The Structure of *E. coli* Signal Recognition Particle Revealed by Scanning Transmission Electron Microscopy

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**Running Title:** Domain Organization of *E. coli* SRP

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**Abbreviations:** 2-D, two-dimensional; 3-D, three-dimensional; CPM, 7-diethylamino-3-(4′-maleimidylphenyl)-4-methylcoumarin; FRET, fluorescence resonance energy transfer; IQAD, Iterative Quaternion-based Angular Determination; SRP, signal recognition particle; STEM, scanning transmission electron microscopy; TF, trigger factor.
Abstract

Structural studies on various domains of the ribonucleoprotein signal recognition particle (SRP) have not converged on a single complete structure of bacterial SRP consistent with the biochemistry of the particle. We obtained a three-dimensional structure for *E. coli* SRP by cryo-scanning transmission electron microscopy and mapped the internal RNA by electron spectroscopic imaging. Crystallographic data was fit into the SRP reconstruction and although the resulting model differed from previous models, they could be rationalized by movement through an interdomain linker of Ffh, the protein component of SRP. Fluorescence resonance energy transfer experiments determined interdomain distances consistent with our model of SRP. Docking our model onto the bacterial ribosome suggests a mechanism for signal recognition involving interdomain movement of Ffh into and out of the nascent chain exit site and suggests how SRP could interact and/or compete with the ribosome-bound chaperone, trigger factor, for a nascent chain during translation.
**Introduction**

The mechanism of co-translational protein targeting is conserved in most organisms. The initial step involves binding of the signal recognition particle (SRP)\(^1\) to both a ribosome and a signal sequence on the nascent polypeptide chain emerging from the ribosome. SRP targets the ribosome-nascent chain complexes to membranes via binding to the SRP receptor (Gilmore *et al.*, 1982). Receptor binding and transfer of the nascent polypeptide from SRP to the translocation machinery in the membrane are regulated by the GTPase activities of SRP and SRP receptor (reviewed in Millman and Andrews, 1997).

Depending on the organism, the components of SRP can vary in number and size. Bacteria have the simplest SRP, composed of a single polypeptide and a small RNA molecule. In *E. coli*, these subunits are called Ffh (‘fifty-four homologue’; for its homology to the SRP54 subunit of human SRP) and 4.5S RNA, respectively. Based on sequence comparisons, the primary sequence of Ffh has been arbitrarily divided into the N (amino-terminal), G (GTPase) and M (methionine-rich) regions.

Several structures have been determined for isolated domains of Ffh. Structures of the apo-form of the NG domain of Ffh (Freymann *et al.*, 1997), and various guanine nucleotide-bound forms (Freymann *et al.*, 1999) revealed that the N and G regions, though distinct based on sequence alignments, are structurally organized as a single tertiary domain. The M domain forms another distinct domain that binds 4.5S RNA and contains the putative signal sequence binding site. M isolated from *T. aquaticus* Ffh (Keenan *et al.*, 1998) and human SRP54 (Clemons, Jr. *et al.*, 1999) have been crystallized in the apo form, and *E. coli* M was solved from a co-crystal with a fragment of the 4.5S RNA (Batey *et al.*, 2000). A crystal that contained full-
length Ffh from T. aquaticus (Keenan et al., 1998) revealed the structures of both NG and M from a single organism but the linker between them was not resolved and the crystal packing precluded unambiguous determination of the relative orientation of the two domains.

Recently, a structure for the archaea Sulfolobus solfataricus full-length SRP54 bound to a fragment of SRP RNA was solved crystallographically (Rosendal et al., 2003). The complete backbone of the protein component was resolved but a 19 kDa protein normally found in this SRP was not included. In addition, much of the archaeal SRP RNA was missing including a region corresponding to nucleotide 84 of E. coli SRP RNA, a site that can be cross-linked to the ribosome (Rinke-Appel et al., 2002) and therefore is probably involved in binding SRP to the ribosome. Interestingly, in S. solfataricus SRP54 the relative orientation and position of M and NG were markedly different from all three of the possible arrangements deduced from the T. aquaticus Ffh structure. However, the domain structures were conserved, suggesting that the NG and M domains behave primarily as separate rigid bodies with high structural conservation across different species.

The length of the SRP RNA varies greatly among organisms from approximately 300 nt in mammals and archaea (reviewed in (Poritz et al., 1988)) to 70 nt in Mycoplasma mycoides (Samuelsson and Guindy, 1990). In organisms with a longer SRP RNA, the particle generally contains multiple protein components; indeed, one proposed role of the RNA is to act as a scaffold for particle assembly. The E. coli SRP RNA is 114 nt of which nucleotides 39-64 have been implicated in binding to Ffh. Little is known about the function of the rest of the RNA in E. coli SRP; additional binding proteins have not been identified, and nucleotides 1-32 and 75-114 can be deleted without compromising viability in rich medium (Batey et al., 2000).
Electron spectroscopic imaging (ESI) of mammalian SRP suggested that much of the RNA functions as a scaffold for six SRP polypeptides (Andrews et al., 1985). Based on a structural model deduced from STEM micrographs, it was proposed that SRP could span the ribosome from the peptide bond formation site (for elongation arrest) to the peptide exit site (where SRP binds the emerging signal sequence). This model for SRP binding to the ribosome was recently verified and refined by cryo-electron microscopy of canine SRP bound to the 80S ribosome (Halic et al., 2004) to indicate direct binding of SRP to a region of the ribosome adjacent to the nascent chain exit site and to the elongation factor binding site near the peptide bond formation site. This structure of SRP also suggests that ribosome binding is another important function for the SRP RNA.

Further structural information is required for several forms of SRP to determine the molecular mechanisms of signal peptide binding, regulation and release. We have visualized E. coli SRP using cryo-scanning transmission electron microscopy (STEM) and calculated a three-dimensional (3-D) reconstruction. ESI in the STEM was utilized to locate the RNA molecule in three dimensions within the SRP reconstruction and then we incorporated data from existing structures to generate a high-resolution model of E. coli SRP. This model is consistent with our measurements of interdomain distances by fluorescence resonance energy transfer (FRET) for free SRP in solution. Docking our model on the ribosome revealed a potential mechanism for signal peptide binding that involves rotation of the M domain of Ffh. Motion of M relative to NG is also predicted from FRET data for SRP. Our model also explains why SRP binding to its receptor, FtsY, and TF binding on the ribosome may be mutually exclusive.
Materials and Methods

Preparation of bacterial signal recognition particle

*E. coli* SRP was made by incubating a 5-fold molar excess of recombinant poly-histidine-tagged Ffh (Zheng and Gierasch, 1997) and *in vitro* transcribed 4.5S RNA for 10 minutes at 0°C and then 10 minutes at 37°C (Freymann et al., 1997). RNA-containing complexes were separated from unbound protein by ion-exchange chromatography using Sepharose DEAE Fast-Flow (GE Healthcare) (adapted from (Walter and Blobel, 1983)). This eluate was then applied to a column containing Ni-NTA resin (QIAGEN) to remove unbound RNA. From an initial binding reaction containing 4.0 nmol of Ffh, the typical yield of SRP obtained by this purification procedure was 20% as 300 µL of 2.6 µM SRP (determined by protein quantification).

