Structure of the *E. coli* ribosome–EF–Tu complex at <3 Å resolution by *C*<sub>s</sub>–corrected cryo-EM

Niels Fischer<sup>1*</sup>, Piotr Neumann<sup>2*</sup>, Andrey L. Konevega<sup>3,4,5</sup>, Lars V. Bock<sup>6</sup>, Ralf Ficner<sup>2</sup>, Marina V. Rodnina<sup>5</sup> & Holger Stark<sup>1,7</sup>

Single particle electron cryomicroscopy (cryo-EM) has recently made significant progress in high-resolution structure determination of macromolecular complexes due to improvements in electron microscopic instrumentation and computational image analysis. However, cryo-EM structures can be highly non-uniform in local resolution<sup>1,2</sup> and all structures available to date have been limited to resolutions above 3 Å<sup>3,4</sup>. Here we present the cryo-EM structure of the 70S ribosome<sup>5</sup> (*E. coli*) in complex with elongation factor Tu, aminoacyl-tRNA and the antibiotic kirromycin at 2.65–2.9 Å resolution using spherical aberration (*C*<sub>s</sub>)-corrected cryo-EM. Overall, the cryo-EM reconstruction at 2.9 Å resolution is comparable to the best-resolved X-ray structure of the *E. coli* 70S ribosome<sup>5</sup> (2.8 Å), but provides more detailed information (2.65 Å) at the functionally important ribosomal core. The cryo-EM map elucidates for the first time the structure of all 35 rRNA modifications in the bacterial ribosome, explaining their roles in fine-tuning ribosome structure and function and modulating the action of antibiotics. We also obtained atomic models for flexible parts of the ribosome such as ribosomal proteins L9 and L31. The refined cryo-EM–based model presents the currently most complete high-resolution structure of the *E. coli* ribosome, which demonstrates the power of cryo-EM in structure determination of large and dynamic macromolecular complexes.

Determining the structure of large, dynamic biological macromolecules at a uniformly high resolution provides a challenge both for X-ray crystallography and cryo-EM. Here we have used aberration-corrected cryo-EM in combination with extensive computational sorting to solve...
the structure of the *E. coli* 70S ribosome at high resolution to visualize rRNA modifications and dynamic parts of the ribosome. We prepared the 70S *E. coli* ribosome in the codon recognition state with the cognate ternary complex EF-Tu–GDP–Phe–tRNAPhe stalled on the ribosome by the antibiotic kirromycin. Currently, a high-resolution crystal structure is available for the 70S–EF-Tu complex from *Thermus thermophilus* at 3.1 Å resolution; however, a comparable structure of the complex from *E. coli* is still lacking, which makes it difficult to integrate structures with the results of biochemical, biophysical and genetic experiments, most of which were obtained with *E. coli* ribosomes. Cryo-EM images were recorded in a 300 kV electron cryo-microscope equipped with a spherical-aberration corrector and a direct electron detector operated in integration mode without alignment of intermediate image frames for motion correction. The aberration corrector was specifically tuned to reduce resolution-limiting aberrations and distortions (Extended Data Fig. 1). We applied a hierarchical classification strategy to sort the ribosome images computationally for the known modes of ribosomal motion and potential sources of heterogeneity in ribosome preparations (Extended Data Fig. 2). The final cryo-EM map at 2.9 Å overall resolution was subsequently used to refine the atomic model for the entire ribosome, including metal ions (Mg$^{2+}$, Zn$^{2+}$) and rRNA modifications (Methods and Extended Data Table 1). The local resolution map is relatively uniform in resolution (Fig. 1a). For large portions of the map the resolution is better than 2.9 Å, whereas only few parts of the ribosome, located at the very periphery, are limited to resolutions >3.5 Å. Locally, average resolutions were 2.8 Å for 65%, 2.7 Å for 44% and 2.65 Å for 24% of the map, as determined by the crystallographic measures $F_{SC}/F_{work}$ (Fourier shell correlation) and $C_{work}$ (Pearson correlation coefficient) (Extended Data Fig. 3; see Methods for details). The final 3D map has a similar, or locally even better resolution compared to the available X-ray structures of the *E. coli* ribosome (Fig. 1 and Extended Data Fig. 4a). Also the structural definition of side chains as judged by local real-space correlations between the map and the model is similar to X-ray structures at comparable resolutions (Extended Data Table 2). In the best defined areas of the map Mg$^{2+}$ ions can be visualized along with water molecules in the coordination sphere, indicating an optical resolution of at least 2.8 Å (Fig. 1b).

The present cryo-EM map visualizes for the first time modifications in rRNA, which were not observable in any of the high-resolution X-ray structures of the bacterial ribosome from 2.4–2.8 Å resolution, and were only seen in high-resolution X-ray structures of the *Haloarcula marismortui* 50S subunit at 2.2–2.4 Å resolution (Extended Data Fig. 4). In contrast, even single methyl groups of nucleosides can be clearly visualized in our cryo-EM map (Fig. 1d, f), as well as the non-planar base of dihydrouridine at position 2449 of 23S rRNA (Fig. 1e), while pseudouridines could be identified indirectly by polar residues within hydrogen-bonding distance of the N1 position. In total, we were able to build all 35 constitutive rRNA modifications of the *E. coli* 70S ribosome.

The RNA modifications are clustered at the main functional centres of the ribosome (Fig. 2 and Extended Data Fig. 5). Clusters of several rRNA modifications are essential for ribosome function, whereas individual rRNA modifications have an important role in fine-tuning the active centres of the ribosome, as well as in antibiotic resistance and sensitivity. Six individual RNA modifications at the decoding centre modulate the efficiency and accuracy of translation initiation by stabilizing interactions in the P site and PTC. The present structure shows that the methyl group of m$^6$G966 in 16S rRNA may act as a ‘backstop’ for the initiator tRNA, thereby stabilizing base-pairing with the initiation codon (Fig. 2b and Extended Data Fig. 5a; see Extended Data Fig. 5b for definition of rRNA modifications). The binding platform generated by m$^6$G966 is further affected by base stacking interactions with the methyl group of m$^5$C967. Messenger RNA binding to the P site is stabilized by an intricate network of interactions that involves four modified nucleotides. The P-site codon directly contacts m$^6$Cm1402 and m$^6$U1498 in 16S rRNA, which in turn are held in place by the bulky dimethylamine groups on m$^6$A1519 and m$^6$A1518. The network of long-range interactions provides the basis for the action of the antibiotic kasugamycin, which binds in the P site and requires dimethylation of m$^2$A1519 for its function. In the A site of the decoding centre, the aminoglycoside class of antibiotics directly binds to a monomethylated residue, m$^7$C1407 in 16S rRNA, which is needed for optimum drug activity (Extended Data Fig. 5c).

