Allosteric Response and Substrate Sensitivity in Peptide Binding of the Signal Recognition Particle

Connie Y. Wang* and Thomas F. Miller III

From the Division of Chemistry and Chemical Engineering, California Institute of Technology, Pasadena, California 91106

Background: The SRP is a central component of the co-translational protein targeting pathway. Long timescale computer simulations reveal shifts in the SRP conformational distribution upon nascent protein binding. The binding-induced conformational shifts correlate with the experimentally observed efficiency of protein targeting. The work provides new insight into the mechanism by which SRP allosterically regulates the fidelity of protein targeting.

Results: Long timescale computer simulations reveal shifts in the SRP conformational distribution upon nascent protein binding. The binding-induced conformational shifts correlate with the experimentally observed efficiency of protein targeting. The work provides new insight into the mechanism by which SRP allosterically regulates the fidelity of protein targeting.

Conclusion: The binding-induced conformational shifts correlate with the experimentally observed efficiency of protein targeting. The work provides new insight into the mechanism by which SRP allosterically regulates the fidelity of protein targeting.

Significance: The work provides new insight into the mechanism by which SRP allosterically regulates the fidelity of protein targeting.

We characterize the conformational dynamics and substrate selectivity of the signal recognition particle (SRP) using a thermodynamic free energy cycle approach and microsecond timescale molecular dynamics simulations. The SRP is a central component of the co-translational protein targeting machinery that binds to the N-terminal signal peptide (SP) of nascent proteins. We determined the shift in relative conformational stability of the SRP upon substrate binding to quantify allosteric coupling between SRP domains. In particular, for dipeptidyl aminopeptidase, an SP that is recognized by the SRP for co-translational targeting, it is found that substrate binding induces substantial changes in the SRP toward configurations associated with targeting of the nascent protein, and it is found that the changes are modestly enhanced by a mutation that increases the hydrophobicity of the SP. However, for alkaline phosphatase, an SP that is recognized for post-translational targeting, substrate binding induces the reverse change in the SRP conformational distribution away from targeting configurations. Microsecond timescale trajectories reveal the intrinsic flexibility of the SRP conformational landscape and provide insight into recent single molecule studies by illustrating that 10-nm lengthscale changes between FRET pairs occur via the rigid-body movement of SRP domains connected by the flexible linker region. In combination, these results provide direct evidence for the hypothesis that substrate-controlled conformational switching in the SRP provides a mechanism for discriminating between different SPs and for connecting substrate binding to downstream steps in the protein targeting pathway.

Protein targeting pathways govern the delivery of both secretory and integral membrane proteins to their appropriate cellular destinations (1). Co-translational protein targeting relies on the recognition of the N-terminal signal peptide (SP) of nascent protein sequences. This process requires the signal recognition particle (SRP), a protein-RNA complex, and universally conserved component of the co-translational protein targeting machinery (2). The SRP binds to the ribosome/nascent-protein complex (RNC) and targets nascent proteins for either translocation or membrane integration via the Sec translocon (3–9). Nascent proteins that undergo targeting via the co-translational pathway exhibit h - helical secondary structure (10, 11). However, the detailed mechanism by which SP binding triggers selection for co-translational targeting remains unclear. In this work, we use free energy (FE) calculations and microsecond timescale trajectories to investigate the coupling between SP binding and SRP conformational dynamics and to provide new insight into the role of SP binding in co-translational protein targeting.

Both structural (12–21) and biochemical work (22–24) suggest that the SRP exhibits multiple stable conformations that are important for protein targeting. The conserved functional core of the SRP (Fig. 1A) is composed of an RNA component and a multidomain protein component (25, 26). The protein component consists of a methionine-rich M domain that contains both the SP-binding site and the RNA-binding site, as well as an NG domain that contains a catalytic GTPase (27–29). The M and NG domains are connected by a flexible 30-residue linker region (20). Fig. 1, A and B, represents two conformations of the SRP that have been proposed to play a central role in the initial step of SP binding to the SRP (19). Recent crystal structures reveal that although the structure of the individual domains is quite similar, the relative arrangement of the domains differs substantially between the two conformations (19, 20). Throughout this paper, we refer to these two conformations as syn and anti, indicating the relative orientation of the SRP RNA and the NG domain.

3 The abbreviations used are: SP, signal peptide; SRP, signal recognition particle; RNC, ribosome/nascent-protein complex; FE, free energy; MD, molecular dynamics; FEP, free energy perturbation; LJ, Lennard-Jones; PB, Poisson-Boltzmann; DPAP, dipeptidyl aminopeptidase.

