The Cryo-EM Structure of a Complete 30S Translation Initiation Complex from Escherichia coli

Patricia Julián1, Pohl Milon2, Xabier Agirrezabala1, Gorka Lasso1, David Gil1, Marina V. Rodnina2, Mikel Valle1,3,

1 Structural Biology Unit, Center for Cooperative Research in Biosciences (CIC bioGUNE), Parque Tecnológico de Bizkaia, Derio, Spain, 2 Department of Physical Biochemistry, Max Planck Institute for Biophysical Chemistry, Göttingen, Germany, 3 Department of Biochemistry and Molecular Biology, Faculty of Science and Technology, University of the Basque Country, Bilbao, Spain

Abstract

Formation of the 30S initiation complex (30S IC) is an important checkpoint in regulation of gene expression. The selection of mRNA, correct start codon, and the initiator fMet-tRNAfMet requires the presence of three initiation factors (IF1, IF2, IF3) of which IF3 and IF1 control the fidelity of the process, while IF2 recruits fMet-tRNAfMet. Here we present a cryo-EM reconstruction of the complete 30S IC, containing mRNA, fMet-tRNAfMet, IF1, IF2, and IF3. In the 30S IC, IF2 contacts IF1, the 30S subunit shoulder, and the CCA end of fMet-tRNAfMet, which occupies a novel P/I position (P/I1). The N-terminal domain of IF3 contacts the tRNA, whereas the C-terminal domain is bound to the platform of the 30S subunit. Binding of initiation factors and fMet-tRNAfMet induces a rotation of the head relative to the body of the 30S subunit, which is likely to prevail through 30S subunit joining until GTP hydrolysis and dissociation of IF2 take place. The structure provides insights into the mechanism of mRNA selection during translation initiation.

Introduction

Initiation is the most regulated step of translation, at which the ribosome selects mRNAs according to their translation initiation regions (TIR) and establishes a correct reading frame. In bacteria, translation initiation is promoted by three initiation factors (IF), IF1, IF2, and IF3 [1,2]. The small ribosomal subunit, 30S, is recruited to single-stranded mRNA regions at the TIR [3]. The Shine-Dalgarno sequence (SD) of the mRNA binds to the anti-Shine-Dalgarno sequence (ASD) of 16S rRNA in a cleft between the head and the platform at the back of the 30S subunit, whereas mRNA wraps in a groove that encircles the neck of the 30S subunit [4-6]. IFs bind to the 30S subunit and work synergistically to accelerate initiation and ensure the choice of the correct start codon and initiator tRNA. IF1 is a one-domain compact protein which binds at the A site of the 30S subunit [7]. IF2 is a multi-domain GTPase that plays a major role in the recruitment of fMet-tRNAfMet to the 30S IC. The highly conserved C-terminal half of the protein (CTD) contains the GTP- and fMet-tRNAfMet-binding domains. The role of the less conserved N-terminal (NTD) is not clear, except that it may provide an additional anchor for IF2 on the 30S subunit [8,9]. Docking of fMet-tRNAfMet to the complex of the 30S subunit with mRNA and IFs completes the formation of the 30S IC [10]. The initiator tRNA is held in a characteristic position by two interactions: one involving the tRNA decoding stem which is buried in the P site of the 30S subunit, and the other between IF2 and the tRNA acceptor end [11]. The orientation of fMet-tRNAfMet in the complex differs from the canonical P-site position and was designated as P/I state [12]. IF2 and fMet-tRNAfMet provide a large interaction surface for binding the 50S subunit [11-13]. Formation of the 30S IC constitutes an important step for the selection of a favorable TIR. The ability of the ribosome to screen the TIR and to check for the fidelity of codon-anticodon interaction strongly depends on the presence of IF3 [14]. However, a structure of 30S IC with IF3 is not available.

IF3 consists of two domains, IF3C and IF3N, connected by a linker. The structures of the separate domains are known, but not of the full-length protein. The binding site for IF3 on the ribosome has been examined by a variety of biochemical techniques and by cryo-EM [15-18]. Overall, IF3 appears to bind to the platform of the 30S subunit, but the domain orientation of IF3 and exact binding contacts for each domain remain controversial. IF3 has several functions during translation initiation. It interferes with ribosomal subunit association [19], affects the rates of tRNA association to and dissociation from the P site [20,21], and ensures the fidelity of translation initiation [22-24]. Most importantly, IF3 is crucial for sensing the TIR of mRNA. In the absence of IF3, the ribosome is largely unable to discriminate against unfavorable TIRs or incorrect initiation codons [14,25].
Translation is the process by which a ribosome converts the sequence of a messenger RNA (mRNA)—produced from a gene—into the sequence of amino acids that comprise a protein. Bacterial ribosomes each have one large and one small subunit: the 50S and 30S subunits. Initiation of translation entails selection of an mRNA, identification of the correct starting point from which to read its code, and engagement of the initial amino acid carrier (tRNA). These events take place in the 30S subunit and require the presence of three initiation factors (IF1, IF2, IF3). Formation of this 30S initiation complex precedes joining with the 50S subunit to assemble the functional ribosome. By using a cryo-electron microscopy approach to visualize the structures without fixation or staining, we have determined the structure of a complete 30S initiation complex and identified the positions and orientations of the tRNA and all three initiation factors. We found that the presence of the initiation factors and tRNA induces rotation of the head relative to the body of the 30S subunit, which may be essential for rapid binding to the 50S subunit and for regulating selection of the mRNA. IF3 had not been seen previously in the context of the 30S structure and its visualization gives insight into a potential role in preventing association of the two ribosomal subunits. These findings are important for understanding how the interplay of elements during the early stages of translation selects the mRNA and regulates formation of functional ribosomes.