Electron Microscopy and Data Collection

Grids containing a carbon film were glow discharged. Purified complexes were adsorbed to the carbon films by injecting the sample into an equal volume of 2.5 mM magnesium acetate on the support. The grid was transferred, in succession, to a series of drops of 2.5 mM magnesium acetate and then was washed with droplets of 50 mM ammonium acetate. Residual ammonium acetate was wicked away and allowed to evaporate partially to form a thin layer across the grid, and then the grid was plunged into liquid ethane. Frozen grids were freeze-dried in a STEM (Vacuum Generators, Model HB601UX) at a temperature of -130°C.

‘Low-dose’ dark field electron micrographs of the freeze-dried SRP were obtained at -130°C in the STEM at a magnification of 500,000x with a gun voltage of 100 kV, corresponding to an electron dose of 15 e/Å². Images were acquired digitally by using a 3Å electron beam at
pixel spacings of 3.3Å. Particle images were converted to mass/pixel by calibrating the STEM image intensity to the carbon film thickness. Carbon film thickness was determined by analysis of the energy loss spectrum (Egerton, 1986) obtained on a Zeiss EM902 transmission electron microscope. The STEM images were filtered by a 1 pixel radius Gaussian blur and contrast enhanced for optimal viewing.

ESI images were collected in the STEM using a 6Å electron beam at 3.3Å pixel spacings. At each point, electrons were counted for 4 windows of spectroscopically separated energy loss values (centered at 0 eV, 112.5 eV, 125 eV, and 150 eV) captured simultaneously in parallel.

**Three-dimensional reconstruction of SRP**

Individual single molecule images of *E. coli* SRP were identified and extracted using the SPIDER/WEB image processing software (Frank et al., 1996) resulting in a set of 3,236 particle images. After normalization of the images, they were randomly divided into 16 subsets of 201 particles. For each subset the relative angular orientations of the randomly oriented particles was determined by angular reconstitution using IQAD (Farrow and Ottensmeyer, 1993). The distributions of angles determined by IQAD were used to calculate completely independent 3-D reconstructions of SRP by back-projection of the individual subsets of original unfiltered images. The reconstructions generated for the individual subsets were refined by a projection matching algorithm (Penczek et al., 1994) created with SPIDER. The refined reconstructions were aligned in three dimensions using EMAN (Ludtke et al., 1999) and combined into a single reconstruction using a round of multi-reference projection matching. The reconstruction was further refined using a two-stage modified version of the projection matching technique (Penczek et al., 1994)
created with SPIDER. Briefly, class images were generated by forward projection of the 3-D structure from the previous iteration. For the first stage of the refinement process, the class images are generated at isotropically distributed orientations at a relatively coarse angular interval (10-15°) and used in the projection matching for the initial alignment of each of the particle images. Next, a second set of class images is calculated for each particle image at a finer angular interval (0.3-1.5°) within a matrix defined by two of the top matched class images from the first stage. This two-stage process results in a very fine sampling of the angular space with a much shorter computation time than if the fine angular interval was used throughout the entire projection matching process.

Reconstructions of the STEM images were calculated on Silicon Graphics IndigoII and Origin2000 Workstations (Silicon Graphics, Mountain View, CA, USA). Visualization of structure data was done using InsightII (Accelrys Software Inc., San Diego, USA). Ffh structures were aligned using an alignment feature of InsightII.

Electronic Spectroscopic Imaging (ESI)

For each set of ESI images of well separated SRP complexes, the average signals for the 112.5 and 125 and 150 eV loss images (essentially carbon film) were used to calculate a correction factor to normalize the detectors to the electron energy loss spectrum curve approximation: \( I = A \cdot E^{-R} \) for carbon (where \( I \) is intensity, \( E \) is electron loss voltage, and \( A \) and \( R \) vary with respect to specimen thickness and chemical composition (Egerton, 1986)). Individual particles of SRP were identified from low angle scattering images. Particle images were extracted from complementary regions of the corresponding 112.5, 125 and 150 eV loss images. For each pixel
in the particle area of the 150 eV loss image, the background intensity due to the carbon film was extrapolated from the corresponding 112.5 and 125 eV loss pixel intensities. The extrapolated background was subtracted from the actual 150 eV loss pixel intensities, to generate an image from only those electrons that had interacted with and lost energy to the phosphorus L-shell electrons, referred to as the ‘net phosphorus signal’.

Three of the energy loss images (112.5 eV, 125 eV and 150 eV) were summed together and projection matched to the SRP reconstruction generated using low-dose STEM images. The corresponding net phosphorus signal images were reconstructed into a 3-D structure by back-projection based on the angular orientation assigned from the summed energy loss images. The resulting reconstructions are inherently noisy due to both quantum noise and radiation damage and, therefore, are presented at an arbitrary threshold that illustrates a relatively contiguous signal representing a 3-D phosphorous localization map.

Fluorescence resonance energy transfer (FRET)

Ffh was expressed from the pGEV2 plasmid (kind gift of A. Gronenborn (Huth et al., 1997)) resulting in a fusion protein with the ‘solubility enhancing tag’ GB1 from the Streptococcal protein G at the N-terminus and a hexa-histidine tag at the C-terminus of Ffh. Mutant versions of the Ffh fusions were expressed in the BL21 (DE3) strain of E. coli cells and purified by ion-exchange chromatography followed by nickel-affinity chromatography. The signal peptide binding activity was verified for each of the mutants (Cleverley and Gierasch, 2002). A 43 nt fragment of 4.5S RNA fragment was prepared as in (Zheng and Gierasch, 1997).

For the generation of proteins with single tryptophan and cysteine residues for FRET
experiments, a mutant version, referred to as Ffh*, was initially made in which the single tryptophan of the GB1 moiety was replaced with phenylalanine and Cys406 of the Ffh was mutated to serine (original Ffh amino acid numbering is retained for easy reference). The following proteins were created: Ffh* R11W/Q340C, Ffh* R11W/S366C, Ffh* R49W/Q340C, Ffh* R49W/S366C, and Ffh* R141C/M347W. Cysteines were externally labeled with the 7-diethylamino-3-(4'-maleimidylphenyl)-4-methylcoumarin (CPM) fluorophore (Molecular Probes, Inc.) according to the manufacturer’s protocols. The labeling efficiencies were in the 0.75-0.85 range for all mutants as determined from spectrophotometric measurements. These values were incorporated into the data analysis during calculation of the distance distributions.

Time-resolved FRET measurements were carried out on a custom-built single photon counting apparatus provided by the C.R. Matthews lab (UMASS School of Medicine, Worcester, MA). Excitation of fluorescence at 295 nm was accomplished by the triplet output of a mode-locked titanium:sapphire laser and fluorescence decays were monitored at 350 nm (E.F. and L.M.G., in preparation). The analysis of time-resolved decay profiles and the distribution of distances between donor and acceptor fluorophores were carried out using the Savuka program (version 6.0.1). Further details of the FRET experiments are presented in Supplemental Material and will be published elsewhere.