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Allosteric Response to Signal Peptide Binding

SRP conformational changes upon SP binding and reduces the demands of protein conformational sampling. We calculate the allosteric effect induced by SP binding for a range of experimentally studied SPs to investigate the effect of changing the SP amino acid sequence. Furthermore, we reveal details of the large scale SRP conformational rearrangements by performing several microsecond timescale (MD) trajectories. Taken together, these calculations provide insight into the sensitivity of the SRP conformational changes to the identity of the SP. In particular, the results of this study reveal that the conformational preference of the SRP for the anti over the syn conformation upon SP binding agrees with trends found in experimentally measured targeting efficiencies, suggesting that SRP conformational dynamics are substrate-specific and provide a selection mechanism of nascent proteins for the co-translational targeting pathway.

COMPUTATIONAL METHODS

Modeling for FE Calculations—Initial coordinates for the syn conformation of the SRP are available from the high resolution crystal structure of the archael Sulfolobus solfataricus species (Protein Data Bank code 1QZW) (20). Coordinates for the anti conformation of the SRP from S. solfataricus have not been reported in a crystallographic study, so they are instead modeled from the available M. jannaschii structure (Protein Data Bank code 3NDB) (19) as follows. A pairwise sequence alignment of the S. solfataricus and Methanococcus jannaschii sequences is performed using ClustalX (50) and input into MODELLER (51). The MODELLER protocol builds coordinates for the S. solfataricus SRP in the anti conformation that minimizes a set of the spatial restraints generated from the sequence alignment with the existing M. jannaschii structure and a set of stereochemical restraints determined from the CHARMM22 force field (52). The model coordinates are refined with conjugate gradient minimization and molecular dynamics with simulated annealing.

The FE associated with SP binding to either the syn and anti conformations of the SRP is calculated for three SPs, listed in Table 1. The first is dipeptidyl aminopeptidase B (DPAP-WT), a prototypical example of a peptide sequence that is targeted to the co-translational targeting pathway (53). The second SP, DPAP-K464L, is obtained from the DPAP-WT sequence by a single lysine-to-leucine mutation at residue 464. This mutation extends the hydrophobic region of the SP relative to the DPAP-WT. The third SP is PhoA, which is found to bind SRP in vivo systems alternatively select PhoA for the post-translational SecB pathway (54). A crystal structure of the DPAP-WT SP bound to the syn conformation of the SRP is available (Protein Data Bank code 3KL4) (17) and is used to initialize coordinates for simulations of the bound DPAP-WT. The coordinates for bound DPAP-K464L are constructed via direct substitution of the leucine side chain coordinates in DPAP-WT at residue 464 with lysine coordinates, followed by steepest descent minimization to remove steric clashes (55). The coordinates for bound PhoA are constructed by sequence alignment with the DPAP-WT SP and modeled by the MODELLER protocol described previously. We note that the N-terminal end of the SP sequences are capped using a...
neutral acetyl group, rather than a positively charged bare ammonium terminus, to account for the fact that in the experimental crystal structure (17) the N terminus of the SP sequence is bonded to additional residues that are not resolved; and for the simulations to include a bare ammonium terminus at the N-terminal end of the SP would introduce an unphysical charge at that point in the sequence.

Free Energy Sampling—We performed nine sets of calculations to determine the FE of binding for each of the three SPs to the SRP in the syn or anti conformation. For the ith SP, these calculations yield, ΔG(1y)G(a)G(I) and ΔG(1y)G(a)G(I), which are, respectively, the FE of aqueous solvation for the SP, the FE associated with transferring the SP from the vacuum state to binding the solvated SRP in the syn conformation, and the FE associated with transferring from the vacuum state to binding the solvated SRP in the anti conformation. These quantities are then used to compute the FE of SP binding, ΔG(I/y)G(I) = ΔG(1y)G(a)G(I) − ΔG(1y)G(a)G(I). In this study, each calculation of a FE difference was performed using the FE perturbation (FEP) method in two stages.