Results

Reconstruction of a Complete 30S IC

The 30S IC was prepared from E. coli 30S subunits, IF1, IF2, IF3, fMet-tRNAfMet, mRNA, and a GTP analog, GDPNP (Materials and Methods). The mRNA used for complex formation was m002, which contains a 9-nt SD and a 5-nt spacer between the SD and the AUG start codon [27]. The extended SD provides strong interactions with the ASD in the 16S rRNA and drastically reduces the dissociation rate of IF3 [14]; m002 is similar to mRNAs with “enhanced” SD used in previous structural and biochemical studies [11–14,26,28]. An initial set of 48,000 particles resulted in a cryo-EM map that showed density which could be attributed to tRNA and IF2 on the surface of the 30S subunit (Figure 1A) at positions expected from a previous cryo-EM reconstruction of the 30S subunit in complex with initiator tRNA and IF2 [11]. However, the densities in our initial reconstruction did not reveal structural details of the tRNA or IF2, suggesting heterogeneity of the sample due to differences in the occupancy of the 30S subunit with ligands and/or distinct conformational states. Separation into two classes by non-supervised maximum likelihood-based classification (ML3D) [29], a recently developed tool that has been successfully applied to other ribosomal samples [30], and independent image processing yielded the 3D maps shown in Figure 1B and 1C. The EM map for class 1 particles (23% of all particles) showed no density for IFs or tRNA (Figure 1B); Class 2 accounted for the majority (77%) of the particles, and the corresponding 3D map revealed the presence of fMet-tRNAfMet and IFs (Figure 1C). In the latter reconstruction, the putative densities for IF2 and tRNA could be seen at a higher density threshold compared with the map obtained from the total set of images. Structural details of the initiator tRNA from the X-ray structure [31] and of IF2 known from previous cryo-EM reconstructions [11–13] were clearly recognizable. Importantly, high-density threshold rendering revealed density for the SD-ASD duplex for both classes (Figure S1), indicating that class 1 represented 30S-mRNA complexes, rather than vacant 30S subunits. The resolutions estimated at 0.5/0.14 cut-off criteria in the Fourier shell correlation (FSC) were 16.8/14 Å (total set of images), 21/17 Å (class 1, representing the 30S-mRNA complex), and 18.3/15 Å (class 2, representing the 30S IC). Classification into more classes or re-classification of class 2 did not result in improvement of resolution or identification of further conformational states. The limited resolution of the reconstructions may reflect inherent flexibility within the 30S IC induced by IF3-IF1 binding to the 30S subunit, as suggested by kinetic experiments [14].

Segmentation of the Map for the 30S IC

After removing the density corresponding to the 30S subunit from the class 2 map (Figure 1C), positions for tRNAs and IFs could be assigned, which were consistent with previous structural and biochemical studies (Figure 1D and 1E). The main part of the resulting density could be attributed to the IF2-fMet-tRNAfMet complex placed along the shoulder and the cleft between the head and the body of the 30S subunit. The density for IF1 was not obvious within the 30S IC map, because its binding site was covered by the IF2 structure. However, fitting the crystal structure of IF1 bound to the 30S subunit [7] suggested that, in our map, IF1 can be embedded in the boundary between the 30S subunit and IF2 (Figure 1D) in the cleft formed by the 530 loop, helix 44 of 16S RNA, and ribosomal protein S12 [7]. The moderate resolution of the current 3D map and the lack of a complete atomic model for IF2 (see below) precluded an unambiguous definition of the boundary between IF1 and IF2. Additional densities observed close to the initiator tRNA and the platform of the 30S subunit were assigned to IF3 (Figure 1E). Thus, the cryo-EM map of the 30S IC had densities attributable to all the elements of the complete complex (see below for further details of fitting).
The 30S Subunit Head Rotation

To investigate whether the binding of initiation factors and fMet-tRNA<sup>Met</sup> changed the conformation of the 30S subunit, we compared the structures of 30S subunits from classes 1 and 2 (Figure 2). When the maps were superimposed in such a way as to produce a maximal correlation of the 30S bodies, a clockwise rotation of the head toward the E site of the subunit in the 30S IC was found, compared to the 30S mRNA complex (Figure 2A). Measured by the position of the globular domain of ribosomal protein S13 as a reference point, the displacement was about 10 Å, corresponding to a rotation of the 30S head of approximately 4–5°. The direction and extent of the rotation in the 30S IC map are very similar to those found in the 70S IC (Figure 2B) [12], as opposed to the conformation of the 30S subunit in the map attributed to the 30S mRNA complex (Figure 2C).

IF2 Structure and Interactions in the 30S IC

The crystal structure of full-length E. coli IF2 is not available so far. To interpret the density attributed to IF2 and to identify the interactions of IF2 within the 30S IC, a domain homology model of IF2 was constructed (Figure 3). The C-terminal half of the protein (CTD; G2, G3, C1, C2 domains) is highly conserved, whereas the N-terminus (NTD; N1, N2, G1 domains) varies in both amino acid composition and length and is absent in archaea [32]. The NTD is lacking in IF2/eIF5B from Methanobacterium thermoautotrophicum, the crystal structure of which is available [33] and which has been used for homology modeling of IF2 from T. thermophilus and E. coli [11,12]. To model the full-length E. coli IF2, including the NTD which was not visualized previously, we combined several tertiary structures based on sequence homology (Figure S2). Rigid-body fitting was carried out independently for each of the modeled homologous domains. Subsequently, all of them were combined into a single structural model, which was fitted into the density of the 30S IC using MD-based flexible fitting (Materials and Methods). In the MD simulations, secondary structure elements were conserved, and most of the large changes are related to long loops, especially in the NTD region. During the flexible fitting the cross-correlation between atomic coordinates and the EM map improved significantly (from 0.87 to 0.95 estimated at the current resolution). The evolution of the fitting...
was also monitored by comparison with the initial IF2 model, and in the last cycles the RMSD values between the initial and flexible-fitted models reached a plateau of 8 Å. The overall arrangement of domains in the CTD of the present model of IF2 (Figure 3) is in agreement with the previous models based on cryo-EM [11–13], except for some differences that are discussed below. The expected volume of the modeled NTD domains (irrespective of the uncertainties of their modeled positions) nicely accounted for the observed cryo-EM density indicating the location of the NTD at the 30S subunit surface.

