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Structural insights into initial and intermediate steps of the ribosome-recycling process

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
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Supplementary Information includes 10 figures, 2 tables, and 3 movies.

Abstract

The ribosome recycling factor (RRF) and elongation factor G (EF-G) disassemble the 70S post-termination complex (PoTC) into mRNA, tRNA, and two ribosomal subunits. We have determined cryo-EM structures of the PoTC•RRF complex, with and without EF-G. We find that domain II of RRF initially interacts with universally conserved residues of the 23S rRNA helices 43 and 95, and protein L11 within the 50S ribosomal subunit. Upon EF-G binding, both RRF and tRNA are driven toward the tRNA-exit (E) site, with a large rotational movement of domain II of RRF toward the 30S ribosomal subunit. During this intermediate step of the recycling process, domain II of RRF and domain IV of EF-G adapt unusual conformations. Furthermore, binding of EF-G to the PoTC•RRF complex reverts the ribosome from ratcheted to unratcheted state. These results suggest that (i) the ribosomal intersubunit reorganizations upon RRF binding and subsequent EF-G binding could be instrumental in destabilizing the PoTC, and (ii) the modes of action of EF-G during tRNA translocation and ribosome recycling steps are markedly different.

Keywords: Ribosome recycling factor (RRF), elongation factor-G (EF-G), RRF-EFG interactions, conformation of EF-G during ribosome recycling, reverse ratcheting of the ribosome

Introduction

After the termination step of translation, deacylated tRNA and mRNA remain bound to the bacterial 70S ribosome. This complex is called the post-termination complex (PoTC), which must be disassembled so that each of its components can participate in the next round of protein synthesis. In bacteria, disassembly of PoTC is carried out by the reaction that requires the binding of the ribosome recycling factor (RRF) to PoTC, subsequent binding of the elongation factor G (EF-G) in complex with guanosine 5'-triphosphate (GTP), followed by GTP hydrolysis (Hirokawa *et al*, 2006). This reaction drives the dissociation of two ribosomal subunits (Karimi *et al*, 1999; Zavialov *et al*, 2005; Peske *et al*, 2005; Hirokawa *et al*, 2005) and release of mRNA and deacylated tRNA (Hirokawa *et al*, 2005). The initiation factor 3 (IF3) has also been implicated (Seshadri and Varshney, 2006) to play a role in the splitting of the 70S ribosome or in keeping the split subunits separated (Hirokawa *et al*, 2005). IF3 was also suggested to be involved in the release of deacylated tRNA from the 30S subunit (Karimi *et al*, 1999; Peske *et al*, 2005) (but see Hirokawa *et al*, 2006).

Atomic structures of RRF from several bacterial species (Selmer *et al*, 1999; Kim *et al*, 2000; Toyoda *et al*, 2000; Nakano *et al*, 2003; Saikrishnan *et al*, 2005; Yoshida *et al*, 2001) are known. Each of these structures shows two distinct domains: domain I has a three-helix bundle, while domain II is a three-layer $\beta/\alpha/\beta$ sandwich, which is connected to domain I through highly flexible linkers. Relative positions of the two domains among these atomic structures are not the same. Therefore, an inherent inter-domain flexibility of RRF has been proposed to be important for its function (Selmer *et al*, 1999; Yoshida *et al*, 2001). The binding position of RRF on the ribosome has been studied by using a variety of biochemical and structural biology techniques, such as the hydroxyl radical probing (HRP) (Lancaster *et al*, 2002), cryo-electron microscopy (cryo-EM) (Agrawal *et al*, 2004; Gao *et al*, 2005; Barat *et al*, 2007), and X-ray crystallography (Wilson *et al*, 2005; Borovinskaya *et al*, 2007; Weixlbaumer *et al*, 2007; Pai *et al*, 2008; Dunkle *et al*, 2011). These studies have revealed a predominant binding position of RRF on the ribosome, henceforth referred to as position 1 (or P1). Another weak and transient binding

position, referred to as position 2 (or P2), was captured on a small fraction of the dissociated large (50S) ribosomal subunits in one of the cryo-EM studies (Barat *et al*, 2007). In its position 1, domain I of RRF resides on the 50S ribosomal subunit, stretching over the ribosomal aminoacyl (A)- and peptidyl (P)-tRNA binding sites (Hirokawa *et al*, 2002; Lancaster *et al*, 2002; Agrawal *et al*, 2004), while domain II of RRF flexibly occupies the inter-subunit space between ribosomal protein S12 of the small (30S) ribosomal subunit and the L11 stalk-base region of the 50S ribosomal subunit (Agrawal *et al*, 2004; Gao *et al*, 2005; Barat *et al*, 2007; Borovinskaya *et al*, 2007; Weixlbaumer *et al*, 2007; Pai *et al*, 2008; Dunkle *et al*, 2011).

In order to describe the binding positions of RRF on the ribosome in conjunction with inter-domain flexibility of the two RRF domains, we introduced a nomenclature in our previous study (Barat *et al*, 2007). Accordingly, RRF binds to the ribosome in position 1, or position P1/IIa, where “P1” and “IIa” refer to binding position of domain I and relative orientation of domain II, respectively. Subsequently, domain II of RRF adapts a different orientation, referred to as position P1/IIb, suggesting that the domain I of RRF stayed in the same position with an altered relative orientation of domain II. In all previous structural studies of the RRF-bound ribosome complexes, domain II adapts slightly different versions of P1/IIb on the ribosome. In position 2, or P2/IIb, RRF is situated exclusively in the P-site region of the 50S ribosomal subunit such that domain II and the inter-domain elbow region of the bound RRF would make significant steric clash with the 30S ribosomal subunit’s neck and head regions. This implies that RRF could achieve this position during, or immediately after, the dissociation of the two ribosomal subunits (Barat *et al*, 2007). Cryo-EM structures of the 50S•RRF•EF-G•GDPNP complex have also been studied, using the pre-dissociated 50S ribosomal subunit for the formation of the complex (Gao *et al*, 2005; Gao *et al*, 2007). However, capturing both RRF and EF-G simultaneously on the PoTC, or on the vacant 70S ribosome, for structural studies has been technically challenging, due to rapid disassembly of the PoTC complex in the presence of both RRF and EF-G.