The first stage involves “turning on” the Lennard-Jones (LJ) interactions between the SP and the remaining atoms in the system. The second stage involves turning on the electrostatic interactions between the SP and the remaining atoms. For each stage, independent sampling trajectories were performed with potential energy functions that correspond to different values of the parameter $\lambda$, as shown in Equation 1,

$$V(\lambda) = (1 - \lambda)V_0 + \lambda V_1$$

(Eq. 1)

For the first stage, in which the LJ interactions are turned on, $V_0$ refers to the noninteracting state, and $V_1$ refers to the state with fully interacting LJ potentials. To ensure the use of smooth potential energy functions during this process (56, 57), the LJ potential energy function was turned on using a $\lambda$-dependent soft-core potential for $V_1$, as implemented in Gromacs 4.5.3 (58) with $\alpha = 0.5$, $\sigma = 0.3$, and $p = 1$; the full LJ potential energy function was recovered for $\lambda = 1$. For the second stage, $V_0$ refers to the state with only LJ interactions (i.e. the electrostatically noninteracting state), and $V_1$ refers to the state with both LJ and electrostatic interactions. The first stage utilizes 10 trajectories corresponding to evenly spaced values of $\lambda$ between 0 and 0.09, 30 trajectories corresponding to evenly spaced values of $\lambda$ from 0.1 to 0.245, and 76 trajectories corresponding to evenly spaced values of $\lambda$ from 0.25 to 1. The second stage utilizes 91 trajectories corresponding to evenly spaced values of $\lambda$ from 0 to 0.9 and 20 trajectories corresponding to evenly spaced values of $\lambda$ between 0.91 and 1.

All FE calculations of SP binding to the SRP are performed on a truncated version of the SRP system that consists of the linker region and the M domain (residues 296–432); the NG domain and the RNA are removed to reduce computational cost (Fig. 2). Consideration of this truncated system is warranted by the fact that all available crystal structures indicate that the SP binding pocket is fully contained within the combined M domain and linker region (17, 19); microsecond simulations of the untruncated system are discussed below. We note that previous experimental studies have focused on the binding of SRP to the full RNC complex, whereas the current simulation only considers the binding of the SP to the SRP portion of the RNC complex; this study thus addresses only the relative impact of the SP on the conformational changes of the SRP.

A truncated octahedral simulation cell is used with periodic boundary conditions. Na+ and Cl− counterions are included to achieve electroneutrality of the simulation cell at an ion concentration of 100 mM. The total system size varies for the SRP in the syn and anti conformation due to different box sizes and different numbers of solvent molecules. For FE simulations of SP binding to the SRP in the syn conformation, the total system size is 29,287 atoms. FE simulations of the SP binding to the SRP in the anti conformation have a total system size of 39,482 atoms. For simulations of the SP in an aqueous environment, the total system size is 9893 atoms. In all FE calculations, we employ the AMBER03 force field (59) and the TIP3P model for water (60).

Two sets of harmonic restraints were applied to limit the sampled configuration space for the SRP and SP molecules in the FE calculations. The first set restrains the SP to the SRP, to avoid dissociation of the SP from the SRP during the course of the long sampling trajectories. Specifically, using the PLUMED plugin (61), each SP was restrained to the SRP by six harmonic restraints between the Cα atoms of residues 4, 10, 14, 16, 4, and 16 of each SP to the Cα atoms of residues Leu-428, Met-424, Glu-416, Glu-412, Ala-334, and Lys-373 of the SRP M domain, respectively; the force constant of the harmonic restraints is 55.6 kJ mol$^{-1}$ nm$^{-2}$, and the distance of minimum restraint for each pairwise interaction corresponds to the crystal structure distance of the atom pair in the 3KLA structure (17). The second set of harmonic restraints limits the ensemble of sampled configurations for the SRP to those associated with either the syn or anti structures that are observed in the experimental crystal structures (19, 20). Specifically, for both the syn and anti conformations of the SRP, each Cα atom associated with residues in the linker region of the SRP (residues 296–330) is restrained in absolute space to its corresponding position in the experimental crystal structure, using a force constant of 1000 kJ mol$^{-1}$ nm$^{-2}$. For all sampling trajectories associated with a given FE calculation, the applied restraint potentials retain the same minimum and force constant, even as the other molecular interactions are alchemically modified as part of the FEP method.