In the 30S IC, IF2 makes extensive contacts with 16S rRNA (helices h5 and h14), IF1, and S12 (Figure 4). The model suggests that the contacts to IF1 and ribosomal protein S12 are made by the NTD of IF2 (Figure 4A), in line with the ability of the isolated NTD of E. coli IF2 to bind to the 30S subunit [9,34]. Domain G3 of IF2 interacts with helix h5 of 16S rRNA and domain C1 of IF2 with

Figure 2. Conformation of the 30S subunit within the 30S IC. (A) 30S subunit conformations in reconstructions of particles from class 1 (red mesh) and class 2 (semitransparent yellow). The arrow indicates the clockwise rotation of the 30S subunit head. The inset shows the positions of protein S13 in the two maps. (B) The current 30S IC map (class 2 after ML3D classification) aligned with the cryoEM map for the 70S IC (EM database code 1249 [12]) in two distinct orientations. (C) Comparison of the 30S-mRNA map (class 1 after classification) with the 70S IC map [12]. The alignment between density maps was performed by maximum overlap of the body of the 30S subunits.

doi:10.1371/journal.pbio.1001095.g002
IC does not interfere with the binding of the 50S subunit to the IF2 contact, suggesting that this interaction of IF2 with the 30S (highlighted orange in Figure 4B) are on the face of the helix opposite the residues involved in the connection with the 50S subunit of IF2 is bridge B8, which is established via helix h14 [35]. However, upon 50S subunit joining (Figure 4C). The only bridge in the vicinity present 30S IC does not interfere with intersubunit bridges formed showing the fitted IF2 (ribbon representation). For fMet-tRNAfMet (pdb code 2FMT; [31]), a rigid-body fitting was performed. Domains G2 (orange), G3 (green), C1 (grey), and C2 (purple) of IF2 are indicated. The NTD was modeled as three sub-domains (yellow, blue, and pink). (B) Domain structure of IF2. Colors are as in (A). Below the segment for each domain, the homology T-Coffee score (72) between IF2 from E. coli and the atomic coordinates used is shown.

doi:10.1371/journal.pbio.1001095.g003

Figure 3. Homology model for IF2 guided by the cryo-EM map. (A) Close-up view of the fMet-tRNAfMet-IF2 complex on the 30S IC. Densities for fMet-tRNAfMet and IF2 were rendered semitransparent to show the fitted IF2 (ribbon representation). For fMet-tRNAfMet (pdb code 2FMT; [31]), a rigid-body fitting was performed. Domains G2 (orange), G3 (green), C1 (grey), and C2 (purple) of IF2 are indicated. The NTD was modeled as three sub-domains (yellow, blue, and pink). (B) Domain structure of IF2. Colors are as in (A). Below the segment for each domain, the homology T-Coffee score (72) between IF2 from E. coli and the atomic coordinates used is shown.

helix h14 (Figure 4B). This set of connections between IF2 and the 30S subunit is similar to the one described in the 70S IC [12], whereas in the 30S IC lacking IF3 the contacts between IF2 and the 30S subunit were seen as small connections involving G3 domain from IF2 and helices h5 and h14 from the 16S rRNA [11]. Interactions with IF1 or S12 were not observed, probably due to the shortened NTD in IF2 from T. thermophila. The position of IF2 on the present 30S IC does not interfere with intersubunit bridges formed upon 50S subunit joining (Figure 4C). The only bridge in the vicinity of IF2 is bridge B8, which is established via helix h14 [35]. However, the residues involved in the connection with the 50S subunit (highlighted orange in Figure 4B) are on the face of the helix opposite to the IF2 contact, suggesting that this interaction of IF2 with the 30S IC does not interfere with the binding of the 50S subunit.

Novel P/I Position of fMet-tRNAfMet Met

Comparison of the fMet-tRNAfMet position in the present 30S IC reconstruction with crystal structures of ribosomes with tRNAs in A, P, and E sites (Figure 5) reveals that the anticodon loop of fMet-tRNAfMet is positioned essentially in the P site of the 30S subunit, while the tRNA elbow in the 30S IC is tilted approximately 10° towards the E site (Figure 5A). The tilt is similar to that in the P/I intermediate position for the initiator tRNA previously described in cryo-EM studies of the 30S IC and 70S IC [11,12]. However, unlike the P/I state visualized before, where the acceptor arm of the tRNA was shifted towards the E site [11,12], the CCA end of fMet-tRNAfMet in our reconstruction is orientated towards the A site by its interaction with the C2 domain of IF2 (Figure 5B). The differences in the orientation are clearly seen at the junction between fMet-tRNAfMet and the C2 domain of IF2 in the respective complexes (Figures 5C,D and S3). Notably, the present 30S IC and the 70S IC reported by [12] have been prepared in the presence of all three IFs and are from the same organism and yet the observed P/I positions are clearly distinct (Figure 5C,D). To distinguish between the two states of the initiator tRNA, we name the state observed in our structure P/I1. Compared to the P-site tRNA, the P/I1 position requires a rotation of the acceptor stem of the tRNA of around 15° (Figure 5).