Assembly and peptidyl-transferase activity of the 50S subunit require a network of six modified nucleosides at the entry site of the peptide exit tunnel (Fig. 2c). Some of these modifications also modulate antibiotic sensitivity of the ribosome, which can now be explained at a structural level. For instance, resistance against the antibiotic linezolid caused by the loss of methylation of m$^2$A2503 can arise from the lack of stabilization by the m$^2$A2503 methyl group on the stacking interaction with A2059, which keeps A2503 in a position that allows the antibiotic binding to the peptidyl-transferase centre. The stacking interaction of the A2503 methyl group with the base of A2059 stabilizes A2503 in a conformation compatible with binding of the antibiotic linezolid (magenta, superposition from PDB ID: 3CPW). The nitrogen in m$^2$G966 forms an additional hydrogen bond that stabilizes the nucleotide conformation.

**Figure 2** | Modifications of rRNA. a, Clusters of rRNA modifications (yellow and blue) at the functional centres of the ribosome. Initiator fMet-tRNAPhe in the P site is shown in green. PTC, peptidyl-transferase centre; DC, decoding centre. b, Close-up of the decoding centre. Methyl group of m$^6$C967 stacks onto the m$^6$G966 base. The methyl group of m$^6$G966 (yellow) restricts the mobility of the initiator tRNA (green). An interaction network of four modified nucleotides stabilizes the mRNA (orange) binding to the P site. c, Cluster of rRNA modifications in the peptidyl-transferase centre. The essential Gm2251 in 16S rRNA stabilizes the mRNA (orange) binding to the P site. d, Cluster of rRNA modifications in the peptidyl-transferase centre. The stacking interaction of the A2503 methyl group with the base of A2059 stabilizes A2503 in a conformation compatible with binding of the antibiotic linezolid (magenta, superposition from PDB ID: 3CPW).

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m^8A1618 and its direct environment have also been recently predicted as the most promising novel drug target by a computational approach^22.

Generally, obtaining uniform resolution for large macromolecules and especially for bound factors is difficult for cryo-EM and X-ray crystallography^23–25, because of mobility and/or occupancy problems. Owing to extensive computational sorting of images, our cryo-EM map is comparable to local resolution variations to crystallographic maps. Higher B factors in X-ray structures are also in line with structural elements exhibiting the highest mobility in molecular dynamics simulations (Extended Data Fig. 6), suggesting that structural dynamics is the prevailing factor limiting local resolution. In X-ray crystallography, those flexible regions may be resolved when stabilized by crystal packing interactions. In cryo-EM, however, computational sorting of images can be employed to improve the local resolution of dynamic structural features. In the cryo-EM map, the local resolution of the EF-Tu–GDP-kirromycin–Phe–tRNA^Phe complex was improved by sorting for ligand occupancy, resulting in a well-defined density at only slightly lower local resolution compared to the ribosome core. Notably, we find a clear density for the antibiotic kirromycin (Fig. 3a), underlining the power of computational sorting in cryo-EM. Whereas the overall architecture of the EF-Tu–GDP–kirromycin–Phe–tRNA^Phe complex is similar to that reported by X-ray crystallography^6,26, there are local differences in the conformation of the distorted tRNA in the A/T state, in the orientation of the catalytic His84 residue and in the interactions between domain 2 of EF-Tu and the 30S subunit (Extended Data Fig. 7). Importantly, a shift in domain 2 (residues 219 to 226) of EF-Tu upon ribosome binding appeared crucial for distortion of the 3' end of tRNA and, ultimately, GTPase activation^27. The cryo-EM structure shows a similar distortion of the tRNA, but different interactions of EF-Tu domain 2, indicating that the reported changes in domain 2 may not be essential for the mechanism of catalytic activation. The highly mobile protein L9 is stabilized in crystals in an extended conformation by contacts to a neighbouring ribosome in the crystal lattice (Fig. 3b). In cryo-EM structures, this stabilization is absent and usually only the amino-terminal domain of the protein is structurally well-defined. Nevertheless, we were able to build the complete model for the conformation of L9 on the ribosome in solution, which reveals the contacts of L9 to the 30S subunit (Fig. 3b, c and Extended Data Fig. 8a). Notably, both conformations

![Figure 3](image-url)

**Figure 3** | Structure of ribosome ligands and mobile elements. Overview of the ribosome–EF-Tu complex. Left, cryo-EM map filtered to the local resolution of the factor binding site (3.1 Å). EF-Tu (red), the antibiotic kirromycin (blue), Phe–tRNA^Phe in the A/T state (purple), fMet–tRNA^fMet in the P site (green) and deacylated tRNA^fMet in the E site (orange) are shown. Centre, comparison of the present structure with the X-ray structure of the kirromycin–Phe–tRNA^Phe complex (Extended Data Fig. 8). Notably, both conformations of the kirromycin–Phe–tRNA^Phe complex were improved by sorting for ligand occupancy, resulting in a well-defined density at only slightly lower local resolution compared to the ribosome core. Notably, we find a clear density for the antibiotic kirromycin (Fig. 3a), underlining the power of computational sorting in cryo-EM. Whereas the overall architecture of the EF-Tu–GDP–kirromycin–Phe–tRNA^Phe complex is similar to that reported by X-ray crystallography^6,26, there are local differences in the conformation of the distorted tRNA in the A/T state, in the orientation of the catalytic His84 residue and in the interactions between domain 2 of EF-Tu and the 30S subunit (Extended Data Fig. 7). Importantly, a shift in domain 2 (residues 219 to 226) of EF-Tu upon ribosome binding appeared crucial for distortion of the 3’ end of tRNA and, ultimately, GTPase activation^27. The cryo-EM structure shows a similar distortion of the tRNA, but different interactions of EF-Tu domain 2, indicating that the reported changes in domain 2 may not be essential for the mechanism of catalytic activation. The highly mobile protein L9 is stabilized in crystals in an extended conformation by contacts to a neighbouring ribosome in the crystal lattice (Fig. 3b). In cryo-EM structures, this stabilization is absent and usually only the amino-terminal domain of the protein is structurally well-defined. Nevertheless, we were able to build the complete model for the conformation of L9 on the ribosome in solution, which reveals the contacts of L9 to the 30S subunit (Fig. 3b, c and Extended Data Fig. 8a). Notably, both conformations...
of L9 observed in cryo-EM and crystals appear to be compatible with the binding of elongation factors to neighbouring ribosomes in polysomes as determined by cryo-electron-tomography27 (Extended Data Fig. 8c). Furthermore, we built the model for L31, which is missing in the available E. coli 70S X-ray structures, probably owing to its dynamic nature. L31 bridges the ribosome at the top of the 30S head and the central protuberance of the 50S subunit. Upon 30S subunit ratcheting, the linker region of L31 switches from an extended to a kinked conformation, while L31 maintains its interactions with both subunits (Extended Data Fig. 8b). The low occupancy of L31 can be explained by its flexible binding mode, which may also be important for the function of L31 as a Zn$^{2+}$ reservoir for the cell28.