All fluorometric measurements were carried out in 10 mM HEPES pH 7.6, 300 mM sodium chloride, 10% glycerol, and 5 mM magnesium chloride. Protein was incubated with a 1.5 molar excess of 4.5S RNA fragment to ensure complete formation of ribonucleoprotein particles.
Results

SRP 3-D Reconstruction

We assembled SRP from its constituent protein and RNA. The reconstituted *E. coli* SRP exhibited the expected GTPase activity (data not shown) and the Ffh was shown to bind signal sequences (Cleverley and Gierasch, 2002) therefore we conclude that the particles used for imaging were functional. To obtain sufficient signal-to-noise ratio to visualize the relatively small *E. coli* SRP (approximately 90 kDa), we used a freeze-dried, unstained sample in the dark-field imaging mode of a STEM. Mass determinations from the STEM images confirmed that the particles used for the reconstruction were monomers of SRP with a mass of 77 +/- 15 kDa. The STEM images of SRP appeared as curved rods with one end larger than the other (Figure 1A). Angular assignment of the individual images revealed there was a slight preference for some orientations of SRP binding to the carbon support (see Supplemental Material). Filtered back-projection included orientation density-dependent filtering for the calculation of the SRP reconstruction to ensure that the higher occurrence projections were not over-weighted in the final reconstruction.

The STEM data converged on a 3-D structure for *E. coli* SRP resembling the numeral 9 from a ‘front’ view (Figure 1B, left) with a globular ‘head’ and a ‘tail’ extending 90Å from one side of the head. The tail gives the reconstruction an overall extended conformation with a maximal length of 150Å, and width of 85Å at the head and a minimal width of 42Å along the tail in this orientation. The structure has a slight curvature along the tail but is otherwise flat (thickness 30Å). The resolution of our SRP reconstruction was estimated to be 12Å by Fourier shell correlation (FSC) using the 3σ criteria (Orlova et al., 1997) (see Supplemental Material for
complete FSC plot). There are three possible locations for the RNA molecule in this reconstruction: first, the RNA could be completely localized to the tail, extending most of the length of the particle; second, the RNA could be folded in such a way that it occupies the globular head of the reconstruction; or, third, the RNA could be positioned so that part of it is in the head and part of it is in the tail (see Supplemental Material for a schematic of all of these possibilities). The first two RNA locations are potentially compatible with the remainder of the particle being the protein component, Ffh (i.e. the head region or the tail region). The third position of the RNA would require that two relatively distant regions of the reconstruction (the distal end of the tail and the remaining head volume) would need to be filled by Ffh and it is not readily apparent how this could be achieved in a biologically relevant/possible way (see Supplemental Material for a description of model building). The length of the tail in the reconstruction is consistent with a highly base-paired double-helix. Moreover the repetitive features of an RNA helix can be discerned in the right hand view of the particle tail (arrowheads in Figure 1B) and, although the tail deviates from the ribbon-like structure that would be expected for a perfect double-helical RNA molecule at the resolution of the reconstruction, it corresponds well with a rod-like RNA model that has the expected secondary structure of *E. coli* SRP RNA (see Supplemental Material). However, chemical and enzymatic probing data of *E. coli* SRP and FRET measurements of SRP RNA bound to Ffh suggest that at least in solution the RNA can fold back on itself when in complex with Ffh (Buskiewicz et al., 2005a),(Buskiewicz et al., 2005b) which would be more consistent with the RNA being located in the head of the particle. To resolve this apparent discrepancy and to correctly assign the RNA and protein components in the reconstruction of *E. coli* SRP, it was therefore, essential to localize the RNA
in the structure directly.

SRP RNA mapping

To locate the nucleic acid within *E. coli* SRP, the distribution of phosphorus in the particle was determined by ESI. Electrons inelastically scattered by L-shell electrons of a phosphorus atom predominantly lose 150 eV of energy (Bazett-Jones and Ottensmeyer, 1982) and can be identified with an electron spectrometer attached to the STEM. These electron spectroscopic images of the phosphorus distribution were used to generate a 3-D reconstruction of the phosphorus within SRP. The resolution in these maps is limited by radiation damage and quantum noise but is just sufficient to track double-stranded nucleic acids in particles, including canine SRP (Andrews *et al.*, 1987). The 3-D ESI phosphorus map is quite noisy, as expected from the small number of phosphorus atoms imaged. Therefore, the reconstruction was filtered to 16Å and shown at an arbitrary threshold that generates a mostly contiguous volume. Lowering the threshold further increases the size of the spurious signals (small structures not connected to the main signal) without changing the overall interpretation. Increasing the threshold leads to further fragmentation but the main signals remain consistent with an extended RNA in SRP. Thus we conclude that the map is sufficient to demonstrate that the RNA adopts an extended rod-like structure with a small but significant bend in the middle (Figure 1C). While the phosphorus map provides little information about the RNA structure beyond that it forms an extended rod, it was calculated independently of the total structure. Therefore, the fact that when the SRP reconstruction was displayed together with the phosphorus map, the phosphorus map overlapped the SRP reconstruction (Figure 1D) confirmed that the tail of the SRP reconstruction contains RNA while the head region of SRP is mostly protein. The size of the gap
in the main axis of the phosphorus map and the spurious signals outside of this area suggests the limit to which the phosphorus map can be used to locate the RNA in the particle.

Generation of ec_SRp, a high resolution model of SRP

The reconstructions in Figures 1 and 2 are displayed as having a hard shell which makes interpretation easier but does not permit easy assessment of the sharpness of the edges of the reconstruction. In order to take this into account and dock crystallographic data into the reconstruction we examined the latitude for fitting by decreasing the viewing threshold (i.e. increasing the visualized volume) of the SRP reconstruction. When the threshold was reduced such that the reconstruction volume was increased by approximately 50% (Figure 2A, black areas) it resulted in a ‘filling in’ of the reconstruction without significantly changing the extent of the exterior boundaries. Thus the edges of the reconstruction are quite sharp and the framework/boundary for the docking of the individual domains of E. coli SRP into the reconstruction are quite obvious.

Two factors affect the appearance of the reconstruction at a particular viewing threshold. First, using an average density to analyze a particle containing both RNA and protein should result in portions of the reconstruction containing solely protein to be under-represented (or under-estimated) and the RNA-only portions to be over-represented. Secondly, the volume enclosed assumes that the reconstruction was calculated from a set of images of a particle adopting only one conformation. Areas of conformational heterogeneity in the image population result in lower density regions in the reconstruction due to misalignment of flexible regions when calculating the reconstruction (discussed below). Finally, the reconstruction will be dominated
by the most prevalent structure on the grid with other structures optimally aligned to the
dominant one such that a single unique reconstruction is generated that represents the most
populous SRP structure on the grid (the effect of the reconstruction process on the final result is
described in the *Supplementary Material*).

