in \textit{KT101/ΔrlmE} was completely replaced by plasmid pBR103, which harbors the \textit{rnb} operon with or without the U2552C mutation, to yield a series of \textit{KT103 strains as described (8). Sequences of primers used in these steps are listed in Table S2.

#### Analysis and Preparation of Ribosomes and Subunits by SDS.

\textit{E. coli} strains were cultivated in LB medium at 37 °C or 22 °C with vigorous shaking until \textit{A\textsubscript{540}} reached 0.4–0.6, quickly chilled by incubation on ice for 10 min, and harvested by centrifugation. To inhibit transcription, rifampicin was added to a final concentration of 250 μg/mL. For overexpression of the \textit{DbpA} \textit{R331A} mutant, 0.2% L-arabinose and 100 μg/mL ampicillin were added before inoculation of the preculture. Cell pellets from 50-ml cultures were resuspended in 6 M \textit{Mg(OAc)}\(_2\), 100 mM \textit{NH}_{4}Cl, 20 mM Hepes-KOH (pH 7.6), 6 mM \textit{β}-mercaptoethanol and lyzed by lysozyme treatment (1.5 mg/mL) followed by 1–3 freeze-thaw cycles. Cell lysates were clarified by centrifugation at 20,000 × \textit{g} for 15 min at 4 °C. The cleared lysate (15 \textit{A\textsubscript{520}} units) was layered on top of a sucrose gradient (10–40% [wt/vol]) in RBS buffer A and B separated by field centrifugation. The 

Ribosomal precursors in the gradient were fractionated on a Protein Gradient Fractionator (BIOCOMP), and the \textit{A\textsubscript{520}} was measured using a UV monitor (AC-5200; ATTO).

Ribosomal precursors in the gradient were fractionated on a Protein Gradient Fractionator (BIOCOMP), and the \textit{A\textsubscript{520}} was measured using a UV monitor (AC-5200; ATTO).

Ribosomal precursors in the gradient were fractionated on a Protein Gradient Fractionator (BIOCOMP), and the \textit{A\textsubscript{520}} was measured using a UV monitor (AC-5200; ATTO).
mass spectrometry (Thermo Fisher Scientific) with an Ultimate 3000 system (Thermo Fisher Scientific). The nucleosides were chromatographed on a ZIC-HILIC column (150 mm × 4.6 mm i.d., 3.5 μm; Merck Millipore) connected to a guard column (20 mm × 2.1 mm i.d., 5 μm; Merck Millipore) at a 0.1 mL/min flow rate of a multistep linear gradient of solvent A ([5 mM NH4OAc (pH 5.3) in water]) and B (acetonitrile): 90–40% B for 0–30 min, 40% B for 30–40 min, 40–90% B for 40–45 min, and 90% B for 45–55 min (45). The eluent was directly conducted to an electrospray ion source of the mass spectrometer and scanned with a positive polarity mode over an m/z range of 240–300 through analysis by capillary liquid chromatography (LC) coupled with nano-electrospray (ESI)/mass spectrometry (MS) on a linear ion trap-Orbitrap hybrid mass spectrometer (LTQ Orbitrap XL; Thermo Fisher Scientific) as described (46, 47). The m/z1915-containing fragment ([PAAC mβΑyΑyΑΑGp; MW 3,550.50] and the m/z1915-containing fragment ([PAACPAAPyACGP; MW 3,536.49] were detected as multiply charged negative ions. Other m/z sites were analyzed by a primer extension method (SI Materials and Methods and Fig. 3I). To analyze methylation status at position 25S2, a synthetic DNA (20 pmol) complementary to G2579–G2579 of E. coli 23S RNA (Table S2) was mixed with 2.0 A260 of total RNA or 20 pmol of isolated 23S RNA in the presence of 50 mM Hepes-KOH (pH 7.6) and 100 mM KC1. The mixture was heated at 90 °C for 5 min and then gradually cooled to 37 °C for 2.5 h, to allow the DNAs to form a heteroduplex. After adding 1 μM NaOAc (pH 5.2) to a final concentration of 50 mM, the uncovered regions of 25S RNA were removed by digestion with 20 USPF units of RNase T1 and 0.5 μg of RNase A; digests were incubated for 60 min on ice. The heteroduplex was extracted with phenol/ chloroform and precipitated with ethanol. The precipitate was resolved by 15% denaturing PAGE and visualized by staining with SYBR Gold (Molecular Probes). The heteroduplex was excised from the gel, eluted from the gel piece for 3 h at 37 °C in elution buffer [1 mM EDTA, 0.1% SDS, 0.4 M NaOAc (pH 5.0)], and precipitated with isopropanol. One-tenth of the specimen was digested with RNase T1, and subjected to MS analysis as described above.