In conclusion, our data shows that aberration-corrected cryo-EM allows dynamic macromolecular machines, such as the ribosome, to be visualized at a uniform resolution better than 3 Å with only small variations in local resolution. The ‘purification’ of electron microscopic images by computational sorting appears to be as powerful as the purification of ribosome conformation and composition during crystal growth in situs in local resolution. The ‘purification’ of electron microscopic images by computational sorting appears to be as powerful as the purification of ribosome conformation and composition during crystal growth in situ. The cryo-EM map visualizes bound water molecules, ions and RNA modifications, providing novel insights into ribosome and antibiotic function, and thereby contributing to an improved structural basis for the development of new antibiotics.

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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**METHODS**

**Complex preparation.** To obtain kirromycin-stalled *E. coli* ribosome–EF-Tu complexes, ribosomes from *E. coli* MRE 600, initiation factors (IF1, IF2, IF3), Met-tRNA\(^{\text{Met}}\), EF-Tu and Phe–tRNA\(^{\text{Phe}}\) were prepared as described\(^{-11,12}\). Prior to initiation, the mRNA (GGGAGAGGUAUAAUAGUGUCGACG; the AUG start codon coding for Met and UUC coding for Phe are underlined) was incubated with 0.1 mM EDTA for 90 s at 80 °C and shock cooled in an ice-water bath. 70S ribosomes (3 μM) were incubated with IF1, IF2, IF3 (4.5 μM), mRNA (15 μM), and \(\text{GTP}_0\) (0.755 Å (as determined by comparing the final 3D map with atomic models from X-ray crystallography). The C. correae corector, electron optical aberrations were corrected to residual phase errors of 45 μrad at scattering angles of 12 to 15 mrad (that is, less than 45 μrad phase error at 1.8 to 2.1 Å; Extended Data Fig. 1). Linear geometrical distortions were reduced to <0.1% using the tiltlesapole beam coils of the C. correae corrector. Furthermore, coma caused by the spot-scanning procedure was minimized using the usimagshift correction in the C. correae corrector alignment. Ribosome particle images were extracted in a fully automated manner using template-independent custom-made software (Cowpick, B. Busche and H.S., unpublished data). The 1,603,254 extracted particle images were corrected locally for the contrast-transfer function by classification and averaging and selected according to quality of powerspectra\(^{-14}\), that is, to show Thon rings better than 3.4 Å up to 2.4 Å. In all subsequent steps, the resulting 1,339,775 contrast-transfer-function-corrected particle images were used. First, the particle images were sorted into groups of particles according to: (1) 30S body rotation; and (2) ligand occupancy, using supervised classification by projection matching\(^{15}\) on the basis of a structural library of different ribosome complexes\(^{16}\) (Extended Data Fig. 2). Sorting in both steps was performed using low-pass-filtered reference maps and particle images binned to about 6 Å per pixel. Finally, 3D classification in RELION 1.2\(^{21}\) was used to obtain the final set of 417,201 particle images with bound elongation factor for the refinement to high-resolution (2.9 Å) using the ‘gold-standard procedure’ in RELION 1.2\(^{27}\) (Extended Data Fig. 3). A local resolution map computed from the two unprocessed half-maps by Resmap\(^{-2}\) revealed only few variations in local resolution over the entire ribosome complex (Fig. 1a). We obtained another cryo-EM map of a ratcheted ribosome with tRNAs in hybrid states and showing protein L31 in a distinct conformation by sorting particle images with 10° rotation angles according to ligand occupancy, as described above, resulting in a final homogeneous population of 8,073 particles, which was refined to 6.4 Å resolution (0.143 criterion) using the gold-standard procedure in RELION. Atomic fluctuations obtained from molecular dynamics simulations. The molecular dynamics simulation of the ribosome in a pre-translocation state (pre1a) presented earlier\(^{18}\) was extended to 2 μs. To identify the rigid core of the 50S subunit, the root mean square fluctuation (r.m.s.f.) of each atom was calculated using the program g_rmsf from the GROMACS simulation suite\(^{19}\) after alignment to the 50S subunit, omitting the first 0.5 μs. The rigid core was defined as all 50S atoms excluding those of the tail of the r.m.s.f. histogram starting at the point where the frequency drops to half of the maximum frequency (0.19 nm). Finally, the r.m.s.f. of all atoms after alignment to the rigid core was calculated (Extended Data Fig. 4). The method is well suited to identify the rigid core of the ribosome. Pseudo-crystallographic refinement and model building. For initial model building, the cryo-EM density map was sharpened by applying a B factor of –120 Å\(^2\), filtered to 3.1 Å resolution and masked using a pseudo bulk solvent envelope obtained by merging different versions of the cryo-EM map filtered at different frequencies (12.0 Å, 8.0 Å, 6.0 Å and 4.0 Å) with the RAVE package\(^{20}\) and Chimera\(^{21}\). Fourier transform of the masked cryo-EM density map to reciprocal space structure factors was performed using Crystallography and NMR System (CNS)\(^{22}\) employing phase significance blurring scale factors derived from CNS values in a resolution-dependent manner\(^{23}\). In detail, a modified CNS input file was used for the assignment of FOMs (figure of merit) estimated based on equation (1) for the map obtained from the full data set and equation (2) for the maps obtained from two half sets:

\[
FOM = \frac{2 \times FSC}{FSC + 0.5}
\]

\[
FOM = \frac{FSC}{0.5}
\]

Obtained phase probabilities, written in the form of Hendrickson–Lattman (HL) coefficients, were used for reciprocal space refinement, performed against an MLHL target (maximum likelihood with experimental phase probability distribution) in both CNS and PHENIX\(^{24}\) programs using both X-ray and electron scattering factors, respectively. Both programs employed automatic optimization of weights used to balance the relative contributions of experimental and restraints terms using a grid search. To fulfill the requirements of the crystallographic MLHL refinement, 95% of the reflections were selected randomly for the ‘Rfree’ set, which was kept identical for all refinements. Homology modeling combined with density-guided energy optimization was performed using the Rosetta package\(^{25,26}\)–employing templates and alignments provided by the HHpred server\(^{27}\). Model density maps were generated based on finally refined models, without bulk solvent correction as implemented in PHENIX. Map normalization (mean and standard deviation of density values are 0 and 1, respectively) was performed using MAPPMAP (Rave package).