For every sampling trajectory (each of which is associated with a different value of the $\lambda$ parameter in one of the two stages

### Summary of the sequences and features of studied SPs

| SP   | Sequence | V(Å$^3$)* | Features                      |
|------|----------|-----------|-------------------------------|
| DPAP-WT | GILVLWHTGVLLKKSHP | 3291 | Efficient co-translational targeting sequence |
| DPAP-K464L | GILVLWHTGVLLKSHP | 3178 | Single-point mutation from WT |
| PhoA | JALLLLPFTYPYKIA | 2338 | Inefficient co-translational targeting sequence |

* Excluded volume of the SP, was determined by VMD volmap (82).
of the FE calculation), the initial configuration of the system was subjected to both relaxation and equilibration before data collection. First, the steepest descent minimization (55) is performed to reduce steric clashes to a force tolerance of 100 kJ mol\(^{-1}\) nm\(^{-1}\). Second, equilibration in the NVT ensemble is performed for 3 ns. Third, for FE calculations with the SRP, we perform 3 ns of equilibration in the NPT ensemble; for FE calculations without the SRP (i.e. aqueous environment), we perform 1 ns of equilibration in the NPT ensemble before data collection. Relaxation, equilibration, and sampling are done within the FEP framework implemented in the Gromacs 4.5.3 package (58). The system is evolved using Langevin dynamics within the FEP framework implemented in the Gromacs 4.5.3 package (58). The system is evolved using Langevin dynamics with a damping constant of 1 ps\(^{-1}\) and a temperature of 300 K. The Parrinello-Rahman barostat (62) is applied for pressure coupling at 1 bar. The particle mesh Ewald method (63, 64) is used to evaluate the electrostatic interactions with a real space cutoff of 10 Å. The LJ interactions are switched off between 8 and 9 Å using a cubic spline, and a long range dispersion correction is applied to the energy and pressure during the course of the simulation to correct for the cutoff (65). The simulations are performed with a 2-fs time step with bond distances constrained using LINCS (58). For FE trajectories associated with turning on the SP interactions in the presence of the SRP, the sampling time for each trajectory ranges from 7 to 10 ns. For trajectories associated with turning on the SP interactions in the absence of SRP (i.e. aqueous environment), the sampling time was 4 ns. Over 12 µs of combined simulation time was performed for the FE sampling.

The Bennett acceptance ratio method (66) is used to bidirectionally weight the collected data from trajectories sampled in the \(i\) and \(i+1\) potential and to calculate \(\Delta G_{i+1}\), the FE difference between the \(i\) and \(i+1\) states. \(\Delta G_{i+1}\) is calculated by iteratively solving Equation 2

\[
e^{\beta \Delta G_{i+1}} = \frac{\langle f(-\beta (\Delta U_{i+1} - C)) \rangle_{i+1}}{\langle f(\beta (\Delta U_{i+1} - C)) \rangle_{i}} e^{\langle BC \rangle}
\]

\[
f(x) = \frac{1}{1 + e^x}
\]

\[
C = \Delta G_{i+1} + \frac{1}{\beta} \ln \frac{N_{i+1}}{N_i}
\]

where \(\Delta U_{i+1} = U_{i+1}(x_i) - U_i(x_i)\) is the potential energy of the configuration \(x_i\) sampled with the \(i\) potential and evaluated in the \(i\) potential. \(U_{i+1}(x_i)\) is the potential energy of the configuration \(x_i\) sampled with the \(i\) potential but evaluated in the \(i+1\) potential. \(N_i\) is the number of independently sampled points in trajectory \(i\).

For each FE calculation, the convergence of each sampling trajectory and the overlap between neighboring trajectories were examined by plotting the autocorrelation function of \(\Delta U_{i+1}(t)\) and the relative entropy, defined in Equation 3,

\[
D(p_i)(p_j) = \int p_i \ln \left( \frac{p_i}{p_j} \right)
\]

where \(p_F = P(\Delta U_{i+1})\) and \(p_B = P(\Delta U_{i+1})\) and where \(D = 0\) for identical distributions. In Fig. 3, we show these convergence and overlap measures for the case of DPAP-WT SP bound to...
SRP in the anti conformation, and similar results are seen for all other FE calculations in this study. In Fig. 3A, we find decorrelation times in the range of 1–900 ns that are used to determine the frequency of sampling for the FE calculations. In all cases, the decorrelation time for the individual trajectory is shorter than the total sampling time. The overlap between neighboring trajectories is shown in Fig. 3B and sufficient windows are added such that $D(p_f(p_j)) \leq 1.5$ to ensure good overlap (67).