To construct a high resolution model of SRP, we first assigned the approximate position
of the RNA in the particle as shown in Figure 1E. This position of the RNA restricts the possible
locations for the M domain which was modeled using the structure from the co-crystal of *E. coli*
M and a fragment of SRP RNA (PDB: 1DUL). Various possible locations of M were examined
by rotating the RNA. During the modeling, the RNA was rotated in small increments however,
for illustrative purposes, the results of rotation at 90° intervals are shown in Figure 2B. Even
when the SRP reconstruction is visualized at a conservative threshold (150% of the expected
molecular mass), only one orientation (Figure 2B, #1) has M positioned entirely within the
volume of the reconstruction. When rotated 180° (Figure 2B, #2) the M domain extends into an
invagination in the SRP structure. Similarly, in the other two orientations shown (+/- 90°, Figure
2B, #3 and #4) M extended significantly beyond even the most conservative border of the SRP
reconstruction. The relatively low resolution of the 3-D phosphorus map allows for some
latitude in the vertical positioning of the RNA molecule (see *Supplemental Material*). Therefore
the fit of the M-domain in the reconstruction was examined after vertical translation of the RNA-
bound M structure. An upward shift within the reconstruction positioned the M-domain either in
the low density region (the ‘hole’) located just above position 1 or outside the mass distribution
(similar to positions 3 and 4) or, if it resulted in an acceptable fit of the M-domain within the
particle mass there was too little space left to fit NG within the head of the reconstruction (see

-15-
Supplemental Material). Shifting the RNA downward moves M below the mass of the head region and positions it too far from the NG domain to be compatible an appropriate interdomain linker (see Supplemental Material).

Positioning the M domain as shown in Figure 2B left a relatively constrained space in the head of the SRP reconstruction for the NG domain. The thickness of the particle constrains the NG domain to an orientation with the long axis aligned with the plane of the particle. Therefore, the NG domain could be docked into the remaining volume of the globular head in only two main orientations; one with the N region above the RNA molecule and the G region next to the M domain; and, one with the N region near the M domain and the G region above the RNA molecule (see Supplemental Material). Also, crystallographically determined structures available for the NG domain have a high degree of rotational symmetry. In order to set the rotational orientation of the NG domain, the length of an NG-M linker from the structure of S. solfataricus SRP54 (approximately 40Å) was used to constrain the distance between the carboxyl-terminus of the NG domain and the amino-terminus of the M domain. Surprisingly, there was a fairly limited region of rotational space that was compatible with this linker length, for either of the two N-G orientations (see Supplemental Material). Therefore, the T. aquaticus NG (PDB: 2FFH) was docked into the reconstruction in the two N-G orientations using the optimal rotation determined from the linker length analysis (see Supplemental Material). The N-G orientation with the N region above the RNA gave a more optimal visual fit (see Supplemental Material) and there were no significant molecular clashes of NG with either M or RNA. This orientation of NG was also compatible with docking SRP to FtsY and to the ribosome (see below). Finally, this orientation of NG in Figure 2C (but not the other NG
orientation) results in a positioning of NG relative to M that is compatible with FRET data for the particle in solution (see below). Once the position of the individual domains was assigned, we replaced the structure for the *E. coli* M domain, in which several amino acids are missing from a flexible loop believed to be involved in signal sequence binding, with the complete M structure from *S. solfataricus*. Finally, the remaining RNA fragment was constructed using a portion of a model of extended *E. coli* SRP RNA obtained from the SRP Database (Rosenblad *et al.*, 2003).

In the final model, referred to as *ec_SR*P, the alpha-helical bundle of the N region is situated above the hairpin tetraloop of the RNA and M is adjacent to the GTPase region of NG (Figure 2D and Figure 3A). The SRP RNA adopts a rod-like conformation with a significant bend at the approximate midpoint of the rod. There are about 8 ‘base pairs’ of RNA (~20 Å) that extend beyond the SRP reconstruction volume. In our model, the spacing between nucleotides in the distal half of the RNA is the same as for theoretical double-stranded, helical RNA with a few nucleotides left un-paired to match loops and bulges from the predicted secondary structure. Using the crystallized portion of the *E. coli* SRP RNA (nucleotides 33-74) as a molecular ‘measuring stick’, 22 nucleotide pairs (which includes a symmetric loop and an asymmetric loop) is approximately 50 Å long. Extending this to the full-length SRP RNA (~56 nucleotide pairs predicted to have similar occurrences of bulges and loops) generates an expected length of 128 Å, which, at about 12 Å shorter than the RNA model shown is consistent with the length of the RNA in our reconstruction. It is also possible that the tertiary structure of this region of the SRP RNA adopts a more irregular, complex configuration, such as side-loops extending from the main helix or a bending back of the distal end, which would result in an overall length that is
consistent with our reconstruction. Alternatively, it is possible that this region of the RNA is so mobile that it is not visible due to the averaging that occurs during the reconstruction process.

The diameter of the tail of the reconstruction is elliptical with dimensions of approximately 42Å by 30Å (+/- ~12Å) compared to 25Å for the diameter of the RNA model. The 30Å dimension is consistent with a largely helical RNA at the resolution of our reconstruction. The larger dimension in the reconstruction suggests conformational flexibility in the RNA that could come from mobile loops that extend away from the main RNA helix and/or from movement of the RNA in one dimension. Consistent with these possibilities, the RNA exhibits lower density than expected (see above). At a resolution around 10Å it should be possible to identify the characteristic shape of a double-stranded RNA, although, for a 3-D reconstruction of the *E. coli* 50S ribosomal subunit, with a much better resolution (7.5Å by FSC at 3σ threshold), a ribbon-like structure for the rRNA was not always readily apparent (Matadeen *et al.*, 1999). Furthermore, the SRP RNA is not continuously double-stranded over the length of the molecule. Remarkably, it is still possible to recognize a repeating feature in the reconstruction (Figure 1B, right, indicated by arrows), with an average spacing of 25Å, consistent with the helical repeat distance (*e.g.* the spacing between minor grooves) in the SRP-RNA model. This repeat pattern more closely resembles the predicted structure of SRP RNA than it does a strict RNA double helix consistent with the resolution obtained for our reconstruction (see Supplemental Material).