The initial fit of an atomic model of the *E. coli* ribosome assembled from various crystal structures (PDB codes: 4GD2 (30S), 3RBT (50S), 2B00 (mRNA, tRNA\(^{\text{Met}}\)), 3LU1 (tRNA\(^{\text{Phe}}\)), 1OB2 (EF-Tu) (R. C. Nielsen et al. unpublished data)) was performed using Chimera, followed by rigid body refinement in the PHENIX program. The atomic model was refined with deformable elastic network (DEN) restraints\(^{-28}\) in CNS with alternating cycles of manual rebuilding in Coot\(^{29}\) and monitoring the local fit to the density with RESOLVE\(^{-30}\). In addition the overall refinement process was monitored by calculating C\(_{\text{run}}\) and C\(_{\text{free}}\) (ref. 8), as well as the correlation between the cryo-EM and the model map (FOM\(_{\text{corr}}\)). The one-dimensional structure factor derived by rotational averaging from the initially refined model was used to re-sharpen the raw cryo-EM density map. Prior to Fourier transformation, the re-sharpened, normalized 3D cryo-EM map was solvent flattened using a smoothed model-based envelope (Rave package, Chimera), encompassing the volume within the distance of at least 3 Å from each atom. As the density in the protein region is not supposed to be negative, voxels with negative density in the protein region were set to zero during the solvent flattening process by the MAPMASK program in the CCP4 suite\(^{31}\). Further model improvement and fitting were facilitated by real space refinement (ERRASER\(^{32}\) and phenix.real_space_refine) against the map calculated using the working set of reflections only) and manual corrections in Coot combined with reciprocal space refinement. Modelling of post-transcriptional modifications was performed in Coot and was based on thorough analysis of the cryo-EM map. Modifications resulting from addition or substitution of an atom or atoms in comparison with unmodified bases were modelled if the presence of additional atoms or consequent changes in shape (for dihydrouridine) were supported by the cryo-EM map. Pseudouridines were modelled as indicated by additional polar/hydrogen-bond interactions formed by the additional amine group. For one methylated (m\(^7\)G247 in 23S rRNA) showing no well-defined corresponding density and two pseudouridines (m\(^7\)F46 and m\(^7\)F2457 in 23S rRNA) with no clear additional interactions, modifications were modelled on the basis of ref. 11 and references therein. In all other cases manual modelling of modified nucleotides was performed only if the cryo-EM map was well defined. New stereochemistry definitions for non-standard ligands were generated with phenix.real. A homology model of protein L31 was built in Rosetta using the *T. thermophilus* 23S rRNA (PDB ID: 3J01)\(^{35}\) and constraints from the present cryo-EM model filtered to 4.5 Å in resolution. Models of protein L9 and L31 were manually adjusted to fit the cryo-EM map filtered at 3.9 Å and 4.5 Å resolution, respectively in the programs O (ref. 39) and Coot. The models were further refined in real space using Rosetta followed by phenix.real_space_refine against the map created from working set reflections only. To maintain the intermolecular interactions of L9 and L31 proteins, the atomic models used for real space refinement in phenix.real_space_refine included surrounding proteins and parts of RNA chains within a radius of at least 10 Å. Improved protein density maps were included in the overall model used for the final reciprocal space refinement against 2.9 Å resolution data generated from the sharpened and solvent-flattened 3D cryo-EM map in PHENIX. The final model consisting of 152,718 individual atoms has been refined to 24.08% and 0.922 for Bamfree and C\(_{\text{run}}\) (definition is given below), respectively. The final model exhibits a good stereochemistry
with 85.59% of residues in the most favoured region and 3.30% residues in the disallowed region of the Ramachandran plot, protein side chain outliers of 2.63% and all atom clash score 9.15. Detailed refinement statistics are presented in Extended Data Table 1. The model for another conformer of protein L31 found in a cryo-EM map of the ribosome in complex with two tRNAs in hybrid states was built at 6.5 Å resolution, as described above for L19 and L31.

**Refinement of half maps.** Refinement of the final model against data sets obtained from two half maps was performed at a resolution of 2.9 Å in PHENIX using five to seven cycles of combined positional (real and reciprocal space) and atomic displacement parameter (ADP) refinement combined with automatically identified TLS (translation/libration/screw) groups. To remove possible model bias from the model refined against reflections obtained from the reconstruction using all the particles, a similar strategy was used as for higher-resolution refinements (a 0.5 Å random shift and additional restraints), as described below. The FSC and CC were calculated between the model and the half map used for refinement, as well as between the model and the other half map for cross-validation (Extended Data Fig. 3b).

**Higher-resolution refinements.** The final atomic model of the E. coli ribosome was divided into three sub-models containing about 65%, 43% and 24% of all residues, respectively, with each sub-model exhibiting similar B factors and resolution estimates for the corresponding fragment of the cryo-EM map (Extended Data Fig. 3). The resulting three models were used to calculate smoothed models encompassing the volume of at least a dimer of each atom. Those were used to cut the required portion of the cryo-EM density map using a pseudo bulk solvent flattening procedure before conversion to reciprocal space structure factors, as described above. Phase significance blurring scale factors applied during the Fourier transform were derived from FSC values calculated between fragments of the two half maps masked by model-based envelopes. The refinement of individual models (initially to 2.75 Å, 2.60 Å and 2.50 Å for 65%, 43% and 24% of all residues, respectively) against MLHL targets were performed in PHENIX by gradually decreasing the high-resolution limit in 0.05 Å steps, for example, local resolution refinements starting with 2.50 Å were performed at 2.50 Å, 2.55 Å, 2.60 Å, 2.65 Å, 2.70 Å and 2.75 Å. To eliminate possible bias against the model refined at 2.9 Å resolution, the atoms were displaced by a random translation up to 0.1 Å before performing the full refinement using restraints for secondary structure base-pairing, base-planarity and hydrogen bonding. The pseudo crystallographic high-resolution limit was selected based on several criteria: (1) the overall R and Rwork factors (lower than 1/10 of the highest resolution limit); (2) the Pearson correlation coefficient calculated between F_uncont and F_out (CCoverall) used for refinement should be greater than 0.2 for the highest resolution shell and the overall correlation coefficient (CCoverall) should not be lower than 0.9; (3) the calculated FSCwork value between model map coefficients (F_uncont, Fout, CCoverall) and structure factors derived from the cryo-EM map (F_uncont, Fphason, CCoverall) used for refinement should be greater than 0.5 for the highest resolution shell; (4) the calculated crystallographic R factor for the highest resolution shell should be not greater than 51%. The mentioned statistical values have been calculated in a resolution-dependent manner using PHENIX and SCALF (CCP4 suite) for 20 shells comprising a similar number of reflections (default number of shells for reporting refinement statistics in PHENIX).