To provide an alternative test of the overlap, we use the overlapping histogram method of Bennett (66), which plots $g_f(\Delta U) - g_B(\Delta U)$, as defined in Equation 4,

$$g_f(\Delta U) = \ln p_f(\Delta U) - \frac{1}{2} \beta \Delta U$$

(Eq. 4)

$$g_B(\Delta U) = \ln p_B(\Delta U) + \frac{1}{2} \beta \Delta U$$

In Fig. 3C, $g_B(\Delta U) - g_f(\Delta U)$ is compared with the value of $\beta \Delta G$ calculated via the Bennett acceptance ratio method. For trajectories with sufficient overlap and correct sampling, $g_f(\Delta U) - g_B(\Delta U) = \beta \Delta G$ over a range of $\Delta U$ values, where $p_f(\Delta U)$ and $p_B(\Delta U)$ have significant overlap.

Additional analysis was performed to avoid possible artifacts in the reported FE calculations due to long range electrostatics. The DPAP-WT sequence is the only SP considered in this study that involves a net charge; the SP sequences PhoA and DPAP-K464L are both neutral. Care must be taken when using the particle mesh Ewald description of long range electrostatics to calculate relative free energies that involve the manifestation of a net charge (such as $\Delta G^{\text{DPAP-WT}}_S$ and $\Delta G^{\text{DPAP-WT}}_A$), because calculations of this kind can be sensitive to the size of the total simulation cell (68). Although one might expect this effect to be small in simulations as large as those performed here, which range from 29,287 to 39,482 atoms in size, we have nonetheless performed two robustness tests to explicitly confirm that such system-size artifacts do not impact the results presented here. In the first robustness test, we utilize the method of Lin et al. (68) to compute the leading order correction to the FE differences due to long range electrostatics (Equation 22 of Ref. 68), which accounts for the shift in the electrostatic potential due to the finite system size of the simulation cell. The calculated corrections are 1.89 and 1.71 kJ/mol for $\Delta G^{\text{DPAP-WT}}_S$ and $\Delta G^{\text{DPAP-WT}}_A$, respectively, such that the relative shift in these quantities due to the system-size effect is only 0.18 kJ/mol; because the relative FE of SP binding to the SRP in the syn versus the anti conformation is sensitive only on this small relative shift, the system-size effect associated with the polarization constraint is concluded to be negligible. In summary, the results from both tests suggest that the conclusions drawn in this paper are robust with respect to possible system-size effects associated with long range electrostatics.

**Microsecond Timescale Trajectories of the Untruncated SRP—**

Microsecond timescale trajectories of the SRP system are performed on the Anton supercomputer, a special-purpose machine for long timescale MD simulations (70, 71). Unlike the FE calculations, the Anton trajectories were performed without any truncation of the SRP system; the simulations included the full RNA, M, and NG domains of the SRP. Four trajectories are performed as summarized in Table 3. Trajectory 1, in which the SRP is initialized from the syn conformation without a bound SP, is prepared with coordinates from the 1QZW structure (20). Trajectory 2, in which the SRP is initialized from the syn conformation with a bound DPAP-WT SP, is prepared with coordinates from the 3KL4 structure (17). Trajectory 3, in which the SRP is initialized from the anti conformation without a bound SP, is prepared with coordinates from the 3NDB structure (19) after removal of the model SP. Trajectory 4, in which the SRP is initialized from the anti conformation with a bound model SP composed of 14 leucine and alanine residues, is prepared from the coordinates of the 3NDB structure (19). All structures were solvated with explicit water molecules in an orthorhombic simulation cell. Na$^+$ and Cl$^-$ counterions are added to achieve electroneutrality at a salt concentration of 100 mM for a total size of 168,500 atoms for the SRP in the syn conformation and 164,294 atoms for the SRP in the anti conformation. Differences in system size are due to different box sizes and different numbers of solvent molecules. Interactions are described by the CHARMM27 force field (72) with TIP3P water.

For each microsecond timescale trajectory, the initial coordinates of the system are equilibrated using the NAMD simulation package (73). Equilibration of the initial configuration consists of conjugate gradient minimization to reduce steric clashes, thermalization of the system to 300 K, and 10 ns of simulation in the NPT ensemble. The system is evolved using Langevin dynamics with a damping constant of 1 ps$^{-1}$, which also provides temperature control at 300 K. Pressure is maintained at 1 bar using the Nosé-Hoover Langevin piston (74). Long range electrostatics are treated using particle mesh Ewald with a real space cutoff of 12 Å. LJ interactions are switched off between 10 and 12 Å using a cubic spline (73). A 2-fs time step was used throughout, and all bond lengths are constrained using the SHAKE algorithm (75).