To understand the molecular details of the ribosome recycling process, it is essential to study the interaction between RRF and EF-G on the PoTC. In order to capture the structure of the PoTC•RRF•EF-G complex, together with mRNA and tRNA, we exploited the fact that such a complex can be stabilized by using RRF and EF-G from two different bacterial species, i.e., when *Thermus thermophilus* RRF (ttRRF) and *E. coli* EF-G (henceforth referred to as EF-G) in conjunction with *E. coli* PoTC (PoTC) (Raj *et al*, 2005) are used. The ttRRF•EF-G combination slows down the recycling reaction (Raj *et al*, 2005). However, increasing the concentration of both factors results in rapid disassembly of the PoTC (N.I., T.Y., A.K., R.K.A., and H.K., in preparation). Furthermore, the same species combination of the two factors, from either *T. thermophilus* or *E. coli*, readily disassembles the PoTC (Raj *et al*, 2005). These results suggest that the mode and site of binding of ttRRF on the PoTC are the same as that of an ecRRF and that the two-species combination is biochemically active. We prepared the PoTC•ttRRF complex and then incubated it with EF-G•GTP in the presence of fusidic acid (FA) to obtain the PoTC•ttRRF•EF-G•GDP•FA complex. FA stabilizes EF-G on the 70S ribosome in a post-GTP hydrolysis state (Agrawal *et al*, 1998; Valle *et al*, 2003; Datta *et al*, 2005; Gao *et al*, 2009; Ratje *et al*, 2010), and its antimicrobial activity is primarily due to inhibition of the ribosome recycling step (Savelsbergh *et al*, 2009). The PoTC•ttRRF and the PoTC•ttRRF•EF-G•GDP•FA complexes were analyzed by three-dimensional (3D) cryo-EM technique (Frank *et al*, 2000). The cryo-EM map of the PoTC•ttRRF•EF-G•GDP•FA complex shows densities for both RRF and EF-G on the PoTC and reveals an intermediate binding position of RRF that lies between previously described binding positions P1 and P2. Molecular analysis of the complex revealed interactions between domain II of RRF and domains III-V of EF-G on the PoTC. In addition, both RRF and EF-G adapt hitherto unknown configurations on the ribosome. These findings provide important insights into the mechanism of ribosome recycling, and suggest that the mode of action of EF-G in the ribosome recycling step is different from that in the tRNA translocation step.

Results

Structures of the PoTC•ttRRF complex and the PoTC•ttRRF•EF-G•GDP•fusidic acid complex

We isolated the polysome from growing *E. coli* cells and removed the peptidyl moiety, by treating with puromycin, to obtain the model post-termination complex (PoTC). As described in the introduction section, to capture both RRF and EF-G simultaneously on the PoTC, we used the combination of ttRRF, EF-G and PoTC. In order to establish that the binding position of ttRRF on PoTC is identical to the binding position of ecRRF, we first obtained a cryo-EM map of the PoTC•ttRRF complex (henceforth referred to as *Complex 1*). The map shows distinct mass of densities corresponding to ttRRF, a tRNA at the P/E hybrid site (Moazed and Noller, 1989; Agrawal *et al*, 1999b) and the mRNA (**Figure 1A-C**; and Supplementary Figure S1A for a stereo viewing), when compared with the structure of a vacant 70S ribosome (Schuwirth *et al*, 2005). The overall location of the density corresponding to domain I of ttRRF appears to be similar to those derived in previous studies (Lancaster *et al*, 2002; Agrawal *et al*, 2004; Gao *et al*, 2005; Barat *et al*, 2007; Borovinskaya *et al*, 2007; Weixlbaumer *et al*, 2007; Pai *et al*, 2008; Dunkle *et al*, 2011). However, domain II of RRF is predominantly oriented towards the 50S ribosomal subunit, making contacts with the stalk-base (Sb) region (**Figure 1A, B**). This orientation of domain II of RRF on the 70S ribosome was first inferred from a hydroxyl radical probing (HRP) study (Lancaster *et al*, 2002) and subsequently observed in our previous cryo-EM studies (Agrawal *et al*, 2004; Barat *et al*, 2007), and was referred to as position P1/IIa (Barat *et al*, 2007). However, position P1/IIa has eluded all X-ray crystallographic studies (Borovinskaya *et al*, 2007; Weixlbaumer *et al*, 2007; Pai *et al*, 2008; Dunkle *et al*, 2011). These observations suggest that RRF in position P1/IIa is attained only in physiologically relevant solution conditions, such as HRP (Lancaster *et al*, 2002) and cryo-EM (Agrawal *et al*, 2004; Barat *et al*, 2007) conditions but not in crystallographic conditions, as all X-ray crystallographic

structures of the 70S•RRF complexes invariably find RRF in the position P1/IIb (Borovinskaya *et al.*, 2007; Weixlbaumer *et al.*, 2007; Pai *et al.*, 2008; Dunkle *et al.*, 2011) with domain II of RRF oriented towards the 30S ribosomal subunit protein S12. In the complexes formed between the 70S ribosome and RRF in our previous cryo-EM studies, density for domain II was split between positions P1/IIa and P1/IIb (Agrawal *et al.*, 2004; Barat *et al.*, 2007). Since the inter-species combination of PoTC and ttRRF, allows us to capture the RRF predominantly in position P1/IIa, density corresponding to domain II is the strongest among all ribosome•RRF structures determined thus far. It is therefore possible to accurately dock all atom coordinates of domain II of ttRRF (Toyoda *et al.*, 2000) into the cryo-EM density (Supplementary Figure S2A). A very weak density is also observed in position P1/IIb, suggesting that in a small fraction (~10%) of the total population of Complex 1 domain II of ttRRF is oriented toward the 30S ribosomal subunit (Supplementary Figure S2B, C). Since the binding position of domain I of ttRRF closely matches with that in *E. coli* 70S•RRF complexes (Agrawal *et al.*, 2004; Barat *et al.*, 2007; Pai *et al.*, 2008) and the fact that ttRRF is capable of disassembling the PoTC with ttEF-G (Raj *et al.*, 2005), the predominant P1/IIa position attained by the domain II of ttRRF on the PoTC must represent a functionally relevant binding state.

By incubating Complex 1 with EF-G, GTP and FA, we obtained the PoTC•ttRRF•EF-G•GDP•FA complex (henceforth referred to as *Complex 2*). The computed cryo-EM map of Complex 2 clearly shows distinct mass of densities for RRF, EF-G, tRNA and mRNA (**Figure 1D–F**; see also Supplementary Figure S1B for stereo viewing), when compared with the map of a vacant 70S ribosome (Schuwirth *et al.*, 2005). Both domains of RRF (Selmer *et al.*, 1999), all five domains of EF-G (Ævarsson *et al.*, 1994; Czworkowski *et al.*, 1994), and a tRNA can be readily identified in this map. However, densities of RRF and EF-G appear to be partially fused, suggesting a direct interaction between these two protein factors.

A comparison between the maps of Complex 1 and Complex 2 reveals a striking movement of RRF toward the tRNA-exit (E) site (or towards L1 stalk) upon EF-G binding (**Figure 1**; also described later), suggesting that a direct interaction between ttRRF and EF-G

shifts the ttRRF from position P1/IIa (and position P1/IIb). We designate this new binding position of RRF as an *intermediate* position (or position Pi). During this shift, domain II of ttRRF severs its interaction with the L11 stalk-base (Sb) region of the 50S ribosomal subunit and is rotated toward the 30S ribosomal subunit. The same stalk-base region makes contact with domain V of EF-G in Complex 2 (**Figure 1**; compare panels D and E with panels A and B, respectively). In addition, we observe a strong mass of well-defined density corresponding to the C-terminal domain (CTD) of protein L7/L12 (**Figure 1D, E**) that forms a bridge (or the previously described “arc-like connection” (Agrawal *et al*, 1998; Datta *et al*, 2005)) between EF-G and the stalk-base region of the 50S subunit. The map of Complex 2 contains densities for both the mRNA and a tRNA (**Figure 1D-F**; also see Supplementary Figures S1 and S3). However, as compared to Complex 1, the density corresponding to the anticodon end of the tRNA is significantly broadened (**Figure 1**, compare panel E with panel B) and partially bifurcated between P and E sites, suggesting that the codon-anticodon interaction is affected significantly upon EF-G binding (also described later).

Interactions between the PoTC and ttRRF in Complex 1

In order to determine molecular interactions between the PoTC and ttRRF, we docked the atomic coordinates of the ribosome (Schuwirth *et al*, 2005; Gao *et al*, 2009) and ttRRF (Toyoda *et al*, 2000) into corresponding densities within the cryo-EM map, using the MDFF program (Trabuco *et al*, 2008; Trabuco *et al*, 2009). This fitting reveals that domain I of ttRRF is situated within the cavity formed by helices 69, 71, 80 and 93 of the 23S rRNA (**Figure 2A**; see also Supplementary Movie S1), while its domain II interacts with the stalk-base of the 50S ribosomal subunit, involving helix 43 of the 23S rRNA and the N-terminal domain (NTD) of the ribosomal protein L11. Two amino acid residues (Tyr46 and Gln54), located at the tip of domain II of RRF, appear to make direct contacts with the ribosome components. Tyr46 (*E. coli* # Tyr45 [or ecTyr45]; See Supplementary Figure S4 for aligned sequences) is in close vicinity to Ser20 and Pro21 of protein L11, while Gln54 (ecGln53) would make a protein-RNA contact

with the highly conserved nucleotide residue A1067 within helix 43 of the 23S rRNA (**Figure 2A**). Arg110 (ecArg109) in the elbow region of ttRRF lies in close proximity to amino-acid residue His76 of protein S12. Tyr121 (ecGlu120) of domain I of ttRRF would contact U1915 of helix 69 (H69) of the 23S rRNA. In addition, Asp97 (ecSer96) of domain II of ttRRF lies in close proximity to two other conserved nucleotides, A2660 and G2661, within the α -sarcin-ricin stem-loop (H95) of the 23S rRNA (**Figure 2B**). Arg119 (ecArg118) lies close to C1947 and G1948 of helix71 (H71) of the 23S rRNA. Thus, while the interactions between domain I of RRF and the ribosome are essentially the same as derived in previous X-ray crystallographic studies (Borovinskaya *et al*, 2007; Weixlbaumer *et al*, 2007; Pai *et al*, 2008; Dunkle *et al*, 2011), interactions between domain II and the ribosome are found to be dramatically different from what has been reported in those studies. This is due to the fact that we captured the ttRRF predominantly in P1/IIa configuration, while P1/IIb is the preferred configuration in crystallographic studies. It is likely that the P1/IIa configuration precedes the P1/IIb configuration on the PoTC (see Discussion).

Interactions among the ribosome, ttRRF and EF-G in Complex 2

Similarly to Complex 1, we flexibly docked the atomic structures of ribosome (Schuwirth *et al*, 2005; Gao *et al*, 2009), ttRRF (Toyoda *et al*, 2000), and a homology model of the *E. coli* EF-G, which was built using the structural templates of ttEF-G (Ævarsson *et al*, 1994; Czworkowski *et al*, 1994), into corresponding densities within the cryo-EM map of Complex 2 (**Figure 2C**; Supplementary Figure S2D for a stereo viewing, and Supplementary Movie S2). Based on these fitting results, we examined the interactions among ttRRF, EF-G, and the ribosomal components. Similarly to Complex 1, domain I of ttRRF is situated within the cavity formed by helices 69, 71, 80 and 93 of 23S rRNA (**Figure 2C**). However, upon EF-G binding specific contacts between RRF domains and the ribosome change significantly (Supplementary Table S1A).

Several amino acid residues within domains III, IV and V of EF-G interact with domain II of RRF (**Figure 2D**; Supplementary Table S1B). Lys484 on domain III of EF-G appears to contact Ser93 (ecAsn92) of ttRRF. Lys608 and Leu611 of EF-G domain IV, near the linker region between domains IV and V, would contact Tyr46 (ecTyr45) and Asn74 (ecSer73) of ttRRF, respectively. Glu661 and Arg638 on domain V of EF-G is in close vicinity to amino acid residues Gly96 (ecGly95) and Gln73 (ecArg72) of ttRRF. Domain II of ttRRF also contacts H69 of the 23S rRNA. Thus, domain II of RRF is somewhat sandwiched between EF-G and the ribosome components. In addition, Tyr121 (ecGlu120) in domain I of RRF appears to switch partner from H69 nucleotide U1915 in position P1 to A1916 in position Pi (Supplementary Table S1A).

As pointed out earlier, there is a marked shift in overall position of ttRRF (**Figure 3A**), as can be inferred by comparing the densities corresponding to the elbow region of ttRRF, before and after EF-G binding (**Figure 3B**). Domain I of ttRRF is shifted toward the E site. Flexible fitting shows that domain I of ttRRF is shifted by $\sim 8 \text{ \AA}$ such that the long axis of domain I of ttRRF is not perfectly parallel to that in position P1 (**Figure 3C**). Furthermore, the relative orientation of domain II of RRF is different from previously described IIa and IIb configurations (Barat *et al*, 2007) (Supplementary Figure S5). Therefore, this intermediate RRF binding position is referred to as position Pi/IIIi, where IIIi refers to a unique relative orientation of domain II of ttRRF (Supplementary Figures S5 and S6). Thus, in addition to pushing the RRF toward the E site, the binding of EF-G to Complex 1 also induces a rotational shift in domain II of ttRRF, from the 50S-subunit side to the 30S-subunit side (curved arrows in **Figure 3**).

Because of a pre-bound ttRRF, ecEF-G is unable to attain its previously known binding position on the ribosome in the 70S•EF-G•GDP•FA complex (Agrawal *et al*, 1998; Datta *et al*, 2005; Gao *et al*, 2009; Ratje *et al*, 2010). Domains IV and V of EF-G are situated more toward the shoulder of the 30S subunit within the intersubunit space, when compared with their positions in the ec70S•ecEF-G•GDP•FA complex (Datta *et al*, 2005) (**Figure 4**). Domain IV of

EF-G adapts a skewed conformation, where both its tip and the proximal end, i.e., the end continuing with domains III and V, are shifted by 8-9 Å towards the shoulder of the 30S ribosomal subunit (**Figure 4C**). However, there are no appreciable changes of domain I and II of EF-G as compared to what has been observed during EF-G-dependent tRNA translocation on the ribosome. Nevertheless, these observations suggest that EF-G can adopt more than one conformational state on the ribosome in post-GTP hydrolysis state and that the mechanism of interaction of EF-G with the RRF-bound PoTC is significantly different from that with the pre-translocational ribosome complex.

Dynamic movements of domain II of RRF

Domain II of RRF has been suggested to be the main functional component of the RRF structure (Guo *et al*, 2006). We have indentified three configurations of domain II on the ribosome, namely IIa, IIb, and Iii. The binding positions of domain I of RRF on the ribosome are essentially the same in P1/IIa and P1/IIb configurations. It is possible that the domain II exists in a dynamic equilibrium between these two configurations. We find that specific ribosomal components switch their binding partners from domain II of RRF (in both positions P1/IIa and P1/IIb) to specific domains of the subsequently bound EF-G. The cleft region between tip of the 23S rRNA helix 43 and NTD of protein L11 in the stalk-base region of the 50S subunit, previously occupied by domain II of ttRRF in position P1/IIa within Complex 1 (**Figure 5A**), is subsequently occupied by domain V of EF-G in Complex 2 (**Figure 5B**). Similarly, the interaction between protein S12 of the 30S ribosomal subunit and domain II of RRF in position P1/IIb (Agrawal *et al*, 2004; Gao *et al*, 2005; Barat *et al*, 2007; Borovinskaya *et al*, 2007; Weixlbaumer *et al*, 2007; Pai *et al*, 2008; Dunkle *et al*, 2011) would be severed upon EF-G binding. The domain II of RRF in position Pi/Iii has moved away from protein S12. The space previously occupied by domain II of RRF in its P1/IIb position (**Figure 5C**) is partially occupied by domain III of ecEF-G in Complex 2 (**Figure 5D**). Furthermore, the interactions between the tip of the 23S rRNA helix 95 (the SRL) and domain II of RRF in position P1/IIa

(**Figure 2B**) would be disrupted upon its transition to P1/IIb as well as upon EF-G binding. These results clearly show that RRF has moved toward the E site upon EF-G binding.

Conformational changes of ribosomal components that interact with RRF and EF-G

The ribosomal components that interact with domain II of RRF also undergo large conformational changes during the transition from P1/IIa (or P1/IIb) to Pi/IIi. For example, H43 in the stalk-base region moves towards the solvent side by ~ 5 Å upon binding of ttRRF alone (**Figure 5A**). Upon subsequent binding of EF-G, the stalk-base shifts back toward the inter-subunit space by ~ 5 Å and the CTD of protein L7/L12 forms a bridge between the NTD of L11 and the G' domain of EF-G (**Figure 5B**), corroborating our previous study (Datta *et al*, 2005) and indicating that the CTD of L7/L12 adapts this relatively stable position on the ribosome in a post-GTP hydrolysis state, both during the translocation and recycling steps. In addition, we observe a distinct movement in density corresponding to protein S12 on the inter-subunit face of the 30S-subunit body. Upon ttRRF binding (both in P1/IIa and P1/IIb configurations) S12 moves towards the 16S rRNA helix 44 (**Figure 5C**), and reverts back, close to its original position, upon subsequent binding of EF-G (**Figure 5D**).

Upon RRF binding to the PoTC, helix 69 (H69) of the 23S rRNA undergoes a conformational change. The tip of H69 is shifted by ~ 4 Å toward the rest of the 50S ribosomal subunit, apparently severing its contact with helix 44 of the 16S rRNA (**Figure 5C**) to disrupt the bridge B2a (Yusupov *et al*, 2001). This movement in H69 is consistent with some of the previous cryo-EM (Agrawal *et al*, 2004; Barat *et al*, 2007) and X-ray crystallographic (Borovinskaya *et al*, 2007; Pai *et al*, 2008) studies; however the overall movement is less pronounced than what was reported in those X-ray studies. The altered position of tip of H69 remains unchanged between P1/IIa and P1/IIb states and upon subsequent binding of ecEF-G to Complex 1 (**Figure 5D**). In other words, the position of H69 is found to be the same in both complexes 1 and 2. Except for the helix 69, no other rRNA helices that are known to interact

with domain I of RRF (e.g., H71, H80 and H93) show any noticeable movement in either of our complexes, when compared with their respective positions in the map of control PoTC.

Global conformational changes of the ribosome and tRNA movement

In addition to the conformational changes described in the previous section, there are global conformational changes of the ribosome in both complexes 1 and 2 (Supplementary Figure S7, Movie S3). This includes a marked movement of the L1 stalk due to RRF binding and upon subsequent binding of EF-G (**Figure 6A, B**). While RRF binding moves the L1 stalk toward the intersubunit space, the subsequent binding of EF-G moves it in the opposite direction, i.e, towards the solvent side. The deacylated tRNA, which is exclusively situated in the P/E state in Complex 1 (**Figure 6A**), is present in both P/E and E/E states in Complex 2 (**Figure 6B**; also see Supplementary Figure S8), suggesting that the anticodon end of the tRNA has been pulled into the E site in a significant population of Complex 2, along with outward movement of the L1 stalk upon EF-G binding. In addition, the ribosomal components corresponding to intersubunit bridges B1a-c and B3 undergo large conformational changes upon ttRRF binding (Complex 1), corroborating our previous study (Barat *et al*, 2007), and upon EF-G binding (Complex 2) (Supplementary Figure S7). Furthermore, the 70S ribosome in Complex 1 is in the ratcheted state (Supplementary Figure S7), where the 30S subunit is found rotated in an anticlockwise direction relative to the 50S subunit (Agrawal *et al*, 1999a; Frank and Agrawal, 2000). This observation is consistent with previous studies (Gao *et al*, 2005; Barat *et al*, 2007). However, in Complex 2, the 30S subunit is not rotated in an anticlockwise direction relative to the 50S subunit. In other words, subsequent binding of EF-G to Complex 1 brings back the 30S subunit to its unratcheted state (**Figure 6C, D**; Supplementary Movie S3), which is unusual for the EF-G-bound 70S ribosome. It should be noted that the movement of the 30S subunit head is in a direction opposite to the earlier described L1-stalk movement that takes place upon binding of EF-G (indicated by arrows in panels B and D of **Figure 6**). This observation also implies that

modes of action of EF-G are markedly different between tRNA translocation and ribosome recycling steps of protein synthesis.

Discussion

Capturing both RRF and EF-G simultaneously on the PoTC is essential for understanding the mechanism of the ribosome recycling step. Since the mRNA and deacylated tRNA remain bound to the ribosome (**Figure 1D-F**) and the ttRRF has advanced (**Figure 3**) toward the tRNA-exit (E) site, the PoTC•ttRRF•EF-G•GDP•FA complex (Complex 2) represents an intermediate state of the ribosome recycling that is stalled by FA immediately after EF-G dependent GTP hydrolysis. The Pi/III position of ttRRF, captured in this study in conjunction with eEF-G, is significantly different from both previously identified positions P1/IIa and P1/IIb. In position Pi/III, the tip of domain 1 of ttRRF is shifted slightly upward (**Figure 3B, C**) such that it is situated between the tip of domain I at positions P1 and the previously described short-lived position P2 (Barat *et al*, 2007), indicating the path of RRF during its transition from positions P1 to P2 on the ribosome. A comparison of mRNA densities between the PoTC•ttRRF complex (Complex 1) and Complex 2, along entire mRNA path (Yusupova *et al*, 2006; Supplementary Figure S3), shows that the contact points between mRNA and the ribosome remain largely unaffected at this stage. However, there is a marked shift in the tRNA's anticodon from the P site to the E site in a subpopulation of Complex 2 (**Figure 6B**; Supplementary Figure S8). This shift in the tRNA's anticodon appears to be coupled to the movement of the L1 stalk towards the solvent side, which could facilitate the ultimate tRNA release from the PoTC. Since the codon-anticodon interaction is expected to be relatively weak at the E site (Jenner *et al*, 2007), it is possible that the mRNA and tRNA releases from the PoTC are triggered during this intermediate state.

The inter-subunit ratchet-like rotation (Agrawal *et al*, 1999a; Frank and Agrawal, 2000) is essential for the tRNA translocation (Horan and Noller, 2007). Normally, during EF-G-dependent tRNA translocation, EF-G binding stabilizes the ribosome in ratcheted state (Agrawal *et al*, 1999a; Frank and Agrawal, 2000). In sharp contrast, we find that EF-G binding to the Complex 1 reverts the ribosome to its unratcheted state. It is possible that during its initial binding to the Complex 1 the tip of domain IV of EF-G

instantaneously pushes the 30S subunit head toward the E site, facilitating the movement of tRNA anticodon from P to E site. However, due to the presence of ttRRF, the rest of domain IV, along with the insertion domain V, of EF-G is not able to occupy its usual binding position on the ribosome (**Figure 4**). This steric constraint would force the tip of domain IV to retract back, and in the process, pull back the 30S subunit head to an unratcheted state (**Figure 6C, D**). Furthermore, there are marked differences in contact points of EF-G with tRNA and RRF between EF-G-dependent tRNA translocation and recycling processes. During tRNA translocation, the tip of EF-G interacts with the anticodon of tRNA (Gao *et al*, 2009), whereas during recycling, there is no direct interaction between EF-G and tRNA, instead, the junction between domains III-V of EF-G interacts with domain II of RRF (**Figure 2C, D**). Thus, modes of EF-G action are strikingly different during tRNA translocation and recycling steps of protein synthesis. In fact, in mammalian mitochondria, a separate isoform of EF-G is recruited exclusively for the recycling step (Tsuboi *et al*, 2009). Similarly, existence of two separate isoforms of EF-G, one each for the tRNA translocation and recycling steps, has also been reported in a bacterial species (Suematsu *et al*, 2010).

Previous cryo-EM (Agrawal *et al*, 2004; Barat *et al*, 2007) and X-ray crystallographic studies (Borovinskaya *et al*, 2007; Pai *et al*, 2008) have shown a large movement in the 23S rRNA helix 69 (H69) upon RRF binding to the 70S ribosome. The tip of H69 has been found to be moving away from the 16S rRNA helix 44 (h44) of the 30S subunit, implying disruption of an intersubunit bridge B2a to facilitate the subunit dissociation during the recycling process. These findings corroborate previous studies showing that the rate of ribosomal subunits association is directly affected by deletion (Ali *et al*, 2006) or mutations (Hirabayashi *et al*, 2006) of H69. In addition, several conformational changes associated with components of other intersubunit bridges, such as B1a-c, and B3 were also noted (Barat *et al*, 2007). We observe similar conformational changes involving bridges B1a-c and B2a in the maps of both complexes 1 and 2 (Supplementary Figure S7). These changes might contribute to the dissociation of ribosomal subunits during the recycling steps. Interestingly, one of the X-ray crystallographic studies of the 70S•RRF complex (Weixlbaumer *et al*, 2007) such conformational changes, including those previously described for H69, were not reported. In that study, an anticodon stem-loop (ASL) was pre-bound in the ribosomal P site. Since an intact tRNA molecule, including the CCA end, is required for tRNA to

adapt the P/E state (Joseph and Noller, 1998), it is likely that ASL stabilized the bridge B2a in that 70S•ASL•RRF complex by maintaining a P/P-like state.

Previous cryo-EM studies of a complex of RRF and EF-G with the pre-dissociated 50S subunit (Gao *et al.*, 2005; Gao *et al.*, 2007) placed the highly flexible domain II of RRF in a position that is firmly occupied by the 16S rRNA helix 44 of the 30S subunit in its natural substrate, PoTC. Since a pre-dissociated 50S subunit is not a substrate for ribosome recycling, and since RRF bound to the pre-dissociated 50S subunit cannot be removed by EF-G while RRF bound to the 70S ribosome is removed by EF-G (Kiel *et al.*, 2003), the binding of RRF and EF-G to pre-dissociated 50S subunit is less likely to represent an actual step of the process of recycling.

Based on the results presented here and previous studies (Agrawal *et al.*, 2004; Barat *et al.*, 2007), we outline the possible sequence of events of ribosome recycling (**Figure 7**). The PoTC carries mRNA and a deacylated tRNA, which fluctuates between P/P and P/E states. This fluctuation is coupled to the unratcheted and ratcheted states, respectively, of the ribosome (**Figure 7A**; also see Dunkle *et al.*, 2011). Upon RRF binding, the PoTC is locked in a ratcheted state, accompanied with distortion (or partial destabilization) of an intersubunit bridge, B2a. Since the ratcheted state of ribosome is observed irrespective of the relative orientation of domain II (Gao *et al.*, 2005; Barat *et al.*, 2007; Dunkle *et al.*, 2011), interaction with domain I must be responsible for locking the ribosome in the ratcheted state. It is likely that RRF binds the PoTC in P1/IIa configuration, as domain II in this case is extended more to the solvent side with interaction with the functionally important and flexible stalk-base region of the 50S subunit (**Figure 7B**). This initial interaction may trigger a movement of RRF's flexible domain II to the P1/IIb configuration such that tip of domain is reoriented into the intersubunit space and toward protein S12 of the 30S subunit (**Figure 7C**), as also proposed in our previous study using ecRRF on the ecPoTC (Barat *et al.*, 2007). While the possibility that RRF is in a dynamic equilibrium between P1/IIa and P1/IIb configurations cannot be ruled out and the question that which of the two configurations present the substrate for the subsequent binding of EF-G to Complex 1 remains unresolved, a spontaneous P1/IIa to P1/IIb transition presents a logical sequence of events as position of domain II of RRF in P1/IIb is close to that attained after EF-G binding (Supplementary Figures S5 and S6). Upon EF-G binding, RRF advances to an

intermediate position Pi/III and the anticodon of the deacylated tRNA from P to E site, with mRNA staying firmly bound to the ribosome, while the 30S subunit reverts back to an unratcheted state (**Figure 7D**). Since the reverse ratchet (or the unratcheted state) is not an energetically favored state for the EF-G-bound ribosome (see Munro *et al*, 2009), Complex 2 represents a high-energy state that ultimately leads to disassembly of the PoTC by pushing RRF to previously observed position P2 (Barat *et al*, 2007; **Figure 7E**). As previously suggested (Barat *et al*, 2007), RRF at position P2 has an extremely low affinity for the dissociated 50S subunit, and is spontaneously released after the ribosomal subunit dissociation.

Materials and Methods

Preparation of ribosomal complexes

The *E. coli* polysome, EF-G and *Thermus thermophilus* (tt)RRF were prepared as described previously (Raj *et al*, 2005). The PoTC•ttRRF complex (Complex 1) was obtained by incubating 32 nM polysome, 100 μ M puromycin with 41 μ M ttRRF in the buffer R, containing 50 mM Tris-HCl (pH 7.5), 10 mM Mg(OAc)₂, 25 mM KCl for 20 min at 30 °C. Binding of ttRRF to the PoTC was ascertained by passing a portion of the reaction mixture through NanoSep 300k Omega to remove unbound ttRRF and then the isolated Complex 1 was subjected to quantitative Western blot using anti-ttRRF antibody (Raj *et al*, 2005). The PoTC•ttRRF•EF-G•GDP•FA complex (Complex 2) was prepared by incubating Complex 1 with 4 μ M EF-G and 1.5 mM GTP for 30 min at 30 °C, followed by the addition of 200 μ M fusidic acid (FA) and continued incubation for another 10 min. As described for Complex 1, a portion of the reaction mixture was analyzed by quantitative Western blot for the presence of ttRRF and EF-G. The cryo-EM grids were prepared immediately after the preparation of the complexes.

Cryo-EM electron microscopy and image processing

Cryo-EM grids were prepared using standard protocol (Grassucci *et al*, 2007) with 32 nM ribosomal complexes, and data were collected on a Tecnai F20 field emission gun electron microscope (FEI) at liquid nitrogen temperature, using an Oxford cryo-holder (Oxford Instruments) at 200 kV and under low dose condition. Images were recorded on film (SO-163, Kodak) at magnification calibrated at 51,282x. Micrographs were scanned by a PhotoScan scanner (Z/I imaging) with a step size of 14 μm , which corresponds to 2.78 \AA pixel size on the object scale. For Complex 1, a total of 153,927 images were manually picked from 195 micrographs, covering from -0.4 to -4.3 μm defocus range. After cross-correlation-based screening, 112,425 images were used in the final reconstruction. For Complex 2, a total of 338,823 images were manually picked from 383 micrographs, covering from -0.4 to -4.0 μm defocus range. After cross-correlation-based screening, 245,866 images were used in the final reconstruction. The methods of supervised classification (Valle *et al*, 2002), using vacant ratcheted- and unratcheted-state ribosome maps (Agirrezabala *et al*, 2008) as two references, and maximum-likelihood 3D classification (Scheres *et al*, 2007) were applied to separate possible conformational states within datasets of both Complex 1 and Complex 2. These analyses suggested that particle images used for both complexes were highly homogeneous. However, the possibility of existence of low levels of conformational heterogeneity in both datasets cannot be ruled out. The SPIDER software (Shaikh *et al*, 2008) was used for all image processing, including particle selection, alignment and 3D reconstruction. Projection-matching (Penczek *et al*, 1994) in conjunction with small-angle alignment (up to 0.2° angular spacing) was employed for the iterative refinements. Resolutions of the final cryo-EM maps of Complex 1 and Complex 2 were 11.1 \AA (at 0.5 cutoff of FSC; or 9.4 \AA at 0.143 cutoff of FSC (Rosenthal and Henderson, 2003)) and 9.9 \AA (at 0.5 cutoff of FSC; or 8.4 \AA at 0.143 cutoff of FSC), respectively (Supplementary Figure S9, Table S2).

Molecular Interpretations

For molecular interpretations, atomic structures of the ribosomes (Schuwirth *et al*, 2005) (PDB IDs; 2AVY and 2AW4), ttRRF (Toyoda *et al*, 2000) (PDB ID; 1EH1) and the homology model of ecEF-G, generated using a protein homology modeling server, SWISS-MODEL (Arnold *et al*, 2006), were used. Initially, coordinates of all domains of RRF and EF-G were individually aligned to the corresponding cryo-EM densities. The independently fitted domains were then linked together using LOOPY (Xiang *et al*, 2002). The manually fitted models were further refined by flexible fitting and energy minimization using MDFF (Trabuco *et al*, 2008; Trabuco *et al*, 2009). Both RRF and EF-G models fit extremely well into corresponding cryo-EM densities, as judged by high cross-correlation coefficient (CCC) values of 0.87 for the ttRRF in Complex 1, and 0.89 and 0.94 for the ttRRF and ecEF-G, respectively, in Complex 2. Densities remained after attributing densities to the ribosome, tRNA, RRF, and EF-G were assigned to mRNA, which matches closely with previously determined path of mRNA (Yusupova *et al*, 2006; Schuette *et al*, 2009). The achievable accuracy of fittings of atomic models into cryo-EM maps has been estimated to be 6 to 10 times better than the resolution of the cryo-EM map (Rossmann, 2000), which would translate to an average value of ~ 1.4 Å in our case. Chimera software (Pettersen *et al*, 2004) was used for all visualization work.

Accession Codes. The cryo-EM maps of both complexes 1 and 2 have been deposited in the EM database (<http://emdep.rutgers.edu>) with accession codes EMD-1915 and EMD-1917, respectively. The extracted factor densities from both complexes have also been deposited with accession codes EMD-1916 and EMD-1918, respectively. The fitted coordinates of ttRRF, ecEF-G, and relevant components of the ribosome structure corresponding to Complex 1 and Complex 2 have been deposited in the protein data bank (www.rcsb.org) under PDB ID codes 3J0D and 3J0E, respectively.

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Author Contributions. H.K., A.K., and R.K.A. designed the research; T.Y. prepared the complexes, collected cryo-EM data, performed image processing, flexible docking, and molecular interpretation. T.R.S. helped with image processing. T.Y. and R.K.A. analyzed and interpreted the data. N.I. provided purified tRRF, EF-G and polysomes and performed biochemical assays. T.Y., H.K., A.K., N.I. and R.K.A. wrote the paper.

Conflict of Interest. The authors declare that they have no conflict of interest.

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Figure Legends:

Figure 1. Segmented cryo-EM maps of the PoTC•ttRRF and PoTC•ttRRF•EF-G•GDP•FA complexes. (A-C) The PoTC•ttRRF complex (Complex 1): (A) The 70S ribosome (yellow, 30S subunit; and blue, 50S subunit) is viewed from the tRNA entry side, with strong densities for both domains (I and II) of ttRRF (red) clearly visible in the inter-subunit space (see also Supplementary Figure S1A for a stereo viewing); (B) the same map is shown from the 50S interface side, where the 30S subunit has been computationally removed to reveal densities corresponding to ttRRF (in position P1/IIa) and deacylated tRNA (green, in the P/E site); and (C) the same complex is shown along with density corresponding to mRNA (dark blue) from the 30S interface side, without the 50S subunit. (D-F) The PoTC•ttRRF•EF-G•GDP•FA complex (Complex 2), shown in matching views with panels A-C: (D) Densities for both ttRRF and eEF-G can be seen in the inter-subunit space, with both domains of ttRRF and all five domains (I-V) of EF-G (orange) readily identifiable (see also Supplementary Figure S1B for a stereo viewing). Position occupied by domain II of ttRRF in panels A-C is now occupied by domain V of EF-G; (E) the same complex is shown from the 50S interface side (without 30S subunit) to reveal densities corresponding to ttRRF (in intermediate position Pi), EF-G, and deacylated tRNA (in the P/E and E sites); and (F) the same complex shown along with density corresponding to mRNA (dark blue; see Supplementary Figure S3 for complete mRNA density) from the 30S interface side, without the 50S subunit. A marked shift in RRF position can be seen when compared with the RRF position in panel C. Landmarks of the 30S subunit: bk, beak; hd, head; pt, platform; sh, shoulder; sp, spur; h44, 16S rRNA helix 44; and S2, protein S2.

Landmarks of the 50S subunit: CP, central protuberance; L1, protein L1 stalk; L7/L12-CTD, C-terminal domain of protein L7/L12; Sb, stalk base (protein L11 region); St, L7/L12 stalk; and H69, 23S rRNA helix 69. In panels D and E, densities corresponding to extended St and L7/L12-CTD are shown at slightly lower threshold values than the threshold value used for displaying the rest of the 50S subunit.

Figure 2. Interactions between the ribosome and ttRRF in Complex 1 (A, B), and between ttRRF and EF-G in Complex 2 (C, D). (A) Flexibly fitted atomic structure of ttRRF (red Ribbons, PDB ID; 1EH1) into the corresponding RRF density (semitransparent pink). Contacts and proximities between the amino acid residues of ttRRF and ribosomal components, such as amino acid residues of proteins L11 and S12 and nucleotides of the 23S rRNA helices 43 (H43) and 69 (H69) are indicated. Domains I and II of the fitted ttRRF coordinates are depicted in red and purple, respectively. (B) Same as in panel A, but rotated around a horizontal axis by ~90 degrees, to reveal interactions between an amino acid residue of domain II of ttRRF and conserved nucleotides of the 23S rRNA helix 95 (H95), and between domain I of ttRRF and helix71 (H71) of the 23S rRNA. (C) Flexibly fitted atomic structures of ttRRF and the homology model of *E. coli* EF-G into the corresponding cryo-EM densities of ttRRF (semitransparent red, in the intermediate position Pi) and EF-G (semitransparent orange), respectively, are shown with the ribosomal components present in immediate vicinity of RRF. EF-G domains I-V are shown in distinctive colors, I (orange), II (brown), III (green), IV (orange), and V (yellow). L27-NTT refers to the N-terminal tail of protein L27. (D) Enlarged boxed area in panel C to reveal ribosomal neighborhood of ttRRF in position Pi and interaction of domain II of ttRRF with EF-G domains III-V. Five select pairs of amino acid interactions are labeled (see also Table S1B for the complete list of amino acids interactions between ttRRF and EF-G). All helices of the 16S and 23S rRNAs are identified with h and H, respectively; while ribosomal proteins of the small and large subunits are prefixed by S and L, respectively. Thumbnails to the lower left of panels A, B, and D depict overall orientations of the ribosome.

Figure 3. Comparison of ttRRF binding positions before and after EF-G binding.

(A, B) Densities corresponding to ttRRF within the cryo-EM maps and (C) flexibly fitted atomic coordinates of Complex 1 (pink) and Complex 2 (red) are superimposed. The overall direction and magnitude of ttRRF movement is depicted by straight arrows. The direction of rotational shift in domain II of ttRRF, upon EF-G binding, is indicated by curved arrows (see also Supplementary Figure S6). Thumbnails to the lower left in panels A and C depict the orientation of the 70S ribosome: In panel A, the ribosome is in top view, whereas in panels B and C it is shown from the L7/L12-stalk side of the 50S subunit to reveal the shift in the elbow region of RRF.

Figure 4. Positions of domain IV and V of EF-G in the presence and absence of ttRRF on the ribosome. (A) Corresponding cryo-EM densities of domain IV and V of EF-G in Complex 2 (orange) and the 70S•EF-G•GDP•FA complex (semitransparent green (Datta *et al*, 2005)) are superimposed. (B) Same as in panel 1, but the transparency has been switched to reveal marked shifts in the position of EF-G domains. Dashed and solid circles in panels A and B highlight the equivalent positions in semitransparent and solid densities. (C) Flexibly fitted coordinates of EF-G domains IV and V into the maps shown in panel A are superimposed. Arrows depict the direction and magnitude of shifts in EF-G domains as compared to their positions in the absence of ttRRF.

Figure 5. Movements of domain II of RRF, and conformational changes of ribosomal components that interact with RRF and EF-G. Fitted atomic structures of the ribosomal components in two functionally relevant regions are shown. (A, B) The L11 stalk-base region; and (C, D) the bridge B2a region, shown along with protein S12. In each panel, fitted structures of two functional states are shown. In panels A and C the ribosomal components of the PoTC

complex are shown as semitransparent Ribbons, while those of Complex 1 are shown as solid Ribbons. Similarly, in panels B and D ribosomal components of complex 1 are shown as semitransparent Ribbons, whereas those of Complex 2 are shown as solid Ribbons. Arrows point to movements in the ribosomal components and RRF. Domain II of RRF in specified positions (panels A, C, and D), domains V and G' of EF-G (panel B), and domain III of EF-G (panel D) are also shown. Since only a weak density for domain II of RRF in P1/Iib configuration was observed in this study, its corresponding position in panel C is based on previous studies (Agrawal *et al.*, 2004; Pai *et al.*, 2008). The position of L7/L12-CTD shown in panel B is similar to that derived in previous cryo-EM (Datta *et al.*, 2005) and X-ray crystallographic (Gao *et al.*, 2009) studies (also see Supplementary Figure S10). Thumbnails to lower left of panels B and D depict the orientations of the 70S ribosome in panels A and B and panels C and D, respectively.

Figure 6. Conformational changes of the ribosome and tRNA movement due to binding of RRF and EF-G. Each structure was aligned, using the core portion of the 50S subunit of the cryo-EM maps as the main guide. **(A, B)** Movement of the L1 stalk (light blue, rRNA helices 77 and 78; dark blue, protein L1) and tRNA (green) during transition from **(A)** Complex 1 to **(B)** Complex 2. **(C, D)** Movement of the 30S subunit head (light brown, 16S rRNA; dark brown, 30S subunit head proteins) during transition from **(C)** Complex 1 to **(D)** Complex 2. Cryo-EM densities are shown as semitransparent grey. In all panels the ribosomes are viewed from the top, in an overall orientation depicted in the thumbnail at the lower right. In panels A and B, only the L1 stalk region (corresponding to left boxed area on the thumbnail) of the 50S subunits, and in panels C and D, only the head portion (corresponding to lower right boxed area on the thumbnail) of the 30S subunits are shown (see also Figure S7 for the overall conformational changes). The inset shown below panel B depicts the split densities for tRNA anticodons between the P (semitransparent) and E (solid) sites in Complex 2. Arrows indicate major movements (see Supplementary Figure S8 for stereo viewing of the tRNA movement).

Landmarks: ac, anticodon end; CCA, acceptor end; H78, helix 78 of the 23S rRNA; S13, small ribosomal subunit protein S13. The rest of the landmarks of the ribosomal subunits are the same as in Figure 1.

Figure 7. Schematic diagram showing steps of the recycling process. (A) The model post-termination complex (PoTC), with 30S (yellow) and 50S (blue) subunit. The head (hd) of the 30S subunit is depicted in a darker-shade block to indicate its relative position with respect to the rest of the 30S subunit body. Three canonical tRNA-binding sites, A, P and E, are depicted by dotted lines on both ribosomal subunits. The deacylated tRNA (green) is shown to fluctuate between P/P (P-tRNA, upper panel) and P/E (P/E-tRNA, lower panel) sites between the two ribosomal populations in dynamic equilibrium. (B) Initial binding of RRF in position P1/IIa, where domain II of RRF is oriented towards the 50S subunit and tRNA is exclusively in the P/E site, with the 30S subunit rotated in an anticlockwise direction (Complex 1). (C) Subsequent reorientation of domain II towards the 30S subunit to attain position P1/IIb. (D) Binding of EF-G shifts RRF to position Pi/IIIi, tRNA anticodon moves from P to E site in a subpopulation of Complex 2, while the head of the 30S subunit is fully rotated back to its original position. (E) Depiction of final disassembly of the PoTC through concomitant movement of RRF to P2/IIb, a position exclusively attained on the 50S subunit during the subunit dissociation (Barat *et al*, 2007). The precise sequence of events that leads to release of all bound components is not known, while binding of IF3 to the 30S subunit keeps the latter from re-associating with the 50S subunit to facilitate the translation re-initiation. Relative positions of the blocks representing 30S subunit body and head in panels A-D depict movements of those 30S-subunit domains with respect to the 50S subunit.