**Preparation of figures.** Figures were generated using Pymol 1.7 (Schrödinger, LLC.), UCSF CHIMERA 1.9⁹, AMIRA 5.2.2 (Visage Imaging, Richmond VIC) and PRISM 5 (GraphPad Software, La Jolla).

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Extended Data Figure 1 | Aberration-corrected cryo-EM. a, Exemplary Zemlin tableau (left) and phase diagram (right) as obtained for the present data set with the CEOS software by correcting electron optical aberrations using the C₃ corrector. The resulting phase errors were less than 45° at ≤2.1 Å (that is, at scattering angles of 12 to 15 mrad) over up to 36 h of image acquisition. The main limiting aberration is axial coma (B₂) and the next limiting aberration would be threefold astigmatism (A₃). b, Local correction for the contrast transfer function. From micrographs (left) areas with individual ribosome particles (yellow frames) were extracted and local power spectra were computed for each of these areas by fast Fourier transform algorithms (FFT). Local power spectra were subjected to principal component analysis (PCA) and classification to average power spectra with similar contrast transfer function parameters that were obtained from different micrographs. Class averages of power spectra reveal an improved signal-to-noise ratio in Thon rings which are clearly visible up to 2.4 Å (right). c, Global power spectrum from a single micrograph showing Thon rings up to 3.5 Å.
Extended Data Figure 2 | Hierarchical sorting of ribosome particle images.
Ribosome particles were sorted in three steps according to: (1) global ribosome conformation (C1), that is, 30S body rotation; and (2) ligand occupancy (C2) and particle quality (C3)(Methods). The asterisk denotes particles assigned to the largest 30S body rotations $\leq -10^\circ$ and $\geq 10^\circ$ which contain particles with extreme 30S rotation angles, but also low-quality particle images.
Extended Data Figure 3 | Resolution curves and model validation of the E. coli 70S ribosome–EF-Tu cryo-EM structure. a, Fourier-shell correlation (FSC) curve (black) for the 70S ribosome cryo-EM reconstruction computed between the masked independent half-maps (half1 and half2) that were obtained by so-called ‘gold-standard’ refinement in RELION. The resolution of the cryo-EM reconstruction is 2.9 Å according to the 0.143 criterion (black dashed line). b, FSC curves computed between cryo-EM maps and model maps generated from refined atomic coordinates. The vertical black dashed line indicates the maximum resolution at which the full atomic models were refined. Black, the FSC curve between the final cryo-EM map (map) and the final model (model); blue, the FSC curve between half map 1 (half1) and the model obtained by refinement only against half map 1 (model1); red, the FSC curve between half map 1 and the model obtained by refinement only against half map 2 (model2). c, FSC curves (FSCwork) between reflections from solvent-flattened cryo-EM map and model as obtained by pseudo-crystallographic refinement of the complete ribosome model (mask1) and three sub-models using different masks corresponding to local variations in resolution (mask2–4; Methods) as shown in h. Coloured numbers indicate the highest resolution used in refinement with the respective mask as indicated by the colour code. For all refinements, the FSC is above the 0.5 threshold (black dashed line) in the highest-resolution shell. Differences to b result largely from solvent-flattening before Fourier transformation for refinement (Methods). d–g, CCwork and Rwork as obtained by refinement using the respective mask (see labels). For a reliable resolution estimate CCwork (ref. 8) is expected to be >0.2 and Rwork <0.51 in the highest-resolution shell. h, Isosurface representations of the mask used for local refinements; % indicates the fraction of atoms of the complete model entailed in the refinement with the respective mask.
Extended Data Figure 4 | Modifications in rRNA. Comparison between cryo-EM and X-ray crystallography. a, Experimental densities. In each row density maps for the same type of rRNA modification are shown (from left to right): for the present cryo-EM map and for the current best resolved bacterial and archaeal ribosome maps determined by X-ray crystallography, that is, the bacterial 70S ribosome from E. coli (Eco70S) at 2.8 Å resolution (PDB IDs: 4TPA and 4TPB); the bacterial 70S ribosome from T. thermophilus (Tth70S) at 2.4 Å resolution (PDB IDs: 4RB5 and 4RB6); and the archaeal 50S subunit from Haloarcula marismortui (Hma50S) at 2.2 Å resolution (PDB ID: 1VQ0). E. coli numbering is used for bacterial ribosome structures. Locations of rRNA modifications as determined by biochemical data are marked by yellow circles, modifications not observed in the density maps are denoted by red arrows and the black arrow designates the non-planarity of dihydrouridine observed in the cryo-EM map.

b, Model-based densities for m^6^2A1518 and m^6^2A1519 showing slight differences due to scattering properties. Densities were computed in CCTBX at 2.65 Å resolution from our final model with atomic-displacement factors kept unchanged using electron (e^- scattering, purple) and X-ray scattering factors (X-ray scattering, blue), respectively. Map thresholds were normalized to show similar density levels for the electron-rich phosphate groups. Accordingly, the absence of densities for modifications in crystallographic maps also at higher resolutions may result from differences in electron and X-ray scattering and in data quality which is affected, for example, by local and global disorder.
Extended Data Figure 5 | rRNA modifications in the E. coli 70S ribosome.