Following equilibration, production trajectories were performed on the Anton system. Harmonic restraints are introduced to the phosphate backbone of the SRP RNA with a force constant of 418.4 kJ mol$^{-1}$ Å$^{-2}$ to prevent conformational rearrangements of the SRP RNA domain and to avoid overall rotations of the complex with respect to the simulation cell that might cause interactions with its periodic images. Molecular dynamics are evolved with the RESPA numerical integration scheme with a 2-fs time step (76). The Berendsen coupling scheme maintains the temperature at 300 K and pressure at 1
bar (77). Long range electrostatics are treated using the k-space Gaussian Split Ewald method (78) with a real space cutoff of 9.45 Å. LJ interactions were cut off at 9.48 Å. All bond lengths are constrained using the M-SHAKE algorithm (79).

RESULTS AND DISCUSSION

Coupling between SP Binding and SRP Conformation—FEP calculations were used to investigate the potential role of SP binding in driving conformational changes in the SRP. For each SP, we consider \( \Delta G^{(i)} \) as the relative FE of SP binding to the SRP in the syn versus the anti conformation. As schematically illustrated using the thermodynamic cycle in Fig. 4, we calculate \( \Delta G^{(i)} \) for a given SP \( i \) using Equation 5,

\[
\Delta G^{(i)} = \Delta G_A^{(i)} - \Delta G_S^{(i)},
\]

where \( \Delta G_A^{(i)} \) and \( \Delta G_S^{(i)} \) are the respective binding FE of the SP to the syn and anti conformations of the SRP. This construction of the thermodynamic cycle avoids the direct calculation of the FE difference between the SRP in the syn and anti conformation, illustrated by the horizontal arrows in Fig. 4.

As explained under “Computational Methods,” harmonic restraints are employed in the FE calculations to limit the ensemble of sampled configurations for the SRP to those associated with either the syn or anti conformations that are observed in the experimental crystal structures (19, 20). We note that a more common implementation of the thermodynamic cycle approach is to first assume that the different conformations of the biomolecule correspond to basins of stability that are separated by a FE barrier and then to fully sample the configuration space associated with those basins of stability. However, well defined FE differences can be computed between any regions of configuration space, and in this study, we focus on calculating the FE differences between configurations in the vicinity of the experimental crystal structure for SRP in the syn and configurations in the vicinity of the experimental crystal structure for SRP in the anti conformation.

The primary quantity of interest in this analysis is \( \Delta \Delta G^{(i)} \), which reports on the degree to which binding of the SP impacts the conformation of the SRP linker region. The results for the three considered SPs are shown in Table 2. For DPAP-WT, a known targeting sequence, we calculate a value of \(-11\) kJ/mol, which indicates substantial coupling of SP binding to the SRP linker region. Furthermore, the negative sign of \( \Delta \Delta G^{(i)} \) for the case of DPAP-K464L indicates that SP binding leads to a significant shift in the SRP conformational distribution toward the anti conformation, which is consistent with the model hypothesis (Fig. 1C).

The computed value of \( \Delta \Delta G^{(i)} \) for DPAP-K464L suggests that a single lysine-to-leucine mutation from the DPAP-WT sequence leads to a modest increase in the induced conformational bias in the SRP distribution. The calculated \( \Delta \Delta G^{(i)} \) is \(-16\) kJ/mol, which is greater in magnitude than the conformational preference of the DPAP-WT SP, although this difference is within a standard deviation of the statistical error. The DPAP-K464L sequence was chosen because the lysine-to-leucine mutation at residue 464 removes a charged residue and extends the hydrophobic core of the SP. Previous experimental studies (53, 80) have shown that increased hydrophobicity leads to more efficient targeting, and our results are thus consistent with the interpretation that this increased hydrophobicity also leads to a greater conformational preference for the anti conformation of the SRP.

Finally, we consider the PhoA SP, which is known to successfully bind to the SRP but which provides inefficient targeting along the co-translational pathway (10, 30, 54). We find (Table 2) that binding of the PhoA SP leads to a reversed shift in the SRP conformational distribution in comparison with DPAP-WT and DPAP-K464L. The calculated \( \Delta \Delta G^{(i)} \) is 13 kJ/mol, indicating that binding of PhoA SP stabilizes the syn conformation of the SRP relative to the anti conformation. This finding is again in agreement with the model hypothesis in which binding of PhoA fails to induce the SRP conformational change that would lead to efficient targeting.