The possibility that there is some conformational heterogeneity within the SRP imaged means that the resolution obtained may not be the same for all areas of the image. For, example, the lower than expected mass in the tail region and the elliptical diameter of the tail suggest that
in the long dimension the resolution may be lower than other areas of the reconstruction.
Filtering the ec_SRP model to different resolutions provides an alternate method for estimating
the resolution of the SRP reconstruction and the extent to which the model represents
reconstruction. The SRP reconstruction (Figure 1B) clearly has more fine structural detail than a
19Å-resolution version of ec_SRP (Figure 3C, right). Also, the decreased density and increased
width of the tail of the reconstruction and the lack of correspondence of other high resolution
features preclude selecting the 7Å-resolution reference (Figure 3C, left) as the model that most
closely resembles the reconstruction. Therefore, the resolution of the model that most closely
resembles the SRP reconstruction is one that has been uniformly reduced to approximately 12Å
(Figure 3C, middle), consistent with the value obtained by FSC at the 3σ threshold. However,
the width of the tail of the reconstruction is not consistent with this model but it is consistent
with mobility of this region in the alignment used for the reconstruction. A more detailed
comparison with different views of the filtered models is consistent with this conclusion (see
Supplemental Material) and supports the relative domain organization in our model (Figure 3A).
The main structural features that constrained domain organization were the phosphorus map and
an approximately 40Å linker between NG and M. Other features were secondary to these
constraints as described above and in the Supplemental Material. During the modeling process
the domain organization reported in other structural analyses of SRP was not used as a
constraint.

We compared ec_SRP to the domain organization reported for structures of full-length
SRP54 from S. solfataricus (56% similarity to E. coli Ffh) and Ffh from T. aquaticus (65%
similarity to E. coli Ffh) post hoc. The structures of the NG and M domains are all similar with
the main differences being their relative positions allowing superposition of the three models (Figure 3D). The structure of S. solfataricus RNA-bound SRP54 revealed that NG and M adopted an L-shape orientation, with the SRP RNA fragment extending from the M domain, giving the whole particle a U-shape. Due to the unresolved NG-M linker of the T. aquaticus apo-Ffh structure (PDB: 2FFH), three potential configurations are generated which could represent full-length T. aquaticus Ffh: A/A (composed of A chain NG and A chain M from the PDB coordinates, respectively), B/A and C/A. In Figure 3D, the B/A domain organization for T. aquaticus is shown, as it is the most consistent with the structure of S. solfataricus SRP54. Comparison of our model for Ffh to the S. solfataricus and T. aquaticus structures (Figure 3D) demonstrates that the organization of M is different in each of the three models but all three can be rationalized by movement of the M domain as a rigid body on a relatively rigid linker arm that is flexible at each end. This is in agreement with a previous proposal that there is flexibility within the NG-M linker (Rosendal et al., 2003) but significantly expands the range of movement to accommodate the S. solfataricus and T. aquaticus configurations. Although the range of motion appears large it is confined to one area on one side of the NG domain (see below).

FRET measurements suggest flexibility in the spatial arrangement of NG and M of Ffh in solution

Fluorescence resonance energy transfer represents an independent method to explore the relative positioning of the NG and M domains of Ffh. Two residues within the N region, R11 and R49, and M347 in the M domain were substituted with a tryptophan (one at a time) to serve as a donor fluorophore. Residues Q340 and S366 in the M domain and R141 in the G region were mutated
to cysteine for labeling with 7-diethylamino-3-(4'-maleimidylphenyl)-4-methylcoumarin (CPM), a fluorophore that acted as an acceptor moiety. The protein was bound to 43 nt of SRP RNA and the extent of FRET was calculated for the various donor-acceptor pairs by measuring the time-resolved intensity decays of Trp11 and Trp49 (as described in Supplemental Material).

The average spatial separation between two sites was evaluated by fitting the lifetime data to a FRET model that assumed a Gaussian distribution of distances. The goodness of fit was estimated by $\chi^2_R$ and was within the range of 1.2 - 1.4. Surprisingly, the Gaussian width was broad with an average value of 22-25 Å strongly suggesting that the distance between M and NG is not fixed in solution. The distance measurements obtained between the respective residues within ec_SRP are in good agreement with the FRET data (Table 1, Figure 4C).

Since multiple positions within Ffh were analyzed by FRET, the distance measurements can be used to provide more complex constraints than one-dimensional measurements. To evaluate the FRET data in three dimensions, individual distances were represented by spheres centered at the alpha-carbons of the appropriate residues within the NG domain (Figure 4). For simplicity, the distances between the donor at residue 11 and the acceptors at residues 340 and 366 were approximated as being equidistant, generating a single sphere centered at residue 11 (Figure 4A, blue sphere). The same assumption was made for the distances between residue 49 and residues 340 and 366 (Figure 4A, green sphere). The intersection of two spheres is a circle whose equation can be solved. This intersection circle was displayed as a toroid (i.e. a circle with 3-D thickness) to account for various sources of uncertainty in the distance measurements such as the broad Gaussian width distribution obtained for the FRET data, the simplification of the distances between the donor residues and residues 340 and 366 and, measurements of the Ffh
structure from alpha-carbon atoms. This toroid (Figure 4B, grey) designates the potential space in which residues 340 and 366 of the M domain are localized. The lack of FRET between residues 141 and 347 defines a third sphere (Figure 4B, red sphere) of prohibited space for residue 347 of the M domain. The distances between the residues used for the FRET measurements are in good agreement with \textit{ec}_SRP (Figure 4C). Significantly, the relative orientations of M in other Ffh homologues (described above) are located close to the grey toroid defined by the FRET data (Figure 4D). This result is consistent with movement of M relative to NG being confined to this region. The \textit{T. aquaticus} configuration fits less well, with the majority of the M domain positioned in the hole of the toroid, so the \textit{S. solfataricus} and \textit{ec}_SRP configurations could define the minimal region of movement but this region could be expanded to include the \textit{T. aquaticus} configuration.

**Discussion**

SRP is a key ‘adaptor’ molecule, performing many linked functions: signal sequence recognition, ribosome binding and regulated release of a nascent chain to the membrane translocation machinery. To understand these roles requires knowledge of the structural states of SRP and identification of the sites of interaction with its binding partners. Here we fit the structures of domains and fragments of SRP from different organisms to the reconstruction of SRP calculated from STEM micrographs to obtain a model for the complete structure of \textit{E. coli} SRP.

Consistent with other structural analyses of SRP, the different domains of SRP could be fit into our reconstruction for \textit{E. coli} SRP only if the relative orientation of NG and M was changed (Figure 3D). Thus, the relative positioning of NG and M is different in every structure
for Ffh and SRP yet solved. These differences may reflect bona fide differences between species or they may indicate a common flexibility among all SRP exacerbated (or revealed) by the different specimen preparations, such as crystal packing, freeze drying on carbon or other conditions. Although there are clearly differences between SRPs from different organisms, the latter case is strongly supported by the broad FRET distance distributions between two fluorophores on the different domains of Ffh. The distance distributions imply that a dynamic range of Ffh conformations, possibly multiple discrete conformations, was sampled which could be conferred by a flexible interdomain linker that permits motion of the M-domain within a limited region (Figure 4). Flexibility is also consistent with recent observations suggesting that the M and NG domain organization changes for E. coli Ffh when it binds RNA (Buskiewicz et al., 2005a). Assuming conformational flexibility exists, the different crystal structures might result from the different specimen preparations or crystallography conditions used for structure determination. Consistent with this possibility, a cross-linking study of the binding of Ffh and FtsY in T. aquaticus suggests an NG-M domain organization that is not compatible with that obtained by crystallography (Chu et al., 2004). These results confirm that M can move relative to NG in SRP, a conclusion that could only be drawn by comparing several different SRP structures obtained from a variety of environments and using different techniques.

Some features of the SRP reconstruction suggest that the RNA has different conformations, as well. The tail of the SRP reconstruction is wider than the SRP RNA in ec_SRP (Figure 3B, see Supplemental Material for further discussion). Also, the density in the tail appears lower than in the head region (Figure 2A) even though double-stranded nucleic acid should be more dense than protein. Washout from the averaging of slightly different
conformations due to local flexibility of the RNA tail of SRP would account for the wider and
less dense tail of the SRP reconstruction. Quantum noise from the fewer number of images from
the direction depicted in Figure 1B, left, could reduce the accuracy of the alignment of this
region of the image and thereby may also contribute to the lower recorded density in the tail of
the reconstruction. Nevertheless, the pattern of intensity expected for double-stranded RNA was
discernable in the narrow view (Figure 1B, right) suggesting that although there was
conformational variation within the images there was a dominant conformation (see
Supplemental Material for further discussion). Consistent with the interpretation that the RNA
changes conformation, the accessibility of nucleotide 27 (located in the kinked region of
ec_SRP) to chemical reagents changes when Ffh binds RNA (Buskiewicz et al., 2005a). The
carbon film specimen support for the STEM imaging had been treated by glow-discharge and
incubated with a magnesium-containing buffer prior to SRP addition in order to increase the
hydrophilicity of the carbon film. The additional charges on the carbon film may have favored
certain conformations of the SRP complex that optimized its contact with the charged surface.
Thus the conformational flexibility of the RNA, especially around nucleotide 27, may be even
greater in solution than recorded in our micrographs. Although it is also possible that adsorption
of the SRP molecules to the carbon film may have induced conformational changes within the
complex, previous studies of electron micrographs of protein molecules of known structures on
carbon film demonstrated that such perturbations were minimal (Andrews and Ottensmeyer,
1982; Harauz et al., 1983).

It is somewhat counterintuitive that a molecule with such a dynamic range of motion
would result in a 3-D reconstruction that represents a single, discrete conformation when
displayed as a hard surface. The major indication that the reconstruction comes from a heterogeneous series of particles is the altered dimension and density of the RNA tail. That even this region has well defined edges indicates that the particles bound the carbon film as a series of discrete structures. The refinement algorithm used to reconstruct the SRP volume (primarily projection matching) causes the reconstruction to converge on the most common structure present within the population of STEM images. By aligning common elements within the structures of the various particles a conformation that is more highly represented in the image population dominates the reconstruction and mobile structural elements will be retained at their positions in the dominant structure while, at other locations, the same structures are averaged out and disappear. Consistent with this, detailed analysis of STEM micrographs of the Ffh protein in the absence of SRP RNA using a variety of reconstruction algorithms revealed an Ffh conformation, similar to the *T. aquaticus* A/A configuration, in almost 60% of the images while a structure resembling the *S. solfataricus* NG-M configuration, was present in 30% of the images. Nevertheless, reconstruction of the entire image population resulted in a single final reconstruction similar to the more abundant one (Mainprize et al., MS in preparation). Therefore, the structure that we have obtained for *E. coli* SRP represents the most abundant conformation present within our image population and other conformations present are either lost or evidenced only by areas of lower than expected density due to the averaging effect of the refinement procedure (See Supplemental Material for further discussion).

In contrast to our finding that the RNA in SRP extends away from the particle in a rod shape, recent FRET data for *E. coli* SRP in solution suggests that the distal half of the RNA folds back onto the protein (Buskiewicz *et al.*, 2005b). It has already been shown by cross-linking to
the large ribosomal subunit that the RNA of *E. coli* SRP adopts an extended conformation when bound to the ribosome (Rinke-Appel *et al.*, 2002) therefore previous FRET studies of SRP in solution have revealed a conformation inconsistent with ribosome-bound SRP. Nevertheless, since this region of SRP RNA (nucleotides 1-32 and 75-114) has been demonstrated to be non-essential (Batey *et al.*, 2000) and contributes at most an accessory function, the precise localization of the distal half of the RNA is unlikely to change the structure of the rest of the particle in a functionally significant way.

The NG-M configuration, deduced from other FRET studies that indicated a folding back of the RNA of SRP in solution, was a more opened version of the *T. aquaticus* A/A configuration. Interestingly, the RNA in this NG-M configuration occludes the FtsY binding site so that further interdomain movement must occur to enable SRP to bind to its receptor. Thus it is possible that the RNA and the M domain are repositioned relative to NG in response to different stages of the SRP pathway. Consistent with this possibility, results from localized hydroxyl radical cleavage of *E. coli* SRP RNA indicated that there was at least one region of Ffh that was closer to the RNA tetraloop when SRP was bound to FtsY than in free SRP (Spanggord *et al.*, 2005). Although the sites of cleavage that were obtained for SRP alone are not inconsistent with our model, there are two sites in the SRP/FtsY complex where they did not see cleavage that are close enough in our model that they might have obtained cleavage. As most of the radical induced cleavage experiments included both SRP and FtsY the model they generated may be more relevant to the complex. Nevertheless the data is not inconsistent with the structure we obtained. The main difference between the *ec_SRP* model and the model they generated is the orientation of the NG domain which is rotated to a position similar to that in *Supplemental*
Material (Figure 4G, right).

SRP-Ribosome Interactions

To explore a possible mode of interaction of SRP with the ribosome, ec_SRP was docked onto the structure of the H. marismortui ribosome by using published cross-linking data for bacterial SRP on the large ribosomal subunit. The RNA was positioned using the location of an RNA cross-link between nucleotide U84 of E. coli SRP RNA and nucleotides 2828-2837 of 23S rRNA (Rinke-Appel et al., 2002) (red spheres, Figure 5A). The binding site for the N region of Ffh was identified based on the cross-linking between two residues in the N region and the L23 protein of the 50S subunit using a cross-linker with a 10Å spacer arm (Gu et al., 2003) (Figure 5A, light yellow). These two sets of cross-linking data provide constraints that align our reconstruction onto the large ribosomal subunit (Figure 5B) such that the particle runs alongside the nascent chain exit site (Figure 5A, black circle) with M aligned at the location of closest approach to this site. These points clearly fix the location and orientation of the long axis of SRP on the ribosome. To our surprise it was only possible to position SRP on the ribosome such that the Ffh M domain is located on the side of SRP opposite from the ribosome (Figure 5D).

Furthermore, the majority of the potential contacts between E. coli SRP and the ribosome appear to be nucleic acid-based. Unexpectedly, the kink in the RNA tail of SRP extends the distal half of SRP RNA away from the ribosome (Figure 5C) where it may recruit non-essential accessory factors. Although the model is placed at a precise angle with respect to the ribosome, there is considerable latitude possible in this angle that would still be consistent with our model and the known biology of SRP. The primary constraint used was not occluding the FtsY binding site on
SRP while minimizing the distance between SRP and the nascent chain exit site on the ribosome.

To obtain insight into the possible functional significance of the interdomain movement of SRP, the structure of *E. coli* trigger factor (TF), another protein known to bind the ribosome adjacent to the nascent chain exit site, was added to our model of SRP docked to the ribosome. The model of ribosome-bound TF was generated as explained previously (Raine *et al.*, 2004). Significantly, the model that resulted from docking the structures of TF and SRP independently explains how both TF and SRP can bind ribosomes simultaneously, such that both can access the nascent chain exit site of the ribosome (Figure 5C).

Consistent with the model in Figure 5C, cross-linking between TF and Ffh was seen only when both were bound to ribosomes (Buskiewicz *et al.*, 2004). Furthermore, the cross-linker in TF was positioned at a site on the side of TF closest to Ffh in our ec_SRP model from which the 10Å spacer arm of the cross-linker could easily bridge between the two proteins. The observation that the cross-linking pattern between Ffh and L23 of the large ribosomal subunit was altered upon TF binding to the ribosome is also consistent with the two binding sites being in close proximity.

In the absence of TF, *E. coli* SRP has been shown to bind virtually any nascent chain as it emerges from the ribosome (Beck *et al.*, 2000), suggesting that SRP may sample all nascent chains not bound by TF. Although the location of SRP on the ribosome is consistent with access to nascent chains, SRP binding to nascent chains in the absence of TF is somewhat artificial as there is good evidence that most, if not all, cytoplasmic ribosomes include bound TF. Moreover, most evidence suggests that only a subset of nascent chains (*e.g.* primarily polytopic membrane proteins) is bound by SRP. Nevertheless, the implication of these findings is that SRP and TF
must both access the nascent chain exit site in order to sample nascent chains as they emerge from the ribosome.

_A model for nascent chain sampling by TF and SRP_

The ribosome-docked _ec_SRP_ suggests that the M domain of Ffh, containing the primary signal sequence binding site, is relatively far from the nascent chain exit site. Thus, nascent chains emerging from the ribosome would first encounter TF and any chains with an affinity for TF would be drawn away from SRP. For SRP to sample nascent chains with low affinity for TF we speculate that interdomain movement within SRP repositions M closer to the nascent chain exit site. Movement of M into and out of the nascent chain exit site would permit SRP to compete with TF for nascent chains containing hydrophobic segments and may also account for why nascent chain binding by SRP and TF often appears mutually exclusive as movement of M back to the position seen in Figure 5D would draw the nascent chain away from TF.

To move M into the nascent chain exit region requires only that the RNA molecule be rotated 90° about the long axis as shown in Figure 5E. This movement of M is consistent with the FRET data as the movement of the M-domain is within the area permitted by the toroid (Figure 4C). More efficient rotation of M may be the elusive role for the RNA in _E. coli_ SRP. If the primary function of the SRP RNA in _E. coli_ is to move M relative to NG then our model showing interactions of SRP to the ribosome for only one half of the RNA may also explain why _E. coli_ can survive with a shortened SRP RNA. Although the SRP RNA is positioned along the ribosome surface it makes relatively few contacts, therefore rotation might be accomplished by the conformational change that occurs in ribosomes translating hydrophobic sequences.
Rotation of M may also result from or regulate GTP binding by SRP. Alternatively (or in addition), this conformational change within SRP could act as a switch to regulate downstream events in the SRP pathway, such as the binding of SRP to FtsY. The binding site of TF on the ribosome is the same as that implicated for binding of the ribosome to the translocation machinery so TF must be removed to clear the translocon binding site. Consistent with one function of FtsY being the displacement of TF from the ribosome (Buskiewicz et al., 2004), incorporating FtsY into our ec_SRP-ribosome model locates the SRP-bound FtsY such that it could directly displace TF (Figure 5F). Interestingly, movement of the G region relative to the N region of Ffh, which is implied by the variation between the structure of apo-Ffh NG and the structure of Ffh NG bound to FtsY (Freymann et al., 1997; Focia et al., 2004), would place the maximal displacement of the G region in a direction such that the proposed signal sequence binding site in SRP would be opened up upon FtsY binding (see Supplemental Material).

One of the main strengths of our ec_SRP model is that it can be used to generate novel, yet testable, predictions regarding the interaction of SRP with both the ribosome and TF. Similar STEM and ESI data was used to predict a mechanism of binding of canine SRP to eukaryotic ribosomes that almost 20 years later was shown to be largely correct (Andrews et al., 1985; Andrews et al., 1987; Halic et al., 2004). Therefore, to complete our understanding of how the various components of the SRP pathway interact and function, additional SRP structures will be required. However, because our model is consistent with structural data currently available, it will be an important tool for analyzing the results of future biochemical and structural investigations of bacterial SRP and for identifying the ligand-induced conformational changes.
that are believed to be the basis for the targeting function of SRP.
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**Figure 1.** STEM and ESI imaging of *E. coli* SRP. A. A representative ‘low-dose’ STEM micrograph of a carbon support containing *E. coli* SRP (left). Putative particles of SRP were extracted as individual images (right) for further structural analysis. Scale bar, 100Å. B. Two views of the reconstruction of *E. coli* SRP generated from low-dose cryo-STEM data. Black arrows indicate repeating ridges in the reconstruction. The reconstruction has been filtered to 12Å and is displayed as a hard surface at a threshold corresponding to approximately 90 kDa. This image display generates sharp edges not expected at the resolution obtained but produces a more interpretable image. C. Two possible locations for the RNA (blue bar) in the reconstruction of SRP (wireframe). D. Three-dimensional reconstruction representing the location of phosphorus atoms within SRP. The reconstruction is presented at the same two views as shown for the complete particle in B. E. The overlap of the SRP reconstruction and the 3-D phosphorus map indicating the relative position of the phosphorus within SRP. Scale bar, 50Å.

**Figure 2.** Determination of the domain organization of the *E. coli* Ffh in our SRP reconstruction. A. The SRP reconstruction visualized at a threshold that corresponds to the molecular mass of *E. coli* SRP (90 kDa, orange) overlaid onto an image at a second threshold that corresponds to 150% of the expected molecular mass (black). The reconstructions are presented at the same two views as in Figure 1. B. The structure of *E. coli* M (green) bound to a fragment of SRP RNA (PDB: 1DUL) was aligned by the RNA component to the region defined by ESI. For illustrative purposes the M domain is shown in four different positions (1-4) related to each other by iterative rotation of 90° about the RNA long axis. The SRP reconstruction (black lines) was visualized at 150% of the expected molecular mass, indicative of the largest
possible volume of the particle. C. The M-RNA structure at position #1, from B, with the NG domain (from *T. aquaticus*, PDB: 2FFH, red) fit into the remaining volume of the reconstruction. The same two orientations of the structure are shown as in B. The N region is situated above the RNA and the G region is above the M domain. Details for the fitting of the NG domain are provided in the Supplemental Material. D) The completed model compared to the reconstruction. The model was constructed from the structure of *T. aquaticus* NG (PDB: 2FFH, red) combined with the structure of *S. solfataricus* M (PDB: 1QZW, green) bound to a fragment of SRP RNA (blue). The remaining SRP RNA (blue) was based on a model of extended *E. coli* SRP RNA (SRP Database). The model is fit into the more conservative reconstruction threshold, corresponding to 90 kDa (orange wireframe), and shown in similar views to those in C.

**Figure 3.** Comparison of the *ec_SRP* model with the reconstruction of *E. coli* SRP. A. Ribbon diagram representation of *ec_SRP* (coloured as in Figure 2D), highlighting potential regions of interest, including the N, G and M regions as well as the tetraloop (t) of the SRP RNA. The amino-terminus of NG and the carboxyl-terminus of M are indicated by blue circles. A black oval indicates the approximate location of the proposed signal sequence binding region of the M domain and the guanine nucleotide binding site of NG is indicated by a green square. B. Alignment of the *ec_SRP* model into the reconstruction of *E. coli* SRP, from Figure 2D. C. The *ec_SRP* model was filtered to three different resolutions (*Left*, 7Å; *Middle*, 12Å; *Right*, 19Å) and displayed at respective threshold values to generate hard surface representations corresponding to approximately 90 kDa (see Supplemental Material for additional views and direct comparison to the reconstruction). D. The relative orientations of M in *ec_SRP* (green),
SRP54 from *S. solfataricus* (PDB: 1QZW, blue) and the B/A configuration of *T. aquaticus* apo-Ffh (adapted from PDB: 2FFH, purple). The slight differences in the respective NG domains are similar to the amount of variation seen in the NG domain from a single organism (*T. aquaticus*) depending on guanine nucleotide occupancy of the G region. For simplicity, only one of the aligned NG domains is shown (grey). The major difference between the three structures is the relative locations of the M domain. To represent a complete polypeptide backbone for the *T. aquaticus* structure, an energy minimized 11 amino acid NG-M linker has been modeled. To show the approximate location of the NG-M linker in the Ffh component of ec_SRP residues 298-318 of *S. solfataricus* SRP54 were positioned between NG and M of ec_SRP.

**Figure 4.** Three-dimensional analysis of FRET-determined interdomain distances. A. Distances obtained by FRET analysis represented as spheres centered at residue 11 (radius 30Å, large blue sphere) and residue 49 (radius 44Å, large green sphere) of the NG domain. The smaller colored spheres indicate the positions of Tryptophan donors at residues 11 and 49. B. A circle representing the intersection of the two spheres from A was calculated and is displayed as a grey toroid. A sphere of radius 50Å (red) was centered on the residue corresponding to 141 in *E. coli* Ffh to represent prohibited space for residue 347 implied from the lack of FRET between these two positions. C. The NG-M configuration defined by ec_SRP is shown with the distances measured from the alpha-carbon atoms of the residues corresponding to the donors and acceptors (Table 1). The arrow is displayed going from the donor position to the acceptor position. The respective residues in M co-localize well with the intersection of the FRET-defined interdomain distances (grey toroid). *Left*, same view as that in Figure 3D. *Right*, after a rotation of 90° about
the Y-axis. D. Comparison of M orientations (colored as in Figure 3D) with the FRET data. Left, same view as that in Figure 3D. Right, after a rotation of ~70° about the Y-axis. Each of the different M domains is localized to the same region defined by the FRET-derived distances. E. The position of M derived from rotation of the SRP RNA by 90° to model signal sequence binding (see Discussion) is also consistent with the FRET data.

**Figure 5.** Docking SRP on the ribosome. A. Surface of the 50S ribosomal subunit from *H. marismortui* (PDB: 1W2B) highlighting subunits L23 (light yellow) and L29 (dark yellow) and the nascent chain exit site (black circle). The nucleotides which correspond to a region of 23S rRNA found to cross-link to bacterial SRP RNA are displayed as red spheres. B. Alignment of the SRP reconstruction (orange) onto the 50S ribosomal subunit maximizing the overlap of regions of SRP with respective cross-linking sites on the ribosome (L23 and rRNA). C. The structure of *E. coli* TF (pink) modeled onto the ribosome along with the SRP reconstruction. TF and SRP are in very close proximity to each other and flank the nascent chain exit site. A second view related to the first by rotation of 90° illustrates the effect of the kink in the RNA structure. D. The SRP reconstruction was replaced with the *ec* _SRP_ model and demonstrates the orientation of M (green) with respect to the other domains of SRP, the large ribosomal subunit and TF. The SRP RNA, NG and M domains are colored as in Figure 3B. E. Rotation of the SRP RNA by 90° moves the M domain of SRP in towards the nascent chain exit site where it can effectively compete with TF for sampling nascent chains as they emerge from the ribosome. F. The structural data of the binding of the NG domain of the SRP receptor, FtsY (PDB: 1RJ9, blue), to *E. coli* Ffh NG was incorporated onto the ribosomal subunit. This location of FtsY is
likely to result in the displacement of TF from the ribosome explaining mutually exclusive
binding of TF and FtsY to ribosome-SRP complexes. Two views related by rotation of 90° are
shown.
Table 1. Spatial separation between different donor-acceptor pairs of *E. coli* Ffh as found in free SRP in solution compared to measurements of various Ffh models

<table>
<thead>
<tr>
<th>Protein</th>
<th>Donor (res. #)</th>
<th>Acceptor (res. #)</th>
<th>Donor-Acceptor Distance, Å</th>
<th>ec SRP&lt;sup&gt;b&lt;/sup&gt;</th>
<th>S. solfataricus&lt;sup&gt;b&lt;/sup&gt;</th>
<th>T. aquaticus&lt;sup&gt;b&lt;/sup&gt;</th>
</tr>
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<tr>
<td>Ffh* R11W/Q340C-CPM&lt;sup&gt;a&lt;/sup&gt;</td>
<td>11</td>
<td>340</td>
<td>30</td>
<td>32</td>
<td>28</td>
<td>29</td>
</tr>
<tr>
<td>Ffh* R11W/S366C-CPM</td>
<td>11</td>
<td>366</td>
<td>29</td>
<td>32</td>
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<td>366</td>
<td>45</td>
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<td>56</td>
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<tr>
<td>Ffh* M347W/R141C-CPM</td>
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<td>141</td>
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<td>42</td>
<td>65</td>
<td>73</td>
</tr>
</tbody>
</table>

a. Amino-terminal fusions of Ffh with GB1. Residues are numbered as in wild-type *E. coli* Ffh.

b. Distance measurements for the Ffh structures/models were made from the alpha-carbon of the respective residues using InsightII. This simplification introduces an uncertainty in the measurement of approximately +/- 3Å.

c. No FRET was detected. Distances of >50Å are not resolved by this method.

d. There is an uncertainty of +/- 3Å in the FRET-determined distance measurements due to an approximate 10% uncertainty in the orientation factor (see Supplemental Material).