Location of IF3 in the 30S IC

Apart from the density for the fMet-tRNAfMet-IF2 complex, a bilobed density was observed connecting the 30S platform and the elbow region of the fMet-tRNAfMet (Figure 1E). Subtracting the volume occupied by the 30S-fMet-tRNAfMet-IF2 complex from the 30S IC reveals a density comprising two domains connected by a linker. According to its size and position, the density was attributable to IF3 (Figure 6). Rigid-body fitting of the atomic coordinates of the N- (IF3N) and C-terminal (IF3C) globular domains of IF3 from Geobacillus stearothermophilus [36] linked by an α-helix yielded a good fit of the EM map. For detailed fitting, IF3C from E. coli was used [37]. The density for IF3 in our cryo-EM map had to be visualized at a lower threshold compared to that of tRNA-IF2 (Figure S4), presumably due to incomplete occupancy of the 30S IC with IF3. Further sorting of the class 2 particles did not separate distinct conformational/occupancy states. The two domains of IF3 were placed in such a way that IF3N contacted the elbow region of Met-tRNAfMet, whereas IF3C was engaged in interactions with the 30S subunit. Cross-correlation measurements support the assignment of IF3 domains, since the alternative arrangement after domain swapping reduces the coefficient from 0.79 to 0.65. IF3C was bound to the 790 loop of h24 of 16S rRNA in the vicinity of fMet-tRNAfMet; however, no direct contact between IF3C and tRNA was found (Figure 6B).

Implications for Subunit Joining

To examine whether the conformation of the 30S IC was suitable for 50S subunit joining, we aligned our 30S IC map with that of the 70S IC [12]. The position for IF3C in the 30S IC would impair the formation of bridge B2b at the interface between the ribosomal subunits (Figure 7A), suggesting that in this arrangement IF3 in the complex physically impairs subunit joining. The fMet-tRNAfMet-IF2 complex provided an extensive surface area for the interaction with the 50S subunit (Figure 7B). Binding of the 50S subunit would position the junction between IF2 and the CCA end of fMet-tRNAfMet close to the peptidyl transferase center. The region containing the sarcin-ricin loop (SRL) of 23S rRNA docked accurately in a cleft formed by domains G2 and C1 of IF2. The GTPase domain, G2, is oriented toward the SRL in the same way as domain 1 of EF-Tu [38]; the contact with the SRL is expected to be important for the GTPase activation of IF2 [39].

Discussion

Structure of the 30S IC

In this study we present the cryo-EM reconstructions of the complete E. coli 30S IC containing initiator tRNA, mRNA, and all three initiator factors, as well as of the 30S subunit in the complex with the mRNA alone. The comparison of the 30S IC and the 30S-mRNA complex revealed that the 30S subunits were present in different configurations in the two complexes. In the 30S IC, the head of the 30S subunit was rotated with respect to the body, which can be attributed to the presence of IFs and fMet-tRNAfMet. Early site-directed crosslinking studies suggested that the formation of the SD-ASD complex in the absence of IFs places the
**Figure 4. IF2 on the 30S IC.** (A) Interactions of the NTD of IF2 with IF1 and S12. Thumbnail shows orientation. (B) Interactions of domains G2 and G3 of IF2 with 16S rRNA. Residues of helix h14 involved in formation of the intersubunit bridge B8 are highlighted in orange. (C) Position of IF2 relative to intersubunit bridges. The binding site for IF2 entails 16S rRNA helices h5 and h14 (green), and proteins IF1 (blue) and S12 (pink). The crystal structure of IF1 bound to the 30S subunit is taken from [pdb code: 1HRO; [7]]. Intersubunit bridges are shown in orange. Bridges in the head of the 30S subunit are highlighted in red and the arrow indicates the direction of the 30S subunit head movement.

doi:10.1371/journal.pbio.1001095.g004

**Figure 5. Orientation of fMet-tRNAfMet bound to the 30S IC.** (A) Comparison of the positions of fMet-tRNAfMet in the 30S IC (red) with the atomic coordinates for A-, P-, and E-site tRNAs (pdb code: 2HGP; [41]). The arrow indicates a tilt of the fMet-tRNAfMet by around 10 degrees compared to the tRNA in the P site. (B) Top view showing the 15 Å rotation of fMet-tRNAfMet in the complex with IF2 (semitransparent green). (C and D) Comparison between the P/I1 and P/I sites. In (C) densities for IF2 and tRNA from the 30S IC are shown semitransparent, allowing us to visualize fitted coordinates for tRNA (red ribbons) and IF2 (green). For comparison, coordinates fitted in the 70S IC (pdb code: 1ZO1 [12]) are represented for IF2 (purple ribbons) and tRNA in the P/I site (blue). In (D) density for IF2 and tRNA is taken from the 70S IC (EM database code 1249) and depicted semitransparent. Atomic coordinates are as in (C).

doi:10.1371/journal.pbio.1001095.g005
mRNA in a “standby” position, from which it is shifted backward, closer to the P site upon binding of initiation factors, in particular IF3, before the interaction with the large subunit takes place [40]. Similarly, crystal structures of the 70S complexes indicated that mRNA moves in the 3′→5′ direction with simultaneous clockwise rotation and lengthening of the SD duplex, bringing it into contact with ribosomal protein S2 [41]. The conformational change in the 30S IC induced by factor binding moves the head of the subunit in the right direction along the pathway of the mRNA and could promote the reported back-tracking of the messenger. At the current resolution, the conformation of the 30S subunit in the 30S IC is the same as in the 70S IC before GTP hydrolysis by IF2, where the 30S subunit is found in a rotated (ratcheted) orientation with respect to the 50S subunit [12] when compared with a 70S post-initiation complex [42]; it should be noted that the origin of initiation components (E. coli) and the mRNAs (extended SD sequence) are very similar in the present study and in [12], allowing for such detailed comparisons. The change in the 30S configuration is similar to that described within other 70S complexes along several steps of translation (see Figure S5).

Figure 6. IF3 on the 30S IC. (A) Density for IF3 (semitransparent orange) fitted using the atomic coordinates for IF3 domains, IF3N (pdb code: 1TIF; [36]) and IF3C (pdb code: 2IFE; [37]). Thumbnail shows the orientation. (B) Same orientation as in (A) with semitransparent 30S subunit. Regions on 16S rRNA and tRNA in the proximity of IF3, based on hydroxyl radical footprinting [17], are shown in navy blue. Arrow indicates helix h24 and the loop around nucleotide 790 in 16S rRNA.

doi:10.1371/journal.pbio.1001095.g006

Figure 7. Docking of the 50S subunit onto the 30S IC. (A) Steric clash between IF3C and helix H69 of 23S rRNA at intersubunit bridge B2b. The present map for the 30S IC was aligned with the 70S IC [12]. (B) IF2 as an important determinant for the 50S subunit association. The position of the sarcin-ricin loop (SRL) at the cleft formed by domains G2 and C1 of IF2 is indicated. In both panels, the thumbnails show the 30S IC docked onto the 50S subunit. Labels on the 50S subunit indicate helices in 23S rRNA (H69, H89, and H95) and the stalk region for protein L1.

doi:10.1371/journal.pbio.1001095.g007
[30, 43–52], but in the current study, the 30S conformation does not require the presence of the 50S subunit. Conformational changes of the 30S subunit were not described in the 30S initiation complex lacking IF3 [11]. This suggests that binding of IFs, in particular IF3, could induce or stabilize the altered 30S subunit conformation. This conformational state of the 30S subunit appears to be retained upon 50S subunit joining until GTP hydrolysis and dissociation of IF2 [26].

The positions of fMet-tRNAfMet and IF2 in the complete 30S IC are similar to, but not identical with, those found in the previously reported reconstructions. As in all available structures, the anticodon stem of the tRNA is buried in the P site of the 30S subunit. However, the position of the tRNA CCA end differs in the reported P/I states. Comparing the 30S IC with the 70S IC from E. coli [both with GDPNP] suggests that, upon binding of the 50S subunit, the orientation of the CCA end of the tRNA changes from the P/I state, pointing towards the A site of the peptidyl transferase center (30S IC; this article), towards the P/I state, which resides between the P and E sites (70S IC; [12]). The position of the C2 domain of IF2 to which the fMet moiety of fMet-tRNAfMet is bound changes accordingly. A somewhat different P/I state was reported for the 30S IC from T. thermophilus formed with GTP in the absence of IF3 [11], which may reflect the known effect of IF3 on the stability of the tRNA binding to the 30S IC. Alternatively, different P/I states may reflect the flexibility of the CCA end of the tRNA. It is possible that all described P/I positions can be sampled during the transition of the initiation tRNA toward the final P site and are important for discrimination of mRNAs with unfavorable TIR (see below).

The density in the cryo-EM map accounted for the entire IF2 and allowed us to map the domain contacts of the factor. As expected, the C2 domain of IF2 binds fMet-tRNAfMet at the single-stranded acceptor end and the fMet moiety of fMet-tRNAfMet is bound changes accordingly. A somewhat different P/I state was reported for the 30S IC from T. thermophilus formed with GTP in the absence of IF3 [11], which may reflect the known effect of IF3 on the stability of the tRNA binding to the 30S IC. Alternatively, different P/I states may reflect the flexibility of the CCA end of the tRNA. It is possible that all described P/I positions can be sampled during the transition of the initiation tRNA toward the final P site and are important for discrimination of mRNAs with unfavorable TIR (see below).

The density in the cryo-EM map accounted for the entire IF2 and allowed us to map the domain contacts of the factor. As expected, the C2 domain of IF2 binds fMet-tRNAfMet at the single-stranded acceptor end and the fMet moiety of fMet-tRNAfMet is bound changes accordingly. A somewhat different P/I state was reported for the 30S IC from T. thermophilus formed with GTP in the absence of IF3 [11], which may reflect the known effect of IF3 on the stability of the tRNA binding to the 30S IC. Alternatively, different P/I states may reflect the flexibility of the CCA end of the tRNA. It is possible that all described P/I positions can be sampled during the transition of the initiation tRNA toward the final P site and are important for discrimination of mRNAs with unfavorable TIR (see below).

The density in the cryo-EM map accounted for the entire IF2 and allowed us to map the domain contacts of the factor. As expected, the C2 domain of IF2 binds fMet-tRNAfMet at the single-stranded acceptor end and the fMet moiety of fMet-tRNAfMet is bound changes accordingly. A somewhat different P/I state was reported for the 30S IC from T. thermophilus formed with GTP in the absence of IF3 [11], which may reflect the known effect of IF3 on the stability of the tRNA binding to the 30S IC. Alternatively, different P/I states may reflect the flexibility of the CCA end of the tRNA. It is possible that all described P/I positions can be sampled during the transition of the initiation tRNA toward the final P site and are important for discrimination of mRNAs with unfavorable TIR (see below).

The density in the cryo-EM map accounted for the entire IF2 and allowed us to map the domain contacts of the factor. As expected, the C2 domain of IF2 binds fMet-tRNAfMet at the single-stranded acceptor end and the fMet moiety of fMet-tRNAfMet is bound changes accordingly. A somewhat different P/I state was reported for the 30S IC from T. thermophilus formed with GTP in the absence of IF3 [11], which may reflect the known effect of IF3 on the stability of the tRNA binding to the 30S IC. Alternatively, different P/I states may reflect the flexibility of the CCA end of the tRNA. It is possible that all described P/I positions can be sampled during the transition of the initiation tRNA toward the final P site and are important for discrimination of mRNAs with unfavorable TIR (see below).

The density in the cryo-EM map accounted for the entire IF2 and allowed us to map the domain contacts of the factor. As expected, the C2 domain of IF2 binds fMet-tRNAfMet at the single-stranded acceptor end and the fMet moiety of fMet-tRNAfMet is bound changes accordingly. A somewhat different P/I state was reported for the 30S IC from T. thermophilus formed with GTP in the absence of IF3 [11], which may reflect the known effect of IF3 on the stability of the tRNA binding to the 30S IC. Alternatively, different P/I states may reflect the flexibility of the CCA end of the tRNA. It is possible that all described P/I positions can be sampled during the transition of the initiation tRNA toward the final P site and are important for discrimination of mRNAs with unfavorable TIR (see below).

The density in the cryo-EM map accounted for the entire IF2 and allowed us to map the domain contacts of the factor. As expected, the C2 domain of IF2 binds fMet-tRNAfMet at the single-stranded acceptor end and the fMet moiety of fMet-tRNAfMet is bound changes accordingly. A somewhat different P/I state was reported for the 30S IC from T. thermophilus formed with GTP in the absence of IF3 [11], which may reflect the known effect of IF3 on the stability of the tRNA binding to the 30S IC. Alternatively, different P/I states may reflect the flexibility of the CCA end of the tRNA. It is possible that all described P/I positions can be sampled during the transition of the initiation tRNA toward the final P site and are important for discrimination of mRNAs with unfavorable TIR (see below).

The density in the cryo-EM map accounted for the entire IF2 and allowed us to map the domain contacts of the factor. As expected, the C2 domain of IF2 binds fMet-tRNAfMet at the single-stranded acceptor end and the fMet moiety of fMet-tRNAfMet is bound changes accordingly. A somewhat different P/I state was reported for the 30S IC from T. thermophilus formed with GTP in the absence of IF3 [11], which may reflect the known effect of IF3 on the stability of the tRNA binding to the 30S IC. Alternatively, different P/I states may reflect the flexibility of the CCA end of the tRNA. It is possible that all described P/I positions can be sampled during the transition of the initiation tRNA toward the final P site and are important for discrimination of mRNAs with unfavorable TIR (see below).
Cryo-EM, Image Processing, and Image Classification
Thin carbon was floated onto Quantifoil grids (Quantifoil Micro Tools GMBH, Jena, Germany). A 3.5-μl aliquot of the sample was placed on each grid. Grids were blotted, plunged in liquid ethane, and stored in liquid nitrogen until data collection. Low-dose images were taken on Kodak SO-163 films in a JEM-2200FS electron microscope (JEOL) operated at 200 kV at a magnification of 50,000. Micrographs were scanned on a Zeiss 1 Photonscan scanner (Zeiss) with a step size of 14 μm. Immediately before grid preparation and vitrification, the mixture was diluted to 30 nM 30S IC with buffer A containing 0.5 mM GDPNP.

Modelling an Atomic Structure for IF2
Homology modeling was carried out using the Swiss-Model server [63]. Since for the entire E. coli IF2 no homolog of known structure was found, proteins and protein domains with the highest sequence similarity corresponding to domains and sub-domains of IF2 were searched by BLAST [64]. Parts of IF2 were modeled based on the respective homolog structure. Residues 1–50 were modeled using the N-terminal subdomain of IF2 from E. coli (pdb code 1ND9; [65]); residues 51–185 with the dynamin-like protein BDLP from S. solfataricus (pdb code 1G7S; [33]); residues 391–559 (corresponding to the G2 domain) and 560–672 (corresponding to the G3 domain) using the code 3CW2; [67]); residues 673–779 using the C1-subdomain of IF2 from B. stearothermophilus (pdb code 1Z9B; [68]); and residues 780–890 (C2 domain) based on the fMet-tRNA^Met-binding domain of IF2 from B. steaothermophilus (pdb code 1D1N; [69]). Independent rigid-body fitting of each modeled region into the IF2 density map was performed using Chimera [62]. The initial relative positions of domains was taken as described in the crystal structure for aIF5B [33], taking into account the interaction between domain C2 of IF2 and the initiator tRNA. Connecting residues were adjusted based on the maxima of the probability functions of the assembled model was carried out using the Swiss-PDB Viewer software [70]. Molecular dynamics-based flexible fitting of the assembled model was carried out by Flex-EM software [71].

Supporting Information

Figure S1  SD-ASD helix on cryo-EM maps. Cryo-EM map for class 1 (A) and class 2 (B) after ML3D classification. The maps are rendered semitransparent and fitted with crystal structure of 30S subunit in the complex with mRNA (pdb code: 1JGO; [5]). The SD-ASD helix from the crystallographic structure is shown in blue. Labels: S7 and S11 indicate positions of ribosomal proteins; h, head of the 30S subunit.

Figure S2 Sequence alignment of IF2 from E. coli with sequences of proteins used for homology modeling. PDB codes for the different atomic coordinates are indicated under the designation of IF2.

Figure S3 Comparison of the current 30S IC map with previous cryo-EM data from initiation complexes. (A) 30S IC with all three IFs (present work, yellow) aligned with the 30S IC from T. thermophilaus lacking IF3 (green; [11]). (B) 30S IC and the isolated density for tRNA-IFs extracted from the cryo-EM map of the 70S IC from E. coli (blue) [12]. Labels: h, head of the 30S subunit; pt, platform; sp, spur. Arrows point to the junction between fMet-tRNA^Met and IF2.

Figure S4 Visualization of the density attributed to IF3 at different thresholds. In panels (A) and (C) the cryo-EM map is depicted solid; in (B) and (D) the map is semitransparent to show the fitted atomic coordinates for IF3 domains: IF3N (pdb code: 1TIF; [36]) and IF3C (pdb code: 2IFE; [37]). The sigma values used for the rendering are indicated. Thumbnail shows orientation.

Figure S5 Comparison of the 30S conformation from class 1 (30S-mRNA) and class 2 (30S IC), both in semi-transparent renderings, with 30S subunits coming from 70S ribosomes from E. coli in rotated (red and solid) and non-rotated (green) states [52]. The conformation of the 30S in the 30S IC is closer to the rotated state. The alignment between density maps was performed by maximum overlapping in the body of the 30S subunits.

Acknowledgments
We thank Wolfgang Wintermeyer for critical reading the manuscript and Sandra Kappler, Astrid Bohm, and Carmen Schillings for expert technical assistance.

Author Contributions
The author(s) have made the following declarations about their contributions: Conceived and designed the experiments: PJ PM MVR MV. Performed the experiments: PJ PM DG. Analyzed the data: PJ PM XA GL MVR MV. Wrote the paper: PJ PM MVR MV.

Accession Numbers
The cryoEM maps for the 30S-mRNA complex (class 1) and for the 30S IC (class 2) have been deposited in the Electron Microscopy Data Bank, http://www.ebi.ac.uk/pdbe/emdb/ (accession numbers 1770 and 1771, respectively).

References
1. Gualerzi CO, Pon CL (1990) Initiation of mRNA translation in prokaryotes. Biochemistry 29: 3881–3889.
2. Laursen BS, Sorensen HP, Mortensen KK, Sperling-Petersen HU (2005) Initiation of protein synthesis in bacteria. Microbiol Mol Biol Rev 69: 101–123.
32. Steffensen SA, Poulsen AB, Mortensen KK, Sperling-Petersen HU (1997) *E. coli* RNA:structure analysis of the infB gene in clinical isolates of *E. coli*. FEBS Lett 419: 281–284.

33. Roll-Mecak A, Gao C, Dever TE, Burke SK (2000) X-Ray structures of the universal translation initiation factor IF2/eIF5B: conformational changes on GDP and GTP binding. Cell 103: 781–792.

34. Moreno JM, Kildsgaard J, Swianowicz I, Mortensen KK, Sperling-Petersen HU (1998) Binding of Escherichia coli initiation factor IF2 to 3OS ribosomal subunits: a functional role for the N-terminus of the factor. Biochim Biophys Acta 1352: 465–471.

35. Yusupova GZ, Yusupov MM, Baucom A, Lieberman K, Earnest TN, et al. (2001) Crystal structure of the ribosome at 5.3 Å resolution. Science 292: 883–886.

36. Biao V, Shu F, Ramakrishnan V (1995) X-ray crystallography shows that translation initiation factor IF3 consists of two compact alpha/beta domains bound by an alpha-helix. EMBO J 14: 4056–4064.

37. Garcia C, Fortier PL, Blanquet S, Lallement Y, Dardel F (1995) Solution structure of the ribosome-binding domain of E. coli translation initiation factor IF3. Homology with the U1A protein of the eukaryotic spliceosome. J Mol Biol 254: 247–259.

38. Schneid TM, Voorhees RM, Kelley AC, Gao YG, Murphy FV, et al. (2009) The crystal structure of the ribosome bound to EF-Tu and aminoacyl-tRNA. Science 326: 689–694.

39. Tomasic J, Vitali LA, Daviër T, Savelberghe A, Spurio R, et al. (2008) Late events of translation initiation in bacteria: a kinetic analysis. Embo J 19: 2121–2136.

40. La Teana A, Gualerzi CO, Brimaoue CF (1995) From stand-by to decoding site. Adjustment of the mRNA on the 30S ribosomal subunit under the influence of the initiation factors. Proc Natl Acad Sci U S A 92: 722–727.

41. Yusupova G, Jenner L, Rees B, Moras D, Yusupov M (2006) Structural basis for messenger RNA movement on the ribosome. Nature 444: 391–394.

42. Gabashvili IS, Agrawal KK, Spahn CM, Grassucci RA, Svergun D, et al. (2008) Solution structure of the E. coli 70S ribosome at 11.3 Å resolution. Cell 100: 537–549.

43. Agirrezabala X, Lei J, Brunelle JL, Ortiz-Meoz RF, Green R, et al. (2008) Visualization of the hybrid state of tRNA binding promoted by spontaneous recycling of the ribosome. Mol Cell 32: 190–197.

44. Agrawal KK, Sharma MR, Kid M, Hirokawa G, Booth TM, et al. (2004) Visualization of ribosome-recycling complex on the Escherichia coli 70S ribosome: functional implications. Proc Natl Acad Sci U S A 101: 18900–18905.

45. Athar MF, Elmasry HH, Kaur H, Ha TA (2000) Spontaneous intersubunit rotation in single ribosomes. Mol Cell 35: 578–588.

46. Ermolichenko DN, Majumdar ZK, Hickerson RP, Spiegel PC, Clegg RM, et al. (2007) Observation of intersubunit movement of the ribosome in solution using FRET. J Mol Biol 370: 530–540.

47. Frank J, Agrawal RK (2000) A ratchet-like inter-subunit reorganization of the ribosome during translocation. Nature 406: 318–322.

48. Gao H, Zhou Z, Rawat U, Huang C, Bouakaz L, et al. (2007) IF3 induces ribosomal conformational changes responsible for dissociation of class I release factor 2. Cell 129: 929–941.

49. Gao N, Zavialov AV, Li W, Sengupta J, Valle M, et al. (2005) Mechanism for the disassembly of the posttermination complex inferred from cryo-EM studies. Mol Cell 18: 663–674.

50. Klascho BP, Miyasikov AG, Van Heel M (2004) Visualization of release factor 3 on the ribosome during termination of protein synthesis. Nature 427: 802–805.

51. Stark H, Rodnina MV, Wieden HJ, van Heel M, Wintzermyer W (2000) Large-scale movement of elongation factor G and extensive conformational change of the ribosome during translocation. Mol Cell 10: 301–305.

52. Valle M, Zavialov A, Sengupta J, Rawat U, Ehrenberg M, et al. (2005) Locking and unlocking of ribosomal movements. Cell 114: 123–134.

53. Guemmegues M, Caserta E, Brandi L, Spurio R, Meunier S, et al. (2000) Mapping the IF3-tRNA(Neo)/Neo binding site of initiation factor IF3. Embo J 19: 5213–5240.

54. Kapoor S, Fabbritti A, Garulli C, Spurio R, Gualerzi CO, et al. (2008) Translation initiation factor IF1 of Bacillus stearothermophilus and Thermus thermophilus substitue for Escherichia coli IF1 in vitro and in vitro without a direct IF1–IF2 interaction. Mol Microbiol 70: 1369–1377.

55. Tapprich WE, Goss DJ, Dahlberg AE (1989) Mutation at position 791 in Escherichia coli 16S ribosomal RNA affects processes involved in the initiation of protein synthesis. Proc Natl Acad Sci U S A 86: 4927–4931.

56. McCarthy JF, Brimaoue CF (1994) Prokaryotic translation: the interactive pathway leading to initiation. Trends Genet 10: 402–407.

57. Chen H, Bjørknes M, Kumar R, Jay E (1994) Determination of the optimal aligned spacing between the Shine-Dalgarno sequence and the translation initiation codon of Escherichia coli mRNAs. Nucleic Acids Res 22: 4953–4957.

58. Ringquist S, Shanfold S, Barrick D, Green L, Bianley J, et al. (1992) Translation initiation in Escherichia coli: sequences within the ribosome-binding site. Mol Microbiol 6: 1219–1229.

59. Antoun A, Pavlov MV, Leppaniemi M, Ehrenberg M (2006) How initiation factors maximize the accuracy of tRNA selection in initiation of bacterial protein synthesis. Mol Cell 23: 185–193.

60. Baxter WT, Leith A, Frank J (2007) SPIRE: the SPIDER reconstruction engine. J Struct Biol 157: 56–63.

61. Scheres SH, Nunez-Ramirez R, Sorzano CO, Carazo JM, Marabini R (2008) Image processing for electron microscopy single-particle analysis using XMIPP. Nat Protoc 3: 977–990.
62. Pettersen EF, Goddard TD, Huang CC, Couch GS, Greenblatt DM, et al. (2004) UCSF Chimera—a visualization system for exploratory research and analysis. J Comput Chem 25: 1605–1612.
63. Arnold K, Bordoli L, Kopp J, Schwede T (2006) The SWISS-MODEL workspace: a web-based environment for protein structure homology modelling. Bioinformatics 22: 195–201.
64. Altschul SF, Madden TL, Schaffer AA, Zhang J, Zhang Z, et al. (1997) Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids Res 25: 3389–3402.
65. Laursen BS, Mortensen KK, Sperling-Petersen HU, Hoffman DW (2003) A conserved structural motif at the N terminus of bacterial translation initiation factor IF2. J Biol Chem 278: 16320–16328.
66. Low HH, Lowe J (2006) A bacterial dynamin-like protein. Nature 444: 766–769.
67. Stolboushkina E, Nikonov S, Nikolain A, Blasi U, Manstein DJ, et al. (2008) Crystal structure of the intact archaeal translation initiation factor 2 demonstrates very high conformational flexibility in the alpha- and beta-subunits. J Mol Biol 382: 680–691.
68. Wienk H, Tomaselli S, Bernard C, Spurio R, Picone D, et al. (2005) Solution structure of the C1-subdomain of Bacillus stearothermophilus translation initiation factor IF2. Protein Sci 14: 2461–2468.
69. Meunier S, Spurio R, Czisch M, Wechselberger R, Guenneugues M, et al. (2000) Structure of the tRNA(1Met)-binding domain of B. stearothermophilus initiation factor IF2. Embo J 19: 1918–1926.
70. Guex N, Peitsch MC (1997) SWISS-MODEL and the Swiss-PdbViewer: an environment for comparative protein modeling. Electrophoresis 18: 2714–2723.
71. Topf M, Lasker K, Webb B, Wolfson H, Chin W, et al. (2008) Protein structure fitting and refinement guided by cryo-EM density. Structure 16: 295–307.
72. Notredame C, Higgins DG, Heringa J (2000) T-Coffee: a novel method for fast and accurate multiple sequence alignment. J Mol Biol 302: 205–217.