Structural Probing of 23S rRNAs. Hydroxyl radical probing was performed essentially as described (48) with slight modifications. Twenty picomoles of 50S subunit or 45S precursor was dissolved in 97 μL of low Mg2+ solution (0.5 mM MgOAc2, 100 mM NH4Cl, 20 mM Hepes-KOH (pH 7.6), 6 mM β-mercaptoethanol) or 97 μL of high Mg2+ solution (10 mM MgOAc2, 100 mM NH4Cl, 20 mM Hepes-KOH (pH 7.6), 6 mM β-mercaptoethanol). To get enough hydroxyl radicals by the Fenton reaction, 1 μL of Fe2+-EDTA solution [100 mM Fe(SO4)2(NH4)2H2O and 200 mM EDTA], 1 μL of 500 mM ascorbic acid, and 1 μL of 10% H2O2 was added to each mixture. The reaction was carried out on ice for 10 min and then terminated by addition of 40 μL of 0.1 M thiourea. Samples were precipitated in ethanol and resuspended in dH2O. rRNAs were extracted by TRizol LS followed by two chloroform treatments, precipitated with isopropanol, and then reprecipitated with ethanol. Primer extension was performed basically as described (49) with a slight modification. Sequences of primers (Table S1) were listed in Table S1. 32P-labeled primer (1–2 × 106 cpm), 0.2 pmol rRNAs, and 0.77 mM dNTPs was incubated at 65 °C for 10 min and then incubated on ice for 1 min to hybridize the primer with the rRNAs. Next, 2 μL of 5X FS buffer (Invitrogen), 0.5 μL of 0.1 M DTT, and 0.5 μL of d[γ-32P]-dATP or d[γ-32P]-dGTP (3.2 μCi/μL) was added, and the mixture was preincubated at 55 °C for 1 min. The reaction was started by addition of 0.5 μL (100 units) of SuperScript III (Invitrogen), incubated at 55 °C for 10 min, terminated by addition of 0.5 μL of 4 M NaOH, and boiled at 95 °C for 5 min to degrade template rRNAs. After the reaction, the cDNAs were ethanol-precipitated and subjected to 8% PAGE in the presence of 7 M urea. The gel was exposed to an imaging plate overnight, and the radiolabeled bands were visualized on an FLA-7000. The intensity of bands in each lane was quantified using the MultiGauge software (FujiFilm). After subtracting the background intensity of non-treated lanes, the intensities with intensity >6000QL were selected. In the high-Mg2+ condition, if the band intensity in the 45S precursor was >1.5-fold higher than the intensity of the corresponding site in the 50S subunit, the site was defined as “45S-specific cleavage.” If the band intensity in the low-Mg2+ condition was >1.5-fold higher than the intensity of the corresponding site in the high Mg2+ condition, the site was defined as “low Mg2+–specific cleavage.” This experiment was repeated several times to confirm the reproducibility of the results.

Enzymatic Formation of the 50S Subunit from the 45S Precursor in Vitro. A crude ribosome fraction containing the 45S precursor was prepared from ΔrmlEΔsrmA, as described above. In vitro methylation of the crude ribosome fraction was performed at 37 °C for 2.5 h in a 60-μl reaction mixture containing 100 mM NH4Cl, 20 mM Hepes-KOH (pH 7.6), 6 mM MgOAc2, 6 mM β-mercaptoethanol, 1 mM ATP, 1 mM GTP, 1 A260 unit of 45S precursor, and 70 μg of the wash fraction (SI Materials and Methods). After the reaction, the buffer was exchanged with RBS buffer A three times by an ultrafiltration using the Amicon Ultra-0.5 100K device (Merck Millipore). Then, the mixtures were resuspended in RBS buffer A, layered on top of a sucrose gradient (10–40% w/vtov) in RBS buffer A and ultracentrifuged in a Beckman SW-41 Rotor at 37,000 rpm for 5 h at 4 °C. For in vitro methylation of the isolated 45S precursor, the reaction was performed at 37 °C for 2.5 h in a 200-μL reaction mixture containing 100 mM NH4Cl, 20 mM Hepes-KOH (pH 7.6), 8 mM MgOAc2, 6 mM β-mercaptoethanol, 1 mM AdoMet, 1 mM ATP, 1 mM GTP, 1 A260 unit of 45S precursor, and 70 μg of the wash fraction (SI Materials and Methods). After the reaction, the buffer was exchanged with RBS buffer A, and the mixture was then analyzed by SDG as described above.

To evaluate the ability of the newly synthesized 50S subunit to associate with the 30S subunit, the 50S subunit isolated from the ΔrmlEΔsrmA strain was added to the reaction mixture immediately after the methylation reaction. The buffer was exchanged with RBS buffer B three times as described above, followed by incubation at 37 °C for 30 min. The mixture was then analyzed by SDG as described above.

ACKNOWLEDGMENTS. We thank the members of the Tsutomu Suzuki laboratory for materials, technical advice, and many fruitful discussions. This work was supported by Grants-in-Aid for Scientific Research on Priority Areas from the Ministry of Education, Science, Sports, and Culture of Japan (to Tsutomu Suzuki) and by a grant from the New Energy and Industrial Technology Development Organization (to Tsutomu Suzuki).