The calculated \( \Delta \Delta G^{(i)} \) values demonstrate that SP binding to the SRP is coupled to the linker region, and its effects on the SRP conformational distribution are dependent on the sequence of the SP. For all three considered SPs, the degree to which we find that SP binding induces conformational changes in the SRP is fully consistent with the hypothesis that the SRP targeting path-

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**Table 2**

| \( i \)     | \( \Delta G_{\text{SRP}}^{(i)} \) | \( \Delta G_{\text{SRP}}^{(i)} \) | \( \Delta G_{\text{SRP}}^{(i)} \) | \( \Delta G_{\text{SRP}}^{(i)} \) | \( \Delta G_{\text{SRP}}^{(i)} \) | \( \Delta G_{\text{SRP}}^{(i)} \) |
|--------------|----------------------------------|----------------------------------|----------------------------------|----------------------------------|----------------------------------|----------------------------------|
| DPAP-WT      | -310 (2)                         | -389 (2)                         | -400 (3)                         | -79 (3)                          | -90 (3)                          | -11 (4)                          |
| DPAP-K464L   | -275 (2)                         | -340 (2)                         | -355 (2)                         | -65 (3)                          | -80 (3)                          | -16 (7)                          |
| PhoA         | -287 (2)                         | -326 (2)                         | -314 (2)                         | -39 (3)                          | -27 (3)                          | 13 (4)                           |

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**Figure 4.** Thermodynamic cycle that enables the quantification of coupling between SP binding and SRP linker conformation. Four states are defined by linker conformation (blue) and the presence or absence of SP (yellow). \( \Delta \Delta G^{(i)} \) is calculated by taking the difference of \( \Delta G_A^{(i)} \) and \( \Delta G_S^{(i)} \) in the dotted boxes. This is equivalent to the difference of \( \Delta G_{\text{syn} \rightarrow \text{anti}} \) and \( \Delta G_{\text{syn} \rightarrow \text{anti}} \) shown along the horizontal arrows.
Allosteric Response to Signal Peptide Binding

![Diagram](image)

**Figure 5**. A, ensemble-averaged structures from SP-bound sampling trajectories with the linker region shown in blue, the fingerloop shown in green, and the SP shown in yellow. B, fingerloop displacement $R$ and the relative orientation of the linker and $\alpha$M1 helix $\phi$ are shown for each SP and SRP pair, revealing the structural differences associated with binding efficient and inefficient SPs.

way is triggered by SP binding that induces conformational changes in the SRP from the syn to the anti conformation. Furthermore, the results explain the counter-intuitive experimental observation that SPs with favorable binding to the SRP need not lead to efficient downstream protein targeting (41); indeed, it is seen here that favorable PhoA binding induces a shift toward the syn conformation of SRP, which hinders the kinetics of complex formation between SRP and its receptor. We additionally note that this connection between SRP conformational shifts upon substrate binding provides a mechanism for SP identity to influence the kinetics of processes at large distances from the SP-binding site.

FE calculations not only provide a quantitative measure of the extent of allosteric coupling between SP binding and SRP linker conformation, but the sampling trajectories also provide details of the structural differences in the SRP upon binding of the different SPs. For each SP considered, Fig. 5A shows the ensemble-averaged configuration of the SRP in the syn and anti conformations. In each case, the fingerloop of the SRP reorganizes to accommodate binding of the SP. The position of the fingerloop, measured in terms of its center-of-mass distance $R$ from the fifth helix of the M domain (αM5) helix at the opposite side of the SP binding pocket, is found to be displaced by 4–9 Å upon SP binding (Fig. 5A). This is in agreement with previous structural and biochemical work (28, 40), which suggests that SP binding displaces the fingerloop and that fingerloop flexibility is crucial for enabling the SRP to bind to SPs of differing sequences.

The three SPs considered here differ substantially in terms of their size, with DPAP-WT and DPAP-K464L excluding more volume than PhoA (Table 1). These differences are reflected in the conformation of the SRP bound to the SP, in terms of both the degree of fingerloop displacement and the relative orientation between the linker region and M domain. These conformational differences are shown in Fig. 5B, where the relative orientation is monitored by the dihedral angle $\phi$ between the helical axis of the linker and the axis of the first helix of the M domain (αM1), as well as by the fingerloop displacement, $R$. For each SP, the observed dihedral angle associated with binding in the syn conformation is small. For the cases of DPAP-WT and DPAP-K464L bound to the SRP in the anti conformation, the value of $\phi$ is much larger than in the syn conformation due to the loss of contact between the linker and the M domain; the smaller PhoA SP likewise exhibits an increased value of $\phi$ in the anti conformation but to a lesser extent. A more striking trend is observed in the fingerloop displacement, $R$. For both of the efficiently targeted sequences, DPAP-WT and DPAP-K464L, the fingerloop is substantially displaced in both the syn and anti conformations, whereas the inefficiently targeted PhoA SP exhibits a reduced displacement of the fingerloop in the anti conformation. The differing degree to which the efficient and inefficient SPs impact these structural features of the SRP suggests a mechanistic basis for the observed correlations between SP binding and SRP conformational changes (Table 2). Although the single amino acid mutation that distinguishes the DPAP-WT and DPAP-K464L, the fingerloop is substantially displaced in both the syn and anti conformations, whereas the inefficiently targeted PhoA SP exhibits a reduced displacement of the fingerloop in the anti conformation. The differing degree to which the efficient and inefficient SPs impact these structural features of the SRP suggests a mechanistic basis for the observed correlations between SP binding and SRP conformational changes (Table 2), and although the binding of these two SPs leads to similar structural changes in the SRP fingerloop displacement and helix orientation (Fig. 5B), we do note that the binding of these two SPs leads to differences in other structural quantities. In DPAP-WT, the positive charge interacts with the negatively charged residues of the fingerloop (Asp-364 and Glu-365) in the syn conformation, but this interaction is not observed in the anti conformation; the DPAP-K464L mutant lacks the positive charge for this interaction with the syn conformation. These different interactions may explain the somewhat stronger bias of the DPAP-K464L mutant for the anti conformation of the SRP upon binding. Regardless, these results illustrate that specific amino acid interactions can play a role in determining the conformational shift of the SRP upon SP binding.

**Large Scale Conformational Changes from Microsecond Trajectories**—Microsecond trajectories are performed with the Anton supercomputer (70) to investigate the conformational dynamics of the SRP (Table 3). Fig. 6 plots the fingerloop displacement, $R$, and the dihedral angle between the helical axis of the linker and $\alpha$M1 helix, $\phi$, as a function of simulation time for all four trajectories; these geometric quantities were previously introduced in the discussion of the FE sampling trajectory.
Allosteric Response to Signal Peptide Binding

### TABLE 3
Summary of microsecond trajectories

| Trajectory | Initial SRP configuration | SP     | Time  |
|------------|---------------------------|--------|-------|
| T1         | syn                       | No SP  | 3.4   |
| T2         | syn                       | With SP| 4.5   |
| T3         | anti                      | No SP  | 4.3   |
| T4         | anti                      | With SP| 3.4   |

Fig. 6 A, fingerloop displacement $R$ as a function of time in the microsecond timescale MD trajectories. $B$, relative orientation of the linker and $\alpha$M1 helix $\phi$ as a function of time in the microsecond timescale MD trajectories.

Allosteric responses of Figs. 5 and 6 reveal similar structural responses of the SRP to binding of the various SPs, confirming that $R$ and $\phi$ are relatively local geometric features that are unchanged by inclusion of the NG domain. For trajectories T1 and T3, neither of which include the bound SP, it is seen in Fig. 6A that the fingerloop occludes the binding pocket, exhibiting a small value for the displacement distance. For trajectories T2 and T4, both of which include the bound SP, it is seen that the fingerloop is significantly displaced (4–9 Å) to accommodate the SP; these results are consistent with the observations in Fig. 5 that SP binding displaces the fingerloop from the binding pocket. Similarly, for trajectories T1 and T2, both of which are initialized in the syn conformation, it is seen in Fig. 6B that the SRP undergoes relaxation to adopt configurations with values of $\phi$ that are consistent with those observed in the equilibrium FE sampling trajectories for SRP in the syn conformation (Fig. 5). For trajectories T3 and T4, both of which are initialized in the anti conformation, it is seen in Fig. 6B that the SRP adopts configurations with values of $\phi$ that are consistent with the loss of contact between the linker region and the M domain, which is again consistent with the structural features that are observed for the equilibrium FE sampling trajectories for SRP in the anti conformation (Fig. 5).

To further characterize the conformational changes observed in the microsecond timescale MD trajectories, Fig. 7 shows snapshots of the system at various times along the trajectories. In addition to illustrating the changes in $R$ and $\phi$ that were discussed previously, Fig. 7 also illustrates the process by which the fingerloop collapses to occupy the SRP binding pocket in the absