Structure of the ribosome with elongation factor G trapped in the pretranslocation state

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During protein synthesis, tRNAs and their associated mRNA codons move sequentially on the ribosome from the A (aminoacyl) to the P (peptidyl) site. This process is catalyzed by the universally conserved ribosome-dependent GTPase (elongation factor G (EF-G) in prokaryotes and elongation factor 2 (EF-2) in eukaryotes). Although the high-resolution structure of EF-G bound to the posttranslocation ribosome has been determined, the pretranslocation conformation of the ribosome bound with EF-G and A-site tRNA has evaded visualization owing to the transient nature of this state. Here we use electron cryomicroscopy to determine the structure of the 70S ribosome with EF-G, which is trapped in the pretranslocation state using antibiotic viomycin. Comparison with the posttranslocation ribosome shows that the small subunit of the pretranslocation ribosome is rotated by ∼12° relative to the large subunit. Domain IV of EF-G is positioned in the cleft between the body and head of the small subunit outwardly to the A site and contacts the A-site tRNA. Our findings suggest a model in which domain IV of EF-G promotes the translocation of tRNA from the A site to the P site as the small subunit spontaneously rotates back from the hybrid state into the nonrotated posttranslocation state.

A/P* hybrid state | translation | single particle analysis | FREALIGN | real-space refinement

The ribosome decodes genetic information and synthesizes proteins in all living organisms. Although the movement of tRNA and mRNA through the ribosome is fundamental for protein synthesis, the molecular mechanism of translocation is not fully understood. Translocation occurs in two sequential steps (1). Subsequent to the peptideyl-transfer reaction, the acceptor ends of the peptideyl- and decacylated tRNAs first move spontaneously relative to the large (50S) ribosomal subunit, from the classic A/A and P/P states into the hybrid A/P and P/E states (A, aminoacyl; P, peptidyl; E, exit). In the second step, elongation factor G (EF-G) catalyzes a coupled movement of the anticodon stem-loops of tRNAs and mRNA on the small (30S) ribosomal subunit, placing tRNAs into the posttranslocation P/P and E/E states. The movement of tRNAs into the hybrid states is accompanied by a counterclockwise rotation of the small (30S) subunit relative to the large (50S) subunit (2–5). Translocation of mRNA and anticodon stem-loops of tRNAs is completed in the presence of EF-G during reverse (clockwise) rotation of the small subunit (6–8).

Although remarkable progress in studies of translocation has been made in recent years, important questions remain unanswered: how does EF-G rectify nonproductive spontaneous ratchet-like intersubunit rotation and tRNA fluctuations to produce efficient translocation? Does EF-G bias diffusion-driven movements of tRNA, or does it promote translocation by converting energy derived from GTP hydrolysis into mechanical movement? Answers to these questions would benefit from the 3D reconstruction of all intermediates of the translocation pathway. Several structures of EF-G–ribosome complexes in which EF-G was bound either to the nonrotated classic state or to the rotated hybrid states of the ribosome have been determined by X-ray crystallography (9–13) and electron cryomicroscopy (cryo-EM) (5, 14, 15). Despite significant differences in ribosome conformations, none of the known EF-G–ribosome structures contain A-site peptidyl-tRNA. In most available structures, domain IV of EF-G, which is known to be critical for translocation, is bound to the 30S A site (5, 9–12). Because the 30S A site cannot be simultaneously occupied by a tRNA and domain IV of EF-G, this conformation of EF-G–ribosome complex is not compatible with a pretranslocation state. An attempt to visualize EF-G bound to the pretranslocation ribosome containing both A- and P-site tRNAs using cryo-EM (16) produced density maps that lacked structural detail and are considered unreliable (17) because of their low (18–20 Å) resolution. A conformation of EF-G–ribosome complex possibly compatible with the pretranslocation state was recently observed in two cryo-EM (15) and crystal (13) structures of EF-G–ribosome complexes containing a single P-site tRNA. In these structures, EF-G displayed a conformation similar to previously observed conformations in other EF-G–ribosome complexes. However, domain IV of EF-G was not fully docked into the A site owing to a large (∼18°) swiveling of the 30S head along the direction of tRNA translocation. Nevertheless, because A-site tRNA was not present, these structures do not directly report on the pretranslocation state containing EF-G and tRNAs in the A and P sites.

Significance

The ribosome decodes genetic information and synthesizes proteins in all living organisms. To translate the genetic information, the ribosome binds tRNA. During polypeptide chain elongation, the tRNA is moved together with the mRNA through the ribosome. This movement is called translocation and involves precisely coordinated steps that include binding of a protein called elongation factor G (EF-G). How exactly EF-G drives translocation is not fully understood. We show in this study a detailed three-dimensional molecular image of the ribosome bound to EF-G and two tRNAs, just before the tRNAs are translocated. The image provides mechanistic clues to how EF-G promotes tRNA translocation.

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Data deposition: The atomic coordinates have been deposited in the Protein Data Bank, www.pdb.org (PDB ID codes 3JST, 3JSU, 3JSW, and 3JSX). The cryo-EM maps have been deposited in the EMDatabank, www.emdatabank.org (access codes EMD-5796–EMD-5800).

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Here we present a cryo-EM reconstruction of the pretranslocation complex at 7.6-Å resolution, obtained by increasing complex stability with viomycin and using a classification algorithm to distinguish between different ribosome conformations in a heterogeneous dataset (18). The conformation of EF-G observed in this structure is significantly different from all previously seen structures of EF-G–ribosome complexes, providing evidence that EF-G undergoes significant structural rearrangement during translocation.

Results

Overall Structure. To gain insight into the mechanism of translocation, we designed experiments to determine the structure of EF-G–ribosome trapped in the pretranslocation state using cryo-EM. Translocation, which normally occurs upon EF-G•GTP binding, was inhibited by adding antibiotic viomycin to *Escherichia coli* ribosomes containing dipeptidyl N-acetyl-Met-Phe-tRNA^Phe^ in the A site and deacylated tRNA^Met^ in the P site. Viomycin also stabilizes the hybrid-state conformation of the ribosome (3, 19, 20) but does not interfere with EF-G binding to the ribosome or GTP hydrolysis by EF-G (21, 22). The EF-G–ribosome complex was assembled in the presence of GTP and fusidic acid (Fus), an antibiotic that inhibits EF-G release after ribosome complex was assembled in the presence of GTP and the ribosome or GTP hydrolysis by EF-G (21, 22). The EF-G–ribosome (3, 19, 20) but does not interfere with EF-G binding to EF-G binding or GTP hydrolysis (21, 22). Surprisingly, there is no clear density in the map that can be attributed to Fus in this structure, EF-G binding is likely stabilized by viomycin, and helix 69 of 23S rRNA as previously seen in the crystal structure of viomycin–ribosome complexes (11, 27). At this site viomycin does not directly contact EF-G, consistent with previous cryo-EM and X-ray structures of EF-G bound to either nonrotated, hybrid-state ribosomes (5, 10–13, 15), we observe domain IV of EF-G in the pretranslocation state outside of the 30S A site, which is occupied by peptidyl-tRNA (Fig. 1C). The conserved loop at the tip of domain IV of EF-G (residues 507–514 in *E. coli* EF-G) makes extensive contact with the anticodon loop of the A-site tRNA (Fig. 1C). In the posttranslocation EF-G–ribosome complex (9), the same loop is inserted into the minor groove of the helix formed by the P-site.
tRNA and mRNA codon. Thus, domain IV of EF-G maintains its interaction with peptidyl-tRNA as tRNA translocates from the A to P site. This observation supports the hypothesis that the interaction between domain IV of EF-G and A-site tRNA facilitates translocation by displacing conserved bases of 16S rRNA G530, A1492, and A1493, which stabilize tRNA binding to the A site by interacting with the minor groove of the codon-anticodon helix (9).

Compared with the posttranslocation complex, the tip of domain IV is located more than 20 Å farther from the A site and toward the periphery of the 30S subunit (Fig. 2A). The large movement of EF-G as a whole between the pre- and posttranslocation states results mainly from a ∼20° rotation of EF-G around the universally conserved sarcin–ricin loop (SRL, nucleotides 2653–2667 of 23S rRNA). The SRL, which interacts with the GTP binding pocket of domain I of EF-G (9, 35), remains static in the transition from the pre- to posttranslocation state of the EF-G ribosome complex. This is consistent with a previously proposed hypothesis that EF-G rotation around the SRL allows a concerted movement of domains III, IV, and V, resulting in movement of the tip of domain IV by up to 30 Å (Fig. 3D). The latter was obtained by real-space refinement of the 30S subunit, the L11 stalk is placed 7 Å farther from the A site toward the E site (Fig. 3D). Another mobile element of the large subunit, the L11 stalk, is moved inward by ∼5 Å and contacts the elbow of the P/E tRNA (Fig. 3C). In contrast to the posttranslocation ribosome where two tRNAs are bound in classic P/P and E/E states, in the pretranslocation ribosome, the L1 stalk, a mobile domain of the large subunit, is moved inward by ∼5 Å and contacts the elbow of the P/E tRNA (Fig. 3C). Another mobile element of the large subunit, the L11 stalk, is placed 7 Å farther from the A site than in the posttranslocation complex (Fig. 3D). Interactions between domain V of EF-G and the conserved loop of helix 43 of 23S rRNA (35) are similar in the pre- and posttranslocation states. Thus, the movement of the L11 stalk during translocation maintains the interactions of the stalk with EF-G as the latter rotates around the SRL (Fig. 3D).

In addition to the rotation around the SRL, EF-G undergoes an interdomain rearrangement. The superposition of domains I–II of EF-G bound to pre- and posttranslocation (9) ribosomes reveals a concerted movement of domains III, IV, and V relative to domains II and I, resulting in a shift of the tip of domain IV by up to 15 Å (Fig. 2B). The superposition of domains I–II of EF-G bound to pretranslocation ribosome and those of EF-G bound to a rotated ribosome with the vacant A site (11–13) also shows a concerted movement of domains III, IV, and V, resulting in a shift of domain IV by 10–15 Å (Fig. S5). These movements are consistent with previous reports implicating structural dynamics of EF-G in translocation. Indeed, restricting the conformational dynamics of EF-G by introducing an intramolecular disulfide cross-link between domains I and V was shown to abolish EF-G translocation activity (37). Furthermore, comparison of the crystal structures of ribosome-free EF-G (38–40) with the crystal structure of EF-G in the posttranslocation ribosome (9) reveals a significant structural rearrangement of domains III, IV, and V, resulting in movement of the tip of domain IV by up to 30 Å and indicating that EF-G is indeed dynamic. Our data demonstrate that domain IV of EF-G in pretranslocation ribosomes adopts an intermediate conformation between free EF-G and EF-G bound to the posttranslocation ribosome (Fig. S5D).

Structure of the Ribosome in the Pretranslocation State. The pre- and posttranslocation EF-G–ribosome complexes differ not only in their EF-G configuration but also in the conformation of the ribosome. Compared with the crystal structure of the posttranslocation EF-G–ribosome complex (9), the small subunit in the pretranslocation ribosome is rotated counterclockwise by ∼12° relative to the large subunit (Fig. 3A). The head of the 30S subunit is swiveled by ∼3° relative to the 30S platform and body toward the E site (Fig. 3B). In contrast to the posttranslocation ribosome where two tRNAs are bound in classic P/P and E/E states, in the pretranslocation ribosome, the large subunit, the L11 stalk, is placed 7 Å farther from the A site than in the posttranslocation complex (Fig. 3D). Interactions between domain V of EF-G and the conserved loop of helix 43 of 23S rRNA (35) are similar in the pre- and posttranslocation states. Thus, the movement of the L11 stalk during translocation maintains the interactions of the stalk with EF-G as the latter rotates around the SRL (Fig. 3D).

In addition to the structure of EF-G bound to nonrotated, classic state ribosomes (9), a number of EF-G–ribosome complexes lacking A-site tRNA and showing various degrees of intersubunit rotation and swiveling of the 30S head have been solved recently using X-ray crystallography (10–13). Our structure of the pretranslocation EF-G–ribosome complex shows an intersubunit rotation that is larger than the rotations seen in any of the EF-G–ribosome crystal structures (12° vs. 7°). In contrast, our pretranslocation EF-G–ribosome complex exhibits a 30S head swivel (3°) that is smaller than that seen in any of the crystal structures. Hence, comparison of ribosome conformations in different EF-G–ribosome complexes supports results of recent kinetic experiments (8) suggesting that intersubunit rotation and head swiveling may occur sequentially rather than simultaneously during ribosomal translocation.

To analyze how EF-G binding affects the conformation of the pretranslocation ribosome, we compared the pretranslocation EF-G–ribosome complex with a structure of the EF-G-free ribosome containing tRNAs in the A/P and P/E states (Fig. S6). The latter was obtained by real-space refinement of the ribosome structure to a 7.6-Å resolution cryo-EM map of a respective class of ribosomes in our dataset (class IV; Methods). The ∼9.5° of
intermediate state of A-site tRNA translocation that we propose to name the A/P* hybrid state (Fig. 4C). A similar conformation of peptidyl-tRNA was observed in cryo-EM reconstructions of bacterial ribosome–tRNA complexes in the absence of EF-G (31), and of pretranslocation mammalian ribosomes obtained in the absence of EF-2 (42), suggesting that the A/P* state may be sampled spontaneously. Indeed, spontaneous sampling of two hybrid states of tRNA binding that differ in the position of the elbow of A-site tRNA was observed in pretranslocation ribosomes using single-molecule fluorescence resonance energy transfer (smFRET) (43). Comparison of pre- and posttranslocation complexes of the bacterial ribosome suggests that translocation of the A-site tRNA into the P site occurs at minimum in three consecutive steps. First, the acceptor end of tRNA shifts on the large subunit after peptidyl transfer (A/P state). Second, the tRNA elbow moves upon EF-G binding, positioning the peptidyl-tRNA in the A/P* state. In the third step, the anticodon stem loop of peptidyl-tRNA translocates into the P/P state during the reverse, clockwise rotation of the small subunit (Fig. 4C).

Discussion

Our cryo-EM reconstruction reveals the structure of a long-hypothesized intermediate of ribosome translocation in which EF-G is bound to the pretranslocation ribosome. We stabilized this translocation intermediate with viomycin, an antibiotic inhibiting translocation. In posttranslocation complexes domain IV of EF-G occupies the 30S A site, thus precluding tRNA binding at this site. In the pretranslocation state reported here EF-G adopts a conformation distinct from the posttranslocation state, permitting tRNA binding in the A site. Stabilization of the EF-G-bound pretranslocation state was possible because viomycin increases the affinity of peptidyl-tRNA to the A site more than 1,000-fold (21, 22). Because viomycin does not contact EF-G and was previously observed in a crystal structure of the ribosome with EF-G bound in the posttranslocation conformation (11), the pretranslocation conformation of EF-G observed in our structure is not directly induced by viomycin. Together with our finding that the structure of the pretranslocation EF-G–ribosome complex is similar to other structures of rotated, hybrid-state ribosomes (2, 4, 31, 32), this suggests that our cryo-EM reconstruction of pretranslocation EF-G–ribosome complex represents an on-pathway intermediate of translocation in the ribosome.

Our structure shows a number of unique features, most notably the ~12° rotation of the small ribosomal subunit and the adoption of the A/P* state by the peptidyl-tRNA in response to EF-G binding, and brings insights into the mechanism of translocation. Fig. 4C provides a minimal description of the translocation mechanism that can be inferred from comparing structures of pre- and posttranslocation states of EF-G–ribosome as well as structure of EF-G-free pretranslocation ribosome. Translocation of mRNA and the anticodon stem-loops of tRNAs was shown to occur during the movement of the small ribosomal subunit from the rotated hybrid state back into the nonrotated classic conformation (7). Comparison of pre- and posttranslocation states of the EF-G–ribosome complex suggests that domain IV of EF-G, which contacts the A-site tRNA, plays a central role in translocation: upon reverse (clockwise) rotation of the small subunit, domain IV of EF-G acts as a steric hindrance for the return of peptidyl-tRNA from hybrid A/P* into classic A/A state (Fig. 4B), pushing A-site tRNA in the forward direction (Fig. 4B). In the absence of EF-G, tRNA fluctuations between classic and hybrid states on the 50S subunit, as well as intersubunit rotation, may occur spontaneously at rates comparable to rates of EF-G-catalyzed mRNA translocation (19, 33, 43, 44). Therefore, it has been hypothesized that thermal energy is sufficient to drive tRNA and mRNA translocation upon spontaneous reverse rotation of the small subunit (15, 32, 45–47). The comparison of pre- and posttranslocation states of EF-G–ribosome complex
now reveals how spontaneous intersubunit rotation may be rectified into translocation by EF-G binding to the ribosome.

Our structure of the pretranslocation EF-G–ribosome complex exposes significant movement of EF-G domains III, IV, and V relative to domains I and II compared with the structures of ribosome-free EF-G and EF-G bound to the posttranslocation ribosome (Fig. S5D). In addition, EF-G as a whole undergoes a ~20° rotation around the SRL of the 50S subunit in the course of translocation (Fig. 2A). The transition of EF-G from the conformation observed in our structure into the conformation observed in the posttranslocation ribosome (9) includes a movement of domain IV toward the A site. This structural rearrangement of EF-G may drive tRNA movement from the A to P site on the small subunit, thus promoting translocation.

Alternatively, movement of domain IV of EF-G to the A site of the small subunit may follow spontaneous movement of peptidyl-tRNA from the 30S A to P site, making tRNA translocation irreversible. Existing FRET and biochemical data do not provide evidence for frequent spontaneous fluctuations of tRNA between A and P sites of the 30S subunit in the absence of EF-G. Therefore, for this translocation mechanism to be possible, EF-G must destabilize binding of peptidyl-tRNA to the A site. The destabilization may then lead to spontaneous movement of tRNAs between binding sites, and EF-G could, in addition to its destabilization action, bias the movement toward the P site. Higher-resolution structures of pretranslocation EF-G–ribosome complexes may provide evidence for the destabilization of A-site tRNA by EF-G. In addition, smFRET experiments may allow distinction between coincident movement of domain IV of EF-G and tRNA, and EF-G movement that follows that of tRNA.

Although comparison of the pretranslocation EF-G–ribosome complex obtained in this work with the structure of posttranslocation EF-G–ribosome complex previously visualized by X-ray crystallography provides important clues about the mechanism of EF-G-induced translocation of tRNA and mRNA, additional intermediates of translocation likely exist. A large (up to 18°) swivel of the 30S head was observed in several EF-G–ribosome structures (11, 13, 15). This movement was hypothesized to open a wide (~20 Å) path for tRNA translocation between the P and E site on the small subunit that is otherwise constricted by RNA residues of the 30S head and platform (48). Recent kinetic studies (8) suggested that intersubunit rotation precedes the 30S head swiveling, whereas mRNA translocation, reverse rotation of body and platform of the small subunit, and back-swiveling of the 30S head seem to occur concurrently. Our pretranslocation EF-G–ribosome structure shows a large (12°) degree of intersubunit rotation and a relatively small (3°) 30S head swivel. Hence, our pretranslocation EF-G–ribosome structure likely represents an early translocation intermediate, formation of which is probably followed by additional 30S head swiveling. Further studies will be required to reconstruct the complete translocation pathway.

Fig. 4. Intermediate states of tRNA translocation. (A) Structural alignment of 23S rRNA in cryo-EM structures of the EF-G-free (light pink) and EF-G-bound (light blue) pretranslocation ribosomes (both determined in this work) reveals that EF-G binding induces the movement of peptidyl-tRNA into the A/P* hybrid state described in this work. For clarity, only EF-G (red), peptidyl-tRNA, and helices 38 (the A-site finger, ASF) and 84 (h84) of 23S rRNA are shown, whereas other structural components of the ribosome are omitted. Peptidyl-tRNA bound to the A/P hybrid state in the EF-G-free ribosome is shown in cyan; peptidyl-tRNA bound to the A/P* hybrid state in the EF-G-bound ribosome complex is shown in dark green. (B) Position of the intermediate tRNA A/P* state in the pretranslocation EF-G-bound ribosome structure (dark green) relative to the A/P state of the EF-G-free hybrid-state ribosome (cyan) and classic A/A- (yellow) and P/P- (orange) tRNAs of the nonrotated, classic-state ribosome (9, 53). Superpositions in A and B were obtained by structural alignment of 23S rRNA from their respective complexes. (C) Schematic depiction of intersubunit rotation and tRNA movement during translocation. After peptidyl-transfer from the P-(orange) to the A-site (yellow) tRNA, the ribosome undergoes spontaneous fluctuations between the nonrotated (classic) and rotated (hybrid) states. Binding of EF-G-GTP (red) induces an extra 2.5° rotation of the 30S subunit (in addition to the 9.5° rotation observed in the EF-G-free hybrid-state ribosome) and shifts peptidyl-tRNA into the A/P* state. Upon subsequent clockwise rotation of the small subunit, domain IV of EF-G moves into the 30S A site and promotes the translocation of tRNAs and their associated mRNA codons on the small subunit. In addition to the two intermediate complexes previously described in this work, the EF-G-GDP (red) intermediate shown in (B) allows the conformation observed in our structure into the conformation observed in the posttranslocation ribosome (9) to be followed by subsequent clockwise rotation of the 30S subunit, thus promoting translocation.

Methods
A detailed description of the study methods can be found in SI Methods.

EF-G-Bound Ribosome Preparation. Fus and viomycin were purchased from Sigma and USP, respectively; tRNA<sup>Met</sup> and tRNA<sup>Phe</sup> from E. coli were purchased from MP Biomedicals and Chemblock, respectively. E. coli ribosomes, aminoacylated tRNAs, His-tagged EF-Tu, and EF-G were prepared as previously described (6). Pretranslocation complex was assembled by incubating 70S ribosomes (0.4 μM) with N-acetyl-Met-tRNA<sup>Met</sup> (0.8 μM) and mRNA m32 (6) (0.8 μM) in buffer
Cryo-EM Data Collection and Image Analysis. More than 1.3 million cryo-EM images of ribosome complexes (Fig. 1) were acquired using an FEI Titan Krios, a cryo-EM microscope fitted with five distinct classes and cryo-EEM maps (classes IV and V) is G®, for each complex. The refined model of EF-G fully accounts for the local cryo-EM density (Fig. 5B). The all-atom rmsd between ribosomal RNAs of each refined model and the starting crystal structure (32) is 1.2 Å; all-atom rmsds between the six best models of EF-G also do not exceed 1.2 Å (Fig. 59), similar to the expected coordinate error of cryo-EM structures at 7.6 Å resolution (32).

Modeling of the Cryo-EM Maps Using Atomic Models. The cryo-electron microscopic (cryo-EM) density of the hybrid-state 70S{RNA}{RF} complex (32) (excluding the RF molecule) was used as a starting model for structure fitting in Chimera (49). A homology model of E. coli EF-G was created using the SWISS-MODEL (50) from Thermus thermophilus EF-G (Protein Data Bank ID 2BM0) (51). Secondary structure elements of the ribosome complexes were manually fitted and then refined using stereochemically restrained rigid-body space-re refinement (25, 26). The correlation coefficient between refined stereochemically restrained rigid-body space-re fineement as a tool in model building. Acta Crystallogr D Biol Crystallogr 58(Pt 5):671–677.


Ribosomal complexes were thawed on ice and diluted (1:2.5) in buffer containing 10 mM Hepes-KOH (pH 7.6), 5 mM MgCl₂, 50 mM NH₄Cl, 2 mM spermidine, 0.1 mM spermine, 6 mM β-mercaptoethanol, viomycin (0.5 mM), GTP (0.5 mM), and fusidic acid (0.5 mM).

C-flat 1.2-1.3 grids (400-mesh) were freshly glow discharged immediately before cryo-plunging with a current of ~20 mA for 45 s in an EMITECH K100X glow discharge unit. Grids were then loaded into an FEI Mark II Vitrobot, which was allowed to equilibrate to 95% relative humidity at 22 °C. Two microliters of 45 s in an EMITECH K100X glow discharge unit. Grids were diluted (1:2.5) in buffer containing 10 mM Hepes-KOH (pH 7.6), 5 mM MgCl₂, 50 mM NH₄Cl, 2 mM spermidine, 0.1 mM spermine, 6 mM β-mercaptoethanol, viomycin (0.5 mM), GTP (0.5 mM), and fusidic acid (0.5 mM).

Electron Microscopy. Sample imaging was performed using an FEI Titan Krios microscope operated at 300 kV. The nominal defocus was varied from 2.8- to 4.4-μm underfocus, with refocusing performed every 10 μm. Images were collected on a Falcon I direct electron detector, with a total dose of 30 electrons/Å² and a calibrated pixel size of 1.04 Å.

Image Processing. Particles were semiautomatically selected using e2boxer’s swarm tool (1), followed by manual curation of the dataset. Defocus parameters were determined using CTFIND3 (2). Boxing was performed using batchboxer (3), with unbinned images having a box size of 320 pixels.

Initial alignment parameters were assigned using IMAGIC (4). Particles were normalized, fivefold binned, phase flipped to account for the contrast transfer function, band-pass filtered with cut-offs of 0.03 and 0.33 (Nyquist units), and masked with a circular mask with radius 0.62 (fraction of half the image size) and a fall-off of 0.08.

An initial reconstruction was obtained from 20,000 particle images that were aligned against a published 70S ribosome structure (5) (EMD-1315). This reference was projected at angular intervals of 7.5° to generate reference images for alignment. The projections were then filtered and masked identically to the experimental images. Particle alignment parameters were determined using multireference alignment implemented in IMAGIC and exported to FREALIGN (6) using IMAGIC’s Imagic2realign tool.

Further processing with FREALIGN was carried out as previously described (7). Initially, refinement and classification were performed on only half of the dataset. The data were refined against a single reference until no further improvement was seen in resolution as indicated by the calculated Fourier shell correlation (FSC) between rounds. Refinement included data up to 35 Å initially and data at higher resolution as refinement progressed, up to a resolution 18 Å in the final round of refinement. Two-fold binned data were then divided into 15 classes using RSAMPLE (7) and classified without refining the alignment parameters, while including data between 150 and 15 Å. This yielded five classes with recognizable features at high resolution [better than 15 Å, FSC = 0.143 (8)]. One of these classes showed density for A/P site and P/E site tRNA, as well as density that appeared at a higher threshold where one would expect EF-G, indicating that it was heterogeneous with respect to EF-G occupancy. We extracted particles that belonged to that class (particles with an occupancy greater than 50% within this class) and repeated classification into three additional classes, using RSAMPLE to generate initial particle assignments. Data between 150- and 12-Å resolution was included in this classification. This classification gave two distinct classes, with and without EF-G, with a resolution between 8 and 9 Å [FSC = 0.143 (8)]. The third class yielded a reconstruction that did not display high-resolution features.

To save computational resources, we used the classification results from the first half of the dataset as a starting point for the classification of the full dataset. The full dataset was first refined against a single class, after which the particles were again classified into 15 classes. The particles in the first half of the dataset were given the class occupancy they had in the first round of classification, whereas the particles in the second half of the dataset initially had 0% occupancy with respect to all of the classes. The classes were classified using fourfold binned data. The classification results were essentially identical to those observed using only the first half of the dataset. The class that showed density for A/P site and P/E site tRNA, as well as density for EF-G, was once again classified further into 3 classes, using RSAMPLE to generate random initial class assignments, yielding two additional classes.

Real-Space Refinement of Structural Models. The crystal structure of the hybrid-state 70S•tRNA•RRF complex (9) (excluding the RRF molecule) was used as a starting model for structure fitting in Chimera (10). A homology model of Escherichia coli EF-G was created using the SWISS-MODEL (11) from Thermo thermophilus EF-G [Protein Data Bank (PDB) ID 2BM0] (12), followed by manual fitting of EF-G domains into the cryo-EM map. Homology models for Escherichia coli proteins L1, L10, L11, and L12 were built using PDB ID 3U4M, IZAV, 1MMS, and 1IDD, respectively. Upon initial fitting of the 70S•tRNA•EF-G structure in Chimera using the “Fit in Map” procedure, the structure was refined in several steps using stereochemically restrained rigid-body real-space refinement, implemented as a module of CNS (13), allowing refinement against cryo-EM maps (14, 15). Refinement was carried out essentially as described previously (16). In the first step of refinement, the ribosomal subunits, tRNAs, and EF-G were treated as rigid groups (5-rigid-group refinement). In the next step, a finer definition of rigid groups was used, in which the ribosome was split into domains known to be able to move with respect to each other. In the final round of rigid-body refinement, secondary structure elements of ribosomal RNA, ribosomal proteins, tRNA, and EF-G were treated as rigid-groups, resulting in a 402-rigid-group definition. At all stages of refinement covalent-bond and non-bonding constraints were used, and the relative weighting of the experimental and stereochemical energy terms were optimized. The correlation coefficient between refined structures and cryo-EM maps (classes IV and V) is 0.86 for each complex. An FSC curve was also calculated against the model and falls below a value of 0.5 at 7.6-Å resolution, in agreement with the earlier resolution estimate (Fig. S2B). The all-atom rmsd between ribosomal RNAs of each refined model and the starting crystal structure (9) is 1.2 Å, similar to the expected coordinate error of cryo-EM structures at 7.6-Å resolution (17). To further validate the position of EF-G, we compared several refined models yielding highest correlation coefficients and resulting from different starting models (alternatively fitted in the map manually or by the “volume-fitting” procedure in Chimera) and from different refinement strategies (different relative weightings of the experimental cryo-EM fit term and stereochemical restraints, and alternative definitions of secondary-structure elements or domains of EF-G). The six structural models of EF-G with best fits to the map are very similar (Fig. S9), with all-atom
rmsds not exceeding 1.2 Å, consistent with the expected coordinate error as discussed above. In sum, the changes in the positions of ribosomal components (such as L1 and L11 stalks), A/P tRNA, and EF-G discussed in our work are significant because they are significantly larger than the coordinate error resulting from the fitting of crystal structures into experimental maps.


Fig. S1. Cryo-EM micrograph showing 2D projections of *E. coli* ribosomes prepared with tRNA, EF-G, fusidic acid, and viomycin.
Fig. S2. Single-particle classification. (A) Sorting of ~1.3 million particles resulted in five distinct classes: (I) pretranslocation ribosomes containing tRNAs in the A (aminoacyl), P (peptidyl), and E (exit) site without EF-G bound (26.7%); (II) posttranslocation ribosomes containing EF-G and tRNAs in the P and E sites (13.4%); (III) ribosomes containing EF-G and tRNA in the P site on the small subunit (6.8%); (IV) pretranslocation ribosomes containing tRNAs in the A and P site on the small subunit (3.5%); and (V) a previously unseen class of ribosomes containing EF-G and tRNAs bound to A and P sites of the small subunit (2.4%). Class averages are shown in surface representation, with 50S subunits colored in light blue, 30S subunits in yellow, tRNAs in green, and EF-G in red. The remaining 47.2% of particles did not show distinct features (rows 2 and 3). (B) Resolution curves for classes I–V of EF-G–ribosome complexes. For convenience, the x axis is labeled with both spatial frequency (Å⁻¹) and resolution (1/spatial frequency, Å). The resolution for each class stated in the text corresponds to an FSC = 0.143 cutoff value (1), shown as a dotted line.

Fig. S3. Close-up views of regions of the structural model of the pretranslocation EF-G–ribosome complex refined into the 7.6-Å cryo-EM map. (A) Side view of peptidyl-tRNA (green) in the A/P* state. (B) Side view of deacylated tRNA (orange) in the P/E state. (C) Codon–anticodon interactions of A/P* and P/E tRNAs with mRNA (yellow). (D) Interactions of the P/E tRNA with the L1 stalk. (E) Intersubunit bridge B1b formed by ribosomal proteins L5 and S13. Cryo-EM density map is shown in gray mesh; 23S rRNA in light blue; proteins of the large subunit in pink; 16S rRNA in light gold; and proteins of the small subunit in wheat.
Fig. S4. Densities for antibiotics viomycin and fusidic acid. Close-up views of the viomycin binding site in (A) the pretranslocation EF-G-bound state (class V, this work); (B) the pretranslocation EF-G-free complex (class IV, this work); and (C) the pretranslocation EF-G-free complex obtained in the absence of viomycin [EMDataBank ID EMD-1541 (1)]. The cryo-EM maps (gray) show clear density for viomycin in its expected binding site (2, 3), whereas the reconstruction obtained in the absence of viomycin (C) shows no density there. 23S rRNA is shown in light blue; A/P tRNA in cyan, A/P* tRNA in dark green; 16S rRNA and mRNA in yellow; proteins of the small subunit in wheat; and viomycin in red. (D) Superposition of the pretranslocation (red, this work) and posttranslocation [blue (4)] EF-G structures highlighting the fusidic acid binding site. Superposition was achieved by aligning domains I–II of the two EF-G structures. Domains of EF-G are labeled with Latin numbers. Fusidic acid, from the posttranslocation EF-G–ribosome structure [PDB ID 2WRI (4)], is shown in yellow. The black box indicates the region shown in E and F. (E) View of the fusidic acid binding site of the pretranslocation EF-G-bound ribosome (class V, this work). (F) View of the fusidic acid binding site of the posttranslocation EF-G-bound ribosome (class II, this work). The map shown in F was amplitude-scaled to the map shown in E using the program diffmap (http://grigoriefflab.janelia.org/diffmap).

Fig. S5. Comparison of conformations of EF-G in various EF-G–ribosome complexes. Domains I–II of the EF-G structures were superimposed. The pre-translocation (this work) and posttranslocation (1) states of EF-G are shown in all panels in red and blue, respectively. (A) EF-G from the *E. coli* 70s*EF-G*GDPCP*viomycin* complex is shown in light orange [PDB ID 4KJ4 (2)]. (B) EF-G from the *T. thermophilus* 70s*tRNA*mRNA*EF-G*GDPCP complex is shown in cyan [PDB ID 4JUW (3)]. (C) EF-G from the *T. thermophilus* 70s*tRNA*mRNA*EF-G*GDP*FUS complex is shown in yellow [PDB ID 4KDG (4)]. (D) EF-G as free protein bound with GDP is shown in green [PDB ID 1DAR (5)]. Domains of EF-G are labeled with Latin numerals.


Fig. S6. A 7.6-Å map of the EF-G-free pretranslocation ribosome complex (class IV). (A) Surface representation of the cryo-EM map. (B) Fitting of the refined model to the 7.6-Å cryo-EM map (gray surface). The backbone structure of the ribosomal 50S subunit is shown in light blue, the 30S subunit in light yellow, and A/P and P/E tRNAs are in cyan and orange, respectively.

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Fig. S7. Comparison of the positions of the small subunit in the EF-G-free (gray) and EF-G-bound (light gold) pretranslocation ribosome complexes (this work). Superposition of 23S rRNA structures (not displayed for clarity) from the EF-G-free and EF-G-bound ribosome complexes reveals that EF-G binding induces a ∼2.5° counterclockwise rotation of the small subunit. Only 16S ribosomal RNA is shown; ribosomal proteins are not displayed for clarity.
Fig. S8. Fit of the refined EF-G structure (red) to the cryo-EM map (gray) shown at contour levels (σ) 1.0 (A), 1.5 (B), 2.0 (C), 2.5 (D), and 3.0 (E). Domains of EF-G are labeled with Latin numbers.
Fig. S9. Comparison of models refined into the density corresponding to EF-G in the pretranslocation EF-G–ribosome complex (class V). The six models of EF-G with the highest correlation coefficients with the map (shown in yellow, red, green, blue, pink, and gold) were obtained by independent real-space rigid-body refinement runs, using alternative starting models and refinement protocols (SI Methods). The all-atom rmsds between the six best models of EF-G do not exceed 1.2 Å, consistent with the expected coordinate error for cryo-EM structures at 7.6-Å resolution as discussed in SI Methods.