a, Stabilizing effects of 16S rRNA methyl groups in the P site of the decoding centre. Numbers in the overview (top left) mark the positions of close-ups (1–3), which show the interactions of the rRNA methyl groups with distances colour-encoded by dashed lines. Close-up 1: Stacking network of m\(^2\)G\(_{966}\) and m\(^5\)C\(_{967}\) stabilizing binding of initiator tRNA. Close-ups 2, 3: rRNA modifications impacting mRNA binding. The universally conserved bulky dimethylamine groups of m\(^6\)m\(^2\)A\(_{1518}\) and m\(^6\)m\(^2\)A\(_{1519}\) stabilize their direct environment by steric encumbrance explaining their requirement for correct packing of 16S rRNA helices 24a, 44 and 45\(^\text{b}\). In particular, the dimethylamine of m\(^6\)m\(^2\)A\(_{1519}\) is involved in medium and long-range repulsive interactions with the backbone of m\(^3\)U\(_{1498}\) and the 2’ O of C\(_{1520}\), while its conformation is mostly determined by the dimethylamine moiety of the adjacent m\(^6\)m\(^2\)A\(_{1518}\) which, in turn, is fixed by short repulsive interactions with O6 of G\(_{1517}\) and O4 of U\(_{793}\). The additional methyl groups of m\(^6\)m\(^2\)A\(_{1519}\) interact with A\(_{792}\), which provides part of the binding site for the antibiotic kasugamycin\(^\text{c}\), accounting for the resistance against kasugamycin upon demethylation of m\(^6\)m\(^2\)A\(_{1519}\)\(^\text{d}\). Furthermore, m\(^6\)m\(^2\)A\(_{1518}\) and m\(^6\)m\(^2\)A\(_{1519}\) impact initiation\(^\text{e}\) possibly via m\(^6\)U\(_{1498}\) whose backbone interacts with m\(^6\)m\(^2\)A\(_{1519}\), while its modified base contacts the mRNA backbone. The methyl groups of m\(^6\)U\(_{1498}\) and m\(^6\)C\(_{1402}\) form part of the binding site for the initiation codon and/or preventing direct hydrogen bonds with the mRNA backbone.

b, Constitutive rRNA modifications in the E. coli 70S ribosome (list adapted from ref. 11 and references therein).

c, rRNA modifications in the A site of the decoding centre and helix 69 of 23S rRNA (H69). The binding site of aminoglycosides in helix 44 of 16S rRNA (h44)—including N4 of m\(^5\)C\(_{1407}\)—is indicated for neomycin B (magenta, superposition from PDB ID: 2ET4)\(^\text{f}\). The three pseudouridines stabilizing H69\(^\text{g}\) by enhancing base stacking\(^\text{h}\) are depicted in blue. The methyl group (yellow) on m\(^3\)Ψ\(_{1917}\) in H69 prevents potential base-pairing with A\(_{1913}\), a residue important for uniform tRNA selection\(^\text{i}\). Note the flipped out conformation of A\(_{1913}\) facilitating interaction with the 2’ OH of m\(^6\)s\(^1\)A\(_{37}\) of the distorted Phe–tRNA\(^\text{m}\) (purple), which, in turn, stacks onto A36 of the tRNA anticodon. d, Methyl group on m\(^6\)G\(_{1835}\) of 23S rRNA enhancing subunit association. The four helices of 23S rRNA that intersect around residue m\(^6\)G\(_{1835}\) and form intersubunit bridges B2b and B2c with helices 24 (h24) and 45 (h45) of 16S rRNA (dark grey) are denoted in different colours: helix 67 (H67), light blue; helix 68 (H68), blue; helix 69 (H69), teal; helix 70 (H70), purple. Inset, contacts of the methyl group on m\(^6\)G\(_{1835}\) with adjacent residues which, in turn, interact with 16S rRNA.

e, Cluster of 23S rRNA modifications in the peptide exit tunnel. The modified rRNA residues, the functionally important nearby tip of protein L22 (teal), P-site fMet–tRNA\(^\text{m}\) (green) and a model of the nascent peptide chain (pink, superposition from PDB ID: 2WWL)\(^\text{j}\) are indicated.
Extended Data Figure 6 | Visualization of structural dynamics of the ribosome by different approaches. a–d, In each panel, the ribosome is shown from the factor binding site on the left and in cut-away view on the right; h denotes the head and b the body of the 30S ribosomal subunit. a, Present cryo-EM map coloured according to local resolution as determined by Resmap. b, Present cryo-EM map coloured according to the B factors obtained from the pseudo-crystallographic atomic model refinement (Methods). c, Model map of the 2.95 Å crystal structure of the *E. coli* 70S ribosome (PDB IDs: 4KJ1 and 4KJ2) coloured according to respective B factors. The black arrow denotes the stabilization of the 30S head region by crystal contacts, whereas the 30S body (white arrow) is less constrained by crystal contacts and shows higher B factors, indicating larger flexibility for this region. d, Snapshot from molecular dynamics trajectory of the *E. coli* 70S ribosome coloured according to root mean squared fluctuations (RMSFs) obtained from the full 2 μs explicit solvent molecular dynamics simulation (Methods). Note the large fluctuations of the 30S head and body (white arrows) of the ribosome in solution not constrained by crystal contacts.
Extended Data Figure 7 | Structure of E. coli EF-Tu–Phe–tRNAPhe bound to the ribosome. a, Detailed comparison of the distorted A/T-site–tRNA interactions between the E. coli and T. thermophilus ribosome–EF-Tu–Kirromycin complexes6,26 and the free E. coli tRNAPhe51. We found significant differences in tRNA conformation and interactions implicated in the GTPase activation mechanism26 that correlate with ribosome binding and differences in organism and tRNA species. Here and below residue numbers refer to E. coli.

b, Overview of the E. coli ribosome–EF-Tu structure. The residues interacting with the EF-Tu ternary complex (depicted in stick representation) generally agree with those seen in the T. thermophilus structures6,26. rRNA helices are denoted as: h44, helix 44 of 16S rRNA; H69, helix 69; and SRL, sarcin–ricin loop of 23S rRNA. The dashed boxes indicate the parts of the structure magnified in c and d.

c, Structural differences in an important ribosome–EF-Tu interaction. In the E. coli structure (Eco, left panel) residues A55 and A368 of 16S rRNA assume different conformations (cyan arrows) and interact differently with the ribosome and EF-Tu–tRNA complex than in the T. thermophilus structure (Tth, right panel, PDB ID: 2WRN). Furthermore, the differences in EF-Tu sequence result in slightly different ribosome–EF-Tu interactions, for example, the e-amino group of lysine 282 in E. coli EF-Tu is within hydrogen-bonding distance of G382 in 16S rRNA, but not the serine at this position in T. thermophilus EF-Tu. Inset on left panel, overlay of the crucial β-turn74 in EF-Tu domain 2 in the free (PDB ID: 1OB2, R. C. Nielsen et al., unpublished data) and ribosome-bound state from E. coli and T. thermophilus6,26,76 (dark grey, complex as indicated) are shown with the corresponding rotamer of the present structure (red). Residues valine 20 and isoleucine 60 of the hydrophobic gate76 are denoted; isoleucine 60 is not resolved in the kirromycin-stalled ribosome–EF-Tu complexes.

d, Dynamics of the catalytic histidine 84 (ref. 75) of EF-Tu. A split density for the side chain of histidine 84 (data not shown) indicates the presence of two rotamers (rot1 and rot2, panel 1) in the present E. coli complex. In the T. thermophilus ribosome-bound EF-Tu structures6,26,76 (dark grey, complex as indicated) are shown with the corresponding rotamer of the present structure (red). Residues valine 20 and isoleucine 60 of the hydrophobic gate76 are denoted; isoleucine 60 is not resolved in the kirromycin-stalled ribosome–EF-Tu complexes.

Extended Data Table 1

| complex | tRNA\(^\text{free}\) | 70S-EF-Tu/Phe-tRNAPhe\(\text{GTP}\) | 70S-EF-Tu-Thr-tRNAThr\(\text{GDP}\) | 70S-EF-Tu-Trp-tRNATrp\(\text{GDP}\) |
|---------|-----------------|-----------------|-----------------|-----------------|
| organism/PDB ID | E. coli/3LOU | E. coli/present model | T. thermophilus/2WRN,2WRO | T. thermophilus/2Y18,2Y19 |
| tRNA structure | G10:G25;G44 | base triple | base triple | base triple |
| A26:U45 | none | WC | none (G:G) | WC |
| G27:C43 | WC | reverse WC | none | (A:U) |
| G15:C48 | reverse WC | reverse WC | none | (D:G) |
| D16:U59 | D:U | none | none (D:A) | none |
| G1:C72 | WC | WC | none, C72 flipped out | WC |
| tRNA-23S rRNA interactions | C56:A1067 | n.a. | parallel base-stacking | T-shaped base-stacking |
| D20:U1066 | n.a. | none | H-bonding distance | none |
| tRNA-16S rRNA interactions | C75:A55 | n.a. | none (A55 flipped in) | stacking (A55 flipped out) |

*Abbreviations: WC, Watson-Crick base pair; none, no base pairing or no interaction; n.a., not applicable

†unmodified; free modified tRNA\(^\text{free}\) from E. coli not available

‡base pair in modified tRNA\(^\text{free}\) from Saccharomyces cerevisiae (PDB ID 1EHZ)
Extended Data Figure 8 | Cryo-EM densities for mobile proteins L9 and L31 and the arrangement of L9 in polysomes. Densities in a and b were obtained by semi-automatic segmentation using the ‘segger’ tool in UCSF CHIMERA\(^41\), normalized and low-pass filtered according to local resolution estimates. a, Cryo-EM map and models of protein L9 (\(~4\,\AA\) local resolution, rendered at 1\(\sigma\)). b, Cryo-EM maps and models of protein L31 in the ground-state of the ribosome (left, \(~4.3\,\AA\) local resolution) and in the rotated state (right, \(~6\,\AA\) local resolution); maps were rendered at 1.5\(\sigma\). c. Model of protein L9 in the context of polysomes. Overviews show the arrangement of neighbouring ribosomes (i-1 and i) in the major t-t form of \(E.\ coli\) polysomes as obtained by fitting the present 70S ribosome structure into the cryo-electron tomography reconstruction\(^27\) in UCSF CHIMERA\(^41\). Left close-up, the conformer of L9 as seen in the present cryo-EM map (L9 cryo-EM, blue) is located close to protein S4 of the neighbouring 30S subunit (30S i) according to the polysome model. The purple arrow indicates the rearrangement of L9 from the cryo-EM conformation to that seen in crystals. The black arrow denotes the location of the mRNA entry channel in the 30S subunit i. Right close-up, the conformer of L9 as seen in the context of ribosome crystals (L9 X-ray, pink) reaches into the ribosomal A-site of the neighbouring 30S subunit and would be compatible with the simultaneous binding of elongation factors in the polysome model. In crystals, protein L9 precludes the binding of elongation factors due to the tighter packing of ribosomes\(^78\). The model of L9 was obtained by superposition of the \(E.\ coli\) 70S ribosome X-ray structure\(^6\) (PDB IDs: 4TP8 and 4TP9) onto ribosome i-1 in the polysome model using UCSF CHIMERA\(^41\).
Extended Data Table 1 | Data collection and model refinement

| E. coli 70S ribosome-EF-Tu complex | Complete model | 65% of model | 43% of model | 24% of model |
|------------------------------------|----------------|--------------|--------------|--------------|
| **Data collection**                |                |              |              |              |
| Particles                          | 417210         |              |              |              |
| Pixel size (Å)                     | 0.75525        |              |              |              |
| Defocus range (µm)                 | 0.7–2.5        |              |              |              |
| Electron dose (e/Å²)               | 40±5           |              |              |              |
| **Reciprocal space data**          |                |              |              |              |
| Space group                        | P1             |              |              |              |
| a, b, c (Å)*                       | 302.1, 302.1, 302.1 |              |              |              |
| a, β, γ (°)                        | 90.0, 90.0, 90.0 |              |              |              |
| Resolution (Å)                     | 2.9            | 2.80         | 2.70         | 2.65         |
| Wilson B (Å²)                      | 44.1           | 46.0         | 45.24        | 43.7         |
| **Refinement†**                    |                |              |              |              |
| Refinement target: MLHL maximum likelihood with experimental phase probability distribution. |                |              |              |              |
| Highest resolution shell is shown in parenthesis. |                |              |              |              |
| Resolution range (Å):‡             | (2.95 – 2.90)  | (2.85 – 2.80) | (2.75 – 2.70) | (2.70 – 2.65) |
| No. Reflections                    | 2366496 (112206) | 2629133 (125775) | 2952674 (139922) | 3101577 (146909) |
| Rwork/B/C (|F|/|F|model) | 0.241/0.922 | 0.221/0.940 | 0.232/0.937 | 0.231/0.939 |
| FSCwork | (0.479/0.298) | (0.494/0.238) | (0.500/0.209) | (0.501/0.252) |
| ML based phase error (*)           | 0.930 (0.541)  | 0.942 (0.500) | 0.938 (0.529) | 0.940 (0.502) |
| Residues with atoms out of density for main chain/side chain# | 28.9 | 27.8 | 29.9 | 29.9 |
| No. Atoms/RSCC★                    | 152719         | 99034        | 66327        | 36067        |
| Protein                            | 49242/0.818    | 14938/0.824  | 6795/0.813   | 32590/0.818  |
| RNA, tRNA                          | 103301/0.872   | 63922/0.873  | 59461/0.872  | 32799/0.871  |
| Ligand, ion                        | 445            | 173          | 71           | 9            |
| B-factors (Å²)                     | 95.4           | 68.4         | 61.9         | 60.8         |
| Protein                            | 110.0          | 74.2         | 66.0         | 69.9         |
| RNA, tRNA                          | 88.5           | 67.5         | 61.5         | 59.9         |
| Ligand, ion                        | 66.5           | 29.1         | 20.1         | 24.9         |
| R.m.s deviations                   |                |              |              |              |
| Bond lengths (Å)                   | 0.004          | 0.007        | 0.007        | 0.006        |
| Bond angles (°)                    | 0.729          | 1.060        | 0.929        | 0.971        |
| Ramachandran plot statistics       |                |              |              |              |
| Most favored (%)                   | 85.59          | 87.4         | 88.56        | 87.53        |
| Disallowed (%)                     | 3.30           | 2.66         | 2.78         | 3.52         |

*For model refinement, the cryo-EM map was cropped from 420 x 420 x 420 pixels to 400 x 400 x 400 pixels.
†Refinement target: MLHL maximum likelihood with experimental phase probability distribution.
‡Highest resolution shell is shown in parenthesis.
§Rwork = \[\frac{\sum |F_{EM} - |F_{model}|}{\sum |F_{EM}|}\], where \(F_{EM}\) are structure factors calculated based on the solvent-flattened cryo-EM map and \(F_{model}\) are structure factors calculated from the refined model. The structure factors belonged to working set which was used for reciprocal space refinement.
#Calculated with RESOLVE.
Extended Data Table 2 | Comparison of densities as observed in cryo-EM and X-ray crystallography by local cross-correlation

| Method/resolution | Cryo-EM @ 2.9 Å E. coli 70S-EF-Tu-tRNA<sub>0</sub> | X-ray @ 2.95 Å E. coli 70S-EF-G 2KJ1 & 2KJ2<sup>®</sup> | X-ray @ 2.8 Å E. coli 70S-quinupristin 4TPA & 4TPB<sup>®</sup> | X-ray @ 2.8 Å T. thermophilus 70S-tRNA 2J00 & 2J01<sup>®</sup> |
|-------------------|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|
| Ribosome complex  |                                               |                                               |                                               |                                               |
| PDB ID            |                                               |                                               |                                               |                                               |
| **Protein**       |                                               |                                               |                                               |                                               |
| Map correlation in region of model | 0.818 | 0.836 | 0.787 | 0.857 |
| Total residues    | 6320  | 6460  | 5722  | 5909  |
| Residues in acceptable density† | 4293  | 5214  | 4019  | 3710  |
| Residues with some weak density‡ | 2027  | 1246  | 1703  | 2199  |
| Residues out of density§ | 3     | 0     | 0     | 0     |
| Residues in very weak density¶ | 103   | 19    | 27    | 49    |
| Residues in weak density¶ | 840   | 316   | 668   | 872   |
| Main-chain out of density | 0     | 1     | 0     | 0     |
| Main-chain in very weak density | 1     | 1     | 0     | 1     |
| Main-chain in weak density | 225   | 77    | 158   | 208   |
| Side-chain out of density | 3     | 3     | 12    | 2     |
| Side-chain in very weak density | 36    | 64    | 58    | 94    |
| Side-chain in weak density | 481   | 508   | 573   | 915   |
| Avg. RSCC for Arg, His and Lys side chains | 0.933 | 1.159 | 0.918 | 0.790 |
| Avg. RSCC for Glu and Asp side chains | 0.514 | 1.090 | 1.091 | 0.588 |
| Avg. RSCC for Gin and Asn side chains | 0.865 | 1.273 | 1.151 | 0.952 |
| Avg. RSCC for aromatic side chains | 1.320 | 1.551 | 1.260 | 1.324 |
| Avg. RSCC for aliphatic side chains | 0.887 | 0.972 | 0.693 | 0.852 |
| **RNA**           |                                               |                                               |                                               |                                               |
| Map correlation in region of model | 0.749 | 0.888 | 0.861 | 0.909 |
| Total residues    | 4800  | 4505  | 4554  | 4494  |
| Residues in acceptable density | 2860  | 3602  | 3021  | 3573  |
| Residues with some weak density | 1940  | 903   | 1533  | 921   |
| Residues out of density | 2     | 0     | 0     | 0     |
| Residues in very weak density | 104   | 36    | 28    | 59    |
| Residues in weak density | 501   | 180   | 596   | 425   |
| Main-chain out of density | 0     | 0     | 0     | 0     |
| Main-chain in very weak density | 4     | 1     | 0     | 0     |
| Main-chain in weak density | 495   | 96    | 189   | 157   |
| Side-chain out of density | 0     | 0     | 0     | 0     |
| Side-chain in very weak density | 0     | 0     | 0     | 0     |
| Side-chain in weak density | 18    | 10    | 58    | 37    |

* Real space correlation coefficients (RSCCs) were calculated with RESOLVE<sup>®</sup> to compare the local model fit to the normalized experimental map.
† ‘acceptable density’: no atom lies in density < 1 s.d. below 1/2 mean for that atom type and no group is < 1/2 mean for that group.
‡ ‘residues with some weak density’: all residues which do not fulfill the requirement for acceptable density.
§ ‘out of density’: density level > 2 standard deviations below 1/2 mean density for the particular residue.
¶ ‘very weak density’: density level < 1/2 mean density for the particular residue.
Φ ‘weak density’: density level < 1/2 mean density for the particular residue.
Overall, the definition of protein and RNA main and side chain densities is similar between the present cryo-EM data and 70S ribosome crystal structures at similar resolutions, suggesting also a similar extent of radiation damage. The RSCC values obtained for specific amino acid side chain groups indicate different sensitivities to radiation damage for specific side chains, with negatively charged side chains being the most sensitive ones both in cryo-EM and X-ray crystallography<sup>®</sup>. The elevated RSCCs calculated for the 2.95 Å E. coli 70S-EF-G crystal structure show the smallest fluctuations between different side chain groups and, thus, the lowest level of radiation damage, most probably owing to multi-crystal merging of 20 partial data sets. In contrast, experimental data for the 2.8 Å E. coli 70S-quinupristin and the 2.8 Å T. thermophilus 70S-tRNA crystal structures were collected from two and one crystal(s), respectively.