An open interface in the pre-80S ribosome coordinated by ribosome assembly factors Tsr1 and Dim1 enables temporal regulation of Fap7

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ABSTRACT

During their maturation, nascent 40S subunits enter a translation-like quality control cycle, where they are joined by mature 60S subunits to form 80S-like ribosomes. While these assembly intermediates are essential for maturation and quality control, how they form, and how their structure promotes quality control, remains unknown. To address these questions, we determined the structure of an 80S-like ribosome assembly intermediate to an overall resolution of 3.4 Å. The structure, validated by biochemical data, resolves a large body of previously paradoxical data and illustrates how assembly and translation factors cooperate to promote the formation of an interface that lacks many mature subunit contacts but is stabilized by the universally conserved methyltransferase Dim1. We also show how Tsr1 enables this interface by blocking the canonical binding of eIF5B to 40S subunits, while maintaining its binding to 60S. The structure also shows how this interface leads to unfolding of the platform, which allows for temporal regulation of the ATPase Fap7, thus linking 40S maturation to quality control during ribosome assembly.

Keywords: ribosome assembly; small ribosome subunit; cryo-EM

INTRODUCTION

Ribosome assembly involves the cotranscriptional processing and folding of four ribosomal (r)RNAs that is coupled to binding of ribosomal proteins (r-proteins) via a machinery of nearly 200 assembly factors (AFs). These AFs represent one way in which the cell ensures that this complex ribonucleoprotein complex assembles faithfully as the ~2000 ribosomes/minute are synthesized in a yeast cell (Warner 1999). Another mechanism by which the cell avoids dead-end or misfolded ribosomes is that ribosome assembly occurs through a series of partially assembled intermediates that serve as checkpoints, first in the nucleolus and then in the cytoplasm (Klinge and Woolford 2019). Although some intermediates were identified biochemically in the 1970s and 1980s, others have only recently been discovered and their structures been determined, in part because of advances in cryo-EM technologies. Ribosomes are central to the cellular function across all of biology, so misfolded ribosomes severely impact the health of the cell because they can lead to errors in translation, stalling on the mRNA (Cole et al. 2009). Visualizing how those intermediates function in quality control of ribosome assembly will advance our understanding of how cells protect themselves from misfolded ribosomes.

After initial transcription and assembly in the nucleolus, largely assembled 40S precursors that retain only seven AFs (Tsr1, Dim1, Nob1, Pno1, Enp1, Ltv1, and Rio2) are exported into the cytoplasm (Baßler and Hurt 2019; Klinge and Woolford 2019). Tsr1 is a large, GTPase-like protein that shares domain homology with other GTPases involved in initiation, like eIF5B or elongation factors EF-Tu (McCaughan et al. 2016), but lacks the GTPase...
active site. Dim1 is a universally conserved methylase, called KsgA in bacteria, that methylates the small subunit rRNA (O’Farrell et al. 2006). Nob1 is a PIN-domain containing endonuclease that is responsible for cleavage of the 3’ end of the 18S precursor (20S rRNA) to create the mature-length 18S rRNA (Pertschy et al. 2009). Pno1 contains two tandem KH homology domains that bind the rRNA (Vanrobays et al. 2008) and acts as a structural partner to Nob1, regulating its endonuclease activity (Woolls et al. 2011). Enp1 and Ltv1 work together to block Rps3 from adopting its mature position to guide assembly of the 40S head (Strunk et al. 2011; Mitterer et al. 2016; Johnson et al. 2017; Collins et al. 2018). Their release from the solvent side of the pre-40S in the cytoplasm, driven by Hrr25/CK1δ phosphorylation (Schäfer et al. 2006; Ghalei et al. 2015; Mitterer et al. 2019), commits the maturing pre-40S to a translation-like intermediate bound to mature 60S subunits (Strunk et al. 2012; Ghalei et al. 2015). Finally, Rio2 is a kinase that binds to the interface side of pre-40S, blocking the eIF1A binding site to prevent premature translation but releasing before formation of the translation-like intermediate (Ferreira-Cerca et al. 2012; Strunk et al. 2012; Huang et al. 2020).

Ribosome maturation is coupled to quality control in a translation-like cycle where pre-40S ribosomes bind mature 60S subunits in an elf58-dependent manner to produce 80S-like ribosomes (Lebaron et al. 2012; Strunk et al. 2012; Ghalei et al. 2017; Huang et al. 2020). These 80S-like intermediates preserve Tsr1 and Dim1 (Strunk et al. 2012; Ghalei et al. 2017; Shayan et al. 2020), and thus present a paradox, because structures of pre-40S ribosomes that contain the above seven AFs show that Tsr1 and Dim1 sterically block 60S binding at the interface (Strunk et al. 2011; Heuer et al. 2017; Johnson et al. 2017; Scaiola et al. 2018). Specifically, the position of Tsr1 within the stable pre-40S intermediate sterically blocks the binding of elf58, which is required for formation of both 80S and 80S-like ribosomes (Lebaron et al. 2012; Strunk et al. 2012; McCaughan et al. 2016). In addition, Tsr1 promotes a conformation of the decoding site helix, h44, which is incompatible with the binding of 60S subunits (Strunk et al. 2011; Heuer et al. 2017; Scaiola et al. 2018). However, its position is variable and contributes to heterogeneity in the pre-40S subunit (Johnson et al. 2017; Thoms et al. 2020). As it is positioned by their position as seen in prior 40S structures. Furthermore, because Dim1 sterically blocks the binding of elf1, thereby preventing premature translation initiation, its release must be regulated during ribosome assembly so translation initiation occurs only after both maturation and quality control are complete. Dim1 release occurs in 80S-like ribosomes and is catalyzed by the ribosome biogenesis ATPase Fap7 that binds Rps14 after testing their ability to faithfully translocate mRNA; bypass of this step allows release of ribosomes defective in reading frame maintenance into the translating pool (Granneman et al. 2005; Ghalei et al. 2017). Moreover, formation of 80S-like ribosomes is a quality control checkpoint that ensures only scanning-competent ribosomes are released into the translating pool, and bypass of these checkpoints releases ribosomes with initiation-defects into the translating pool (Huang et al. 2020). Finally, Nob1-dependent cleavage of the 3’ end of the rRNA occurs while this 80S-like is formed (Strunk et al. 2012). In summary, Tsr1 and Dim1-containing 80S-like ribosomes intermediates form in the final stages of ribosome maturation to ensure only translation-competent ribosomes enter the translation pool.

Despite their importance in quality control and maturation, the structure of 80S-like ribosomes has not been described. Thus, we do not have a structural understanding of how they are formed, despite the presence of the AFs Tsr1 and Dim1 that block subunit joining, and how their formation enables proofreading of scanning competence. Furthermore, how they enable temporal regulation of Fap7-dependent Dim1 release is also unknown. Therefore, we used single-particle cryogenic electron microscopy (cryo-EM) to visualize the structure of 80S-like ribosomes that accumulate upon Fap7 depletion (Ghalei et al. 2017). The structure is validated by biochemical data herein, as well as previous structural and biochemical data, illuminating how 80S-like ribosomes accommodate Dim1 and Tsr1, despite their position blocking subunit joining in earlier intermediates. Furthermore, structural and biochemical data also reveal how 80S-like ribosomes enable quality control.

RESULTS AND DISCUSSION

To better understand how 40S ribosome maturation is quality-controlled via structural and functional mimicry of translation events, we used cryo-EM to visualize the structure of 80S-like ribosomes. This structure was substantially different from mature 80S ribosomes because the interface between the mature 60S and pre-40S subunits is opened up, creating space between the subunits (Fig. 1A). This intermediate contains the interface AFs Dim1 and Tsr1, which are accommodated (together with h44) in the interface because of its opening (Fig. 1B). This significant rearrangement of the subunits answers one of the biggest questions about the structural nature of this intermediate that retains both Dim1 and Tsr1, despite their positions as steric blocks in previous pre-40S intermediates (Strunk et al. 2011; Heuer et al. 2017; Johnson et al. 2017; Scaiola et al. 2018). Tsr1 shifts modestly such that it detaches from the pre-40S head (Fig. 1C). Dim1 remains at
the interface, but is removed from h45 as described below (Fig. 1C).

These ribosome assembly intermediates were accumulated by Fap7 depletion (Strunk et al. 2012; Ghalei et al. 2017) and purified via a TAP-tag on Tsr1 (Lebaron et al. 2012; Ghalei et al. 2017; Mitterer et al. 2019; Shayan et al. 2020), which is fully functional (Supplemental Fig. S1a). SDS-PAGE, mass-spectrometry, and western blot analysis demonstrate that these intermediates retain the AFs Tsr1, Dim1, and Pno1, but Nob1 is substoichiometric (Supplemental Fig. S1b,c). Additionally, 60S ribosomal proteins and ribosome associated factors like the SSA/SSB chaperones were identified, consistent with the presence of 25S rRNA (Ghalei et al. 2017). Importantly, previous biochemical data confirmed that these intermediates are on the pathway because addition of recombinant Fap7/ATP releases Dim1 (Ghalei et al. 2017).

Initial analysis identified that two populations of particles could be visually differentiated in both raw fields of view and in 2D analysis (Supplemental Fig. S2a,b). One refined to a resolution of 3.6 Å and had features like mature 80S ribosomes (Supplemental Fig. S2c,d), similar to another 80S preribosome with features similar to mature ribosomes but lacking the AFs Tsr1, Dim1, and Pno1 (Sciola et al. 2018). In this subclass, the platform, beak, decoding helix, and B3 bridge are in their mature conformation and there is no density for any tRNA (Supplemental Fig. S2e). This subclass is not described further as its significance is unclear. The other subclass had strong 60S, but weaker pre-40S, density (Supplemental Fig. S2b). Further refinement improved the clarity of the pre-40S and 60S to overall resolutions of 3.7 Å and 3.4 Å, respectively (Supplemental Fig. S2c,d). Although the particles showed no preferred orientation, the resolution of pre-40S remained anisotropic (Supplemental Fig. S3a–c). Nevertheless, features at the core of the particle were consistent with those overall resolution measures (Supplemental Fig. S3d). Local classification described below clarified the interface and platform to ~5.5–8.5 Å-resolution (Supplemental Fig. S3e).

Immature subunit interface

One consequence of the opened 80S-like subunit interface is that fewer bridges are formed than in mature 80S (Fig. 2A). For example, the B1a/b/c bridges, which involve the 40S head, are not formed because the pre-40S head is far from the 60S central protuberance. Similarly, pre-40S turns away from 60S on the platform side and toward it on the beak side, thus bridge B7a cannot form because that side of the head is too distant from 60S. Further, the twisted position of pre-40S relative to 60S accommodates the immature position of h44, which is shifted left (Heuer et al. 2017; Sciola et al. 2018), preventing formation of the B3 and B5 bridges at the top of h44. In contrast, eukaryote-specific bridges at the 40S foot are largely maintained. In the case of a B5/B8-like bridge between h44 and Rpl23/uL14 and a B6-like bridge between h44 and Rpl24, the intersubunit connections shift due to the novel orientations of the subunits but involve analogous interactions between similar structural elements (Supplemental Fig. S4).

A striking difference between 80S-like and mature 80S ribosomes is the interaction between H69 and the small subunit. In mature ribosomes, H69 binds h44 (Ben-Shem et al. 2011) to establish the B2a bridge and its deletion results in a dominant lethal phenotype (Ali et al. 2006). This bridge is the target of RRF (Pai et al. 2008), which dissociates ribosomes, as well as some antibiotics (Borovinskaya et al. 2007; Wang et al. 2012; Prokhorova et al. 2017), supporting its importance. In 80S-like ribosomes, H69 binds h24, which moves substantially from its mature position (Fig. 2B). Two helical densities can be assigned to H69 and h24 with high confidence, however the moderate resolution of the subunit interface prevents us from modeling atomic interactions between H69 and h24. The local resolution of one helix is ~5.5 Å, lower than 60S but sufficiently well-resolved to model H69 in its mature position (Fig. 2C; Supplemental Fig. S1e). The tip of the other helix is resolved at ~8.5 Å; however, it connects to the well-resolved base of a repositioned h24, strongly suggesting that the RNA that binds H69 derives from h24 (Fig. 2C; Supplemental Fig. S1f). We validated interactions between Dim1 and h24, as well as Dim1 and 60S, in biochemical experiments described below.
Dim1 stabilizes h24 and binds 60S

In pre-40S, Dim1 sterically blocks the position of the mature B2 bridge between h44 and H69, inhibiting subunit joining in canonical 80S ribosomes (Strunk et al. 2011; Heuer et al. 2017; Johnson et al. 2017; Scaiola et al. 2018). This conflict would still exist if Dim1 remained in that position within 80S-like ribosomes, despite their rearranged interface, because H69 remains the main connection between the subunits. Instead, Dim1 repositions along with h24, moving above H69 to alleviate the steric conflict (Fig. 3A; Supplemental Movie S1). This movement essentially releases Dim1 from pre-40S, maintaining contact only with h24, while promoting a new interaction with 60S, thus priming Dim1 for release.

We created a Dim1 variant and tested its effects on subunit joining to validate the position of Dim1, which, despite local classification, is resolved at ~8.5 Å. The variant was designed to also probe the importance of the altered contacts between Dim1 and h24. In 80S-like ribosomes residues altered in Dim1-IKN (I250A/K253A/N254A) face h24 (Fig. 3B). In pre-40S ribosomes, these amino acids appear to be solvent exposed (Fig. 3C; Mitterer et al. 2019). To test if the altered residues are important for formation of 80S-like ribosomes, we used an in vivo assay in which variants were expressed in a galactose-inducible/glucose-repressible Dim1/Fap7 yeast strain, supplemented with plasmids encoding wild-type or variant Dim1 and inactive Fap7. In strains competent for 80S-like ribosome formation, accumulation of 80S-like ribosomes upon inactivation of Fap7 leads to cosedimentation of pre-18S and 25S rRNA in 80S-sized fractions (Strunk et al. 2012; Ghalei et al. 2017). Dim1-IKN expresses and shifts the equilibrium between 40S and 80S-like ribosomes away from 80S-like ribosomes (Fig. 3D,E), demonstrating a role for these residues in the formation of 80S-like ribosomes. Thus, these biochemical data validate the novel position of Dim1, and demonstrate the importance of its binding to both h24 and 60S for the formation of 80S-like ribosomes, thereby establishing a role for Dim1 in stabilizing 80S-like ribosomes.

Tsr1 promotes the opened interface

In earlier 40S assembly intermediates, Tsr1 binds between the body and head, with its amino-terminal α-helix inserted behind h44 to force it away from the body, blocking formation of canonical 80S ribosomes (Supplemental Fig. S5a; Strunk et al. 2011; Heuer et al. 2017; Johnson et al. 2017; Scaiola et al. 2018). In 80S-like ribosomes, the
carboxy-terminal domain is repositioned toward the beak’s tip via a rigid-body movement, pivoting around Ala63 within the amino-terminal helix, which remains behind h44 in a well-ordered region of Tsr1 (Fig. 3A; Supplemental Fig. S5b,c; Supplemental Movie S2).

To test the importance of the amino-terminal helix of Tsr1, we deleted the amino-terminal most amino acids including Ala63 (Tsr1-ΔN74) and assessed the effect on cell growth and formation of 80S-like ribosomes. In yeast strains where endogenous Tsr1 is under a galactose-inducible/glucose-repressible promoter, providing Tsr1-ΔN74 on a plasmid produces a strong growth phenotype (Supplemental Fig. S5d). In contrast, deleting just the amino-terminal 43 amino acids has no growth defect (Supplemental Fig. S5e). This isolates the functionally important region of Tsr1 to the helix inserted under h44.

Next, we used the in vivo subunit joining assay described above to test if the Tsr1-ΔN74 variant impairs the formation of 80S-like ribosomes. Sucrose-gradient fractionation of Fap7-depleted cells with wild type or Tsr1-ΔN74 demonstrate that the ratio of 20S pre-rRNA in 80S-like ribosomes relative to 40S-sized precursors is strongly reduced (Fig. 4A). This observation is supported by genetic interactions showing that Tsr1-ΔN74 behaves like another recently characterized Tsr1 mutant that blocks subunit joining, Tsr1-RK (Supplemental Fig. S5f–g; Huang et al. 2020). Tsr1-RK and Tsr1-ΔN74 are both rescued by the internal Tsr1-deletion, they are both synthetically sick with Ltv1 deletion, as well as a mutation in Rps15_YRR that affects subunit joining. Consistent with the biochemical data, these genetic interactions suggest that akin to Tsr1-RK, Tsr1-ΔN74 impairs subunit joining.

Moreover, the data also show a depletion of Tsr1-ΔN74 from 80S-like ribosomes, while they are bound efficiently to pre-40S ribosomes (Fig. 4B), again supporting the interpretation that this helix is required for the formation of 80S-like ribosomes. The increased amount of free Tsr1 in the Tsr1-ΔN74 cells is likely due to a combination of factors, including higher expression levels of Tsr1-ΔN74 (Supplemental Fig. S5h) and its inability to form 80S-like ribosomes.
ribosomes. Notably, weaker binding of Tsr1-ΔN74 cannot account for the observed phenotypes, because the growth of cells containing Tsr1-ΔN74 is not affected by the Tsr1 levels (Supplemental Fig. S5h,i). Thus, our biochemical and genetic data demonstrate a role for the amino-terminal helix of Tsr1, which inserts under h44 and pivots in the transition from pre-40S to 80S-like ribosomes in the formation of 80S-like ribosomes, perhaps in part by facilitating this rigid body movement.

The initiation factor eIF5B is required for formation of 80S-like ribosomes (Lebaron et al. 2012), but its binding is incompatible with the position of Tsr1 in previously described earlier pre-40S intermediates (Strunk et al. 2011; Heuer et al. 2017; Johnson et al. 2017; Sciaiola et al. 2018). To visualize if Tsr1 repositioning enables eIF5B binding, we superimposed 60S from the mature 80S-eIF5B complex onto this intermediate to place eIF5B (Fig. 4C,D; Supplemental Fig. S6a,b; Fernández et al. 2013). This analysis suggests that the postulated clashes between Tsr1 and eIF5B (McCaughan et al. 2016) are largely resolved in 80S-like ribosomes. Notably, it is the opened subunit interface that provides space for concurrent binding of eIF5B and Tsr1 at the subunit interface, not Tsr1 repositioning.

In our previous data set of earlier 40S assembly intermediates (Johnson et al. 2017), Tsr1 is similarly rotated in a small population of molecules (Supplemental Fig. S6c). In this low-resolution structure, Tsr1 has a disordered carboxy-terminal domain due to its mobility in isolated pre-40S and/or the low number of particles. In addition, the same position of Tsr1 is also observed in a recent structure of the SARS-CoV2 protein Nsp1 bound to pre-40S subunits (Thoms et al. 2020). Together, these observations suggest that its movement away from h44 to the beak is intrinsic to Tsr1, and not induced, for example, by eIF5B.

The structure of 80S-like ribosomes clarifies multiple roles for Tsr1 in the formation of 80S-like ribosomes versus canonical 80S ribosomes: (i) Tsr1 forces out h44, thereby sterically preventing the subunits from close approach to form the canonical subunit bridges in the head (Fig. 1A; Supplemental Fig. S5b); (ii) Tsr1 stabilizes the interface due to its interaction with both pre-40S and 60S (Supplemental Fig. S6d); (iii) Tsr1 blocks access of eIF5B to its typical binding site on 40S, instead enforcing a position where eIF5B blocks formation of the strong and early-forming B3 bridge (Fig. 4C,D; Lu et al. 2014). Thus, the structure suggests that by modulating the position of eIF5B on the pre-40S Tsr1 steers the subunits away from the canonical B3-containing interface into 80S-like ribosomes.
Head folding in nascent pre-40S

Previous structures show that earlier pre-40S lack Rps10/eS10, Rps26/eS26, and Asc1 (Strunk et al. 2011; Heuer et al. 2017; Johnson et al. 2017; Scaiola et al. 2018), and that the tip of h31 remains unfolded (Heuer et al. 2017; Scaiola et al. 2018). In 80S-like intermediates, Asc1 and Rps10 locate to their mature position (Supplemental Fig. S7), indicating these proteins are recruited prior to Fap7 activity, as expected from biochemical analyses (Strunk et al. 2012). Furthermore, the tip of h31 is visible (Supplemental Fig. S8). In earlier pre-40S intermediates, Tsr1 binds adjacent to the last ordered nucleotide in h31, and Rio2 binds adjacent to the h31's mature location (Strunk et al. 2011; Heuer et al. 2017; Johnson et al. 2017; Scaiola et al. 2018). Thus, we suggest that h31 folding arises from dissociation of Rio2 and detachment of Tsr1 from the head. This coincides with the beak moving to its mature position while the head straightens (Supplemental Fig. S9).

Platform unfolding

Extensive differences between earlier pre-40S subunits and 80S-like ribosomes observed at the platform, which is opened toward 60S via a rigid body motion (Fig. 1A). This movement is presumably facilitated by repositioning of h24 from the platform to the subunit interface, accompanied by disorder in the tip of h23 and partial loss of Rps1 and Rps14, as previously observed (Strunk et al. 2012), requiring local classification to improve its resolution. The other 80S ribosome subclass we observe has a canonical platform (Supplemental Fig. S2e), ruling out artifacts during sample preparation as causative for the rearrangements. Further, the particles do not adopt a preferred orientation in the ice (Supplemental Fig. S3a), as expected if they were interacting with the air-water interface to unfold the platform. We therefore conclude that these changes represent folding transitions during 40S subunit maturation. A recently determined structure of pre-40S, also isolated via TAP-Tsr1 but without the Fap7 deletion, describes similar unfolding of the platform (Shayan et al. 2020) but where the platform shifts toward h44, rather than opening out from the head.

To interpret the platform structure, we placed Rps1•Rps14•Pno1 from the earlier pre-40S intermediates (Scaiola et al. 2018) in the platform density with rigid body fitting, assuming the Rps14•Pno1 dimer remains unaltered (Fig. 5A). To validate this assumption, we produced mutants in the Rps14•Pno1 interface, Rps14-R107E and Pno1-QDF (Q153E/D157R/F237A, Fig. 5B). These variants were tested for growth defects in galactose-inducible/glucose-repressible Rps14 and Pno1 strains, respectively, where both demonstrated growth phenotypes (Supplemental Fig. S10a,b). Because Pno1 and Rps14 are both bound (but do not yet interact) in early 90S precursors (Barandun et al. 2017; Cheng et al. 2017; Sun et al. 2017), we used northern analysis to confirm that these mutations do not substantially affect early maturation events. Pno1 or Rps14 depletion reduce 20S rRNA >20-100-fold, respectively (Supplemental Fig. S10c,d; Barandun et al. 2017; Cheng et al. 2017; Sun et al. 2017). In contrast, 20S rRNA levels are reduced approximately twofold in the Rps14-R107E and Pno1-QDF mutants, and accumulation of 23S rRNA is not observed. Thus, lethal or near-lethal growth defects from these mutants do not arise from early maturation defects. Next, we used sucrose-gradient fractionation to assess if Pno1 binding to 80S-like ribosomes is affected in these mutants. 80S-like ribosomes were accumulated via depletion of Fap7 (Strunk et al. 2012), and western analysis was used to probe the sedimentation of Pno1. As expected from a weakened Pno1•Rps14 interface, the fraction of free Pno1 increases in both mutants relative to their isogenic wild-type controls (Fig. 5C,D). These data strongly suggest that the Pno1•Rps14 interface is maintained in 80S-like ribosomes, supporting placement of the unchanged Rps1•Rps14•Pno1 complex in the platform density. Note that Nob1 was not identified in the EM density, presumably due to its substoichiometric presence (Supplemental Fig. S1b,c) and its mobility, which has precluded its visualization in earlier studies that did not use crosslinking (Strunk et al. 2011; Heuer et al. 2017; Johnson et al. 2017; Scaiola et al. 2018). Similarly, density corresponding to ITS1 is not visible, as also observed in previous structures (Strunk et al. 2011; Heuer et al. 2017; Johnson et al. 2017; Scaiola et al. 2018; Shayan et al. 2020).

Regulation of Fap7 activity at a late stage in ribosomal assembly

The intermediates studied here were purified from Fap7-depleted cells. Consistently, addition of Fap7•ATP to these intermediates leads to Dim1 release (Ghalei et al. 2017). To confirm that the molecules can bind Fap7 stably, we developed a binding assay where we add either Fap7 or Fap7•Rps14 in the presence of either ATP or the nonhydrolyzable ATP-analog AMPPNP to the 80S-like ribosomes herein, and assay for binding by cosedimentation with ribosomes. These data show that Fap7 binds these intermediates. Addition of Rps14 stabilizes binding two- to threefold, consistent with reduced occupancy of Rps14 in the purified particles. In addition, AMPPNP stabilizes binding over ATP (Fig. 6A), consistent with ATPase-dependent release of Dim1 (and presumably Fap7) from 80S-like ribosomes (Ghalei et al. 2017). Unfortunately, we were unable to obtain a structure from these Fap7-bound ribosomes, presumably due to partial occupancy. Nonetheless, we could model Fap7 into the platform by docking the crystal structure of the Fap7•Rps14 dimer to better understand how Fap7 functions in Dim1 release. Two Fap7•Rps14 dimers were present in the asymmetric unit of the crystals.
 Loc’h et al. 2014), but after superimposition of Rps14, one molecule of Fap7 is positioned between Dim1 and Rps14 (Fig. 6B), consistent with biochemical data that show Fap7 bridges these proteins (Ghalei et al. 2017). Intriguingly, the second molecule of Fap7, which was originally proposed to be the biologically relevant dimer (Loc’h et al. 2014), clashes with the platform (Fig. 6C).

To validate the position of Fap7, we created mutations in Fap7 and Rps14 that would affect the interface that allows Fap7 to bridge Dim1 and Rps14. Rps14-K49E and Fap7-RYD (R114E/Y116A/D118K) alter residues that contact their binding partners only in the dockable interface, whereas Rps14-RVM (R41E/V42L/M46A) alters an arginine that binds Fap7 in both (Val42 and Met46 bind Fap7 in the dockable interface) (Fig. 6B, C). Each mutation results in a substantial growth phenotype (Fig. 6D). Furthermore, recombinant Fap7-RYD and Rps14-RVM bind Rps14 and Fap7 more weakly (Fig. 6E). Finally, as in inactive Fap7 (Ghalei et al. 2017), the growth phenotype from Fap7-RYD is partially rescued by a self-releasing Dim1 mutation, Dim1-EKR (Fig. 6F). Together, these experiments support this placement of Fap7 in 80S-like ribosomes. Additionally, this biochemically validated position of Fap7 between Rps14 and Dim1 also validates interpretation of the medium-resolution features of the map: Rps14 on the platform and Dim1 at the subunit interface.

Fap7 binds Rps14 (Hellmich et al. 2013; Loc’h et al. 2014; Ghalei et al. 2017) and the biochemical data above demonstrate the importance of their interaction for 40S maturation and Dim1 release (Ghalei et al. 2017). Rps14 is already bound in the earliest nucleolar 40S assembly intermediates (Barandun et al. 2017; Cheng et al. 2017; Sun et al. 2017), yet Fap7-dependent Dim1 release is one of the last cytoplasmic steps in maturation, raising the question how Fap7 recruitment to nascent ribosomes is regulated over the course of ribosome assembly. Docking of Fap7 onto earlier pre-40S intermediates answers this question because it shows that in the folded platform there is steric conflict between Fap7 and the rRNA, regardless of which Fap7•Rps14 dimer is docked onto the Rps14 position (Fig. 6G). Specifically, the Fap7 binding site on Rps14 that positions Fap7 as a bridge to Dim1 is blocked by h23. In contrast, in 80S-like ribosomes this steric conflict is relieved by...
FIGURE 6. Fap7 can only bind an opened platform. (A) Fap7 binds to purified 80S-like ribosomes. Fap7 binding in the presence and absence of Rps14 and ATP or AMPPNP was assessed in a pelleting assay. Ribosome pellets were probed for Fap7, Tsr1-TAP or Rps10 (left), and bound Fap7 was quantified relative to the amount of 80S-like ribosomes (Tsr1-TAP, right). Data are the average of two replicates and error bars indicate SEM. Arbitrary Units, A. U. (B) Docking Fap7•Rps14 (Fap7, chain G and Rps14, chain F) from PDB ID 4CW7 (Loc’h et al. 2014) onto Rps14 (orange) in 80S-like ribosomes places Fap7 (bright blue) in direct contact with Dim1 (dark blue), as predicted by previous biochemical data (Ghalei et al. 2017). The residues in Fap7-RYD (R114E/Y116A/D118K, cyan) and Rps14-RVM (R41E/V42L/M46A, red) are at the interface between Fap7 and Rps14 (formed between chains G and F). Rps14-K49E is in purple. In contrast, chain C from that structure clashes with the pre-40S platform. (C) An alternative dimer of Fap7•Rps14 reveals that Fap7 (cyan) would be in steric clash with the pre-40S platform. (D) Growth (left) or Fap7 expression levels (western blot, right) of Gal::Fap7 cells containing an empty vector (e.v.), wild-type Fap7, or Fap7-RYD or growth of ΔRps14B; Gal::Rps14A cells containing an empty vector (e.v.), wild-type Rps14, Rps14-RVM, or Rps14-K49E plasmids were compared by 10-fold serial dilution on YPD or YPGal plates. (E) Interface mutations in Fap7 or Rps14 weaken their binding affinity for each other. Shown are Coomassie-stained SDS-PAGE gels of protein binding assays on amylose beads of purified, recombinant MBP-Fap7 or MBP-Fap7-RYD and SUMO-Rps14 or SUMO-Rps14-RVM. In, input; Ft, flow-through; W, final wash; El, elution (left). Quantification of SUMO-Rps14 (S-Rps14) compared to MBP-Fap7 (M-Fap7) in elution normalized to wild-type (right). Data are the average of two replicates and error bars indicate SEM. (F) Doubling time, in minutes, of cells depleted of endogenous Dim1 and Fap7 (Gal::Dim1; Gal::Fap7) and transformed with plasmids encoding Dim1 or Dim1-EKR and either Fap7 or Fap7-RYD. The white column represents the expected doubling time if there was no rescue of Fap7-RYD by Dim1-EKR (See also Ghalei et al. 2017). The height of this column was calculated by multiplying the observed fold differences for each single mutation. The data are the average of 16–17 biological replicates and the error bars represent SEM. Unpaired t-test was performed comparing expected and actual doubling times of cells expressing Fap7-RYD and Dim1-EKR. (*) P-value = 0.030. (G) Both Fap7 and Rps14 interfaces clash in pre-40S ribosomes (Rps14 [orange], h23 [red], rRNA [gray] from PDB 6FA1/EMD-4128 [Sciaiola et al. 2018]). (H) The Fap7 binding site on Rps14 is blocked by h24 in nucleolar 90S preribosomes and pre-40S ribosomes.
opening of the platform, which repositions Rps14, and mobilizes h23 to expose the Fap7 binding site. Thus, unfolding of the platform in 80S-like ribosomes regulates Fap7 binding at a late stage of ribosomal assembly, when it acts to dissociate Dim1 (Fig. 6H).

80S-like ribosomes are formed in an elF5B-dependent manner during the maturation of 40S ribosomes (Lebaron et al. 2012; Strunk et al. 2012). Yet, elF5B is not an essential protein in yeast, raising the question whether there are elF5B-independent pathways in 40S maturation. While it is impossible to exclude such a possibility, we believe that the data herein provide evidence against this model as they demonstrate how 80S-like ribosomes are required for binding of Fap7, which is an essential protein. Thus, we believe that the nonessential nature of elF5B is simply a reflection of the ability to form 80S complexes without elF5B, which must also happen during translation in the elF5B deletion strains.

In summary, the structure of 80S-like ribosomes presented here, validated by genetic and biochemical data herein, as well as previous biochemical (Ghalei et al. 2017), mass spectrometry (Strunk et al. 2012), and crystallographic (Loc’h et al. 2014) data, reveal unexpected features that allow for reconciliation and explanation of many previous observations. (i) Relative to 80S ribosomes, 80S-like ribosomes display an opened interface whose formation is enabled by Tsr1: Tsr1 supports a conformation of h44, the decoding helix, that necessitates an open interface. In addition, Tsr1 blocks the canonical binding mode of elF5B, required for formation of both 80S and 80S-like ribosomes; and stabilizes 80S-like ribosomes by binding both subunits. (ii) Dim1 is accommodated through its repositioning. In its new position it stabilizes an interaction between H69 from 60S with h24 from pre-40S, thus promoting the stability of the novel 80S-like complex interface. (iii) Because h24 is a component of the platform, the interaction between H69 and h24 opens the platform toward 60S and mobilizes constituent RNAs and proteins. (iv) Platform remodeling allows for temporal regulation of the ATPase Fap7, which links dissociation of Dim1 to quality control (Ghalei et al. 2017). (v) As noted previously (Huang et al. 2020), the structure of the 40S in 80S-like ribosomes resembles the structure of scanning subunits, providing a structural explanation for how formation of 80S-like ribosomes tests scanning competence to ensure the fidelity of translation initiation. Thus, this structure explains how formation of 80S-like ribosomes is required for proofreading of 40S ribosome maturation.

MATERIALS AND METHODS

Yeast strains/cloning

Yeast strains (Supplemental Table S1) were produced using PCR-based recombination (Longtine et al. 1998) and confirmed by PCR and western blotting. Mutations in Dim1, Fap7, Tsr1, Pno1, and Rps14-expressing plasmids were introduced by site-directed mutagenesis, confirmed by sequencing.

Growth assays

Doubling times were measured in a Synergy 2 microplate reader (BioTek Instruments) as described (Collins et al. 2018; Huang et al. 2020). Statistical analyses were performed using Prism v.6.02 (GraphPad Software).

Sucrose density gradient analysis

Sucrose gradient fractionations of whole cell lysates, followed by northern and western blot analysis, were performed as described (Strunk et al. 2012). 40S and 80S fractions were quantified using Quantity One 1-D Analysis Software v.4.1.2 (Bio-Rad Laboratories). Pno1 was quantified using ImageJ Software v1.52 (100) (Rasband, W.S., ImageJ, U. S. National Institutes of Health, https://imagej.nih.gov/ij/, 1997-2018).

Antibodies

Antibodies against recombinant Rps10, Dim1, Nob1, Pno1, Rpo2, and Tsr1 were raised in rabbits (Josman, LLC). Antibodies against Rpl3 or eEF2 were gifts from J. Warner or T. Goss-Kinzy, respectively.

Protein binding

MBP-Fap7(-RYD) and SUMO-Rps14(-RVM) were expressed and purified as described (Ghalei et al. 2017). Protein binding assays were performed as described (Ghalei et al. 2017). An amount of 3 μM MBP-Fap7/MBP-Fap7-RYD was mixed with 4.4 μM SUMO-Rps14/SUMO-Rps14-RVM in binding buffer (BB, 50 mM Tris pH 7.5/150 mM NaCl/5% glycerol), applied to amylose resin, washed, and bound proteins eluted in BB + 50 mM maltose.

Ribosome binding assay

80S-like ribosomes were affinity-purified from Tsr1-TAP; Gal::Fap7 cells grown in YPD medium for 16 h as described (Ghalei et al. 2017). An amount of 20 nM 80S-like ribosomes were incubated with 70 nM purified, recombinant Fap7 or Fap7+SUMO-Rps14 complex in 50 μL of buffer (30 mM HEPES-KOH [pH 6.8], 100 mM NaCl, 6 mM MgCl2, 0.5 mM EDTA, and 1 mM DTT). ATP or AMPPNP was added to a final concentration of 0.5 mM. The samples were incubated on ice for 15 min, placed on 400 μL of a 20% sucrose cushion, and centrifuged for 2 h at 400,000g in a TLA 100.1 rotor. The supernatant was removed, pellets resuspended in SDS loading dye and analyzed by western blotting.

Sample purification and Cryo-EM preparation

Fap7 was depleted by growth in YPD for 16 h. An amount of 2.5 mL of lysis buffer (30 mM HEPES-KOH pH 6.8, 100 mM NaCl, 6 mM MgCl2, RNasin, PMSF, Benzamidine, EDTA—free protease
inhibitor tablet (Santa Cruz Biotechnology), Leupeptin, Pepstatin, and Aprotinin) was added to 4 g of lysed frozen cell powder. The cell powder was mixed with 2 mL of Zircona silica beads (Millipore-Sigma) and vortexed 20 sec for homogenization. The frozen cell lysate thawed at 4°C on a rocker. Thawed cell lysate was cleared via two centrifugations at 4°C: first, at 3000 g for five min and second, at 10,000 g for 15 min. The cleared supernatant was incubated with 250 µL of pre-equilibrated IgG beads (GE Healthcare) at 4°C for 1.5 h with gentle rotation. After incubation, the flow through was discarded and beads were washed three times with buffer A (30 mM HEPES-KOH pH 6.8, 100 mM NaCl, 6 mM MgCl₂, 0.075% NP-40, PMSF, Benzamidine) followed by an additional wash buffer B (30 mM HEPES-KOH pH 6.8, 100 mM NaCl, 6 mM MgCl₂, PMSF, Benzamidine). Washed beads were incubated with 250 µL TEV cleavage buffer (30 mM HEPES-KOH pH 6.8, 100 mM NaCl, 6 mM MgCl₂, PMSF, Benzamidine, 1 mM DTT and 0.5 mM EDTA) supplemented with 2.5 µL AcTEV protease (Invitrogen) at 16°C for 2 h with gentle shaking. After incubation, flow through was collected. The concentration and quality of eluate was determined spectrophotometrically using a Nanodrop 1000 (Thermo Scientific). Analysis of mature and 80S-like ribosomes were independently automodeled with a mask on pre-40S (Supplemental Fig. S3I). The platform density follows the contours of an unchanged Rps11•Rps14•Pno1 complex, supported by mutagenesis.

Modeling

60S and pre-40S were modeled from 80S ribosomes (PDB 3J77 [Svidritskiy et al. 2014]) using Chimera (Pettersen et al. 2004) to rigid body fit each subunit followed by manual adjustment in Coot (Emsley et al. 2010). Rpl41, h23, h45, Rps1, Rps26, and Rps14 were removed with adjustments to Rpl19, H69, and the L1 stalk. In pre-40S, h24 was moved as a rigid body to match the contours of the remodeled bridge. Additionally, the head RNA and proteins were positioned into that density. Tsr1 from PDB 6FAI (Scaloia et al. 2018) was fit as a rigid body. Yeast Dim1 was modeled in SwissModel (Waterhouse et al. 2018) and, after rigid-body fitting into the bilobed density, α-helices and β-sheets were manually placed, followed by refinement with “Phenix.real.space.refine” (Adams et al. 2010). The model matched the density with a correlation coefficient (CC) of 0.8 at 7 Å-resolution. Rps11•Rps14•Pno1 was fit into the segmented platform density, matching the density with a CC of 0.6 at 7 Å-resolution. Each model was independently refined with “Phenix.real.space.refine” (Adams et al. 2010). Pre-40S had 89.7% residues in favored Ramachandran regions and 0.2% in outlier regions (13% Clashscore). 60S had 85.9% residues in favored Ramachandran regions and 0.2% in outlier regions (27.9% Clashscore).

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

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Author contributions: J.R. and H.G. prepared samples; J.R. and M.E.S. carried out structure reconstruction; M.D.P., H.H., H.G., and S.C. carried out yeast analyses; M.C.J. performed cryo-EM analysis of pre-40S; K.K. and M.E.S. conceived of the experiments and wrote the paper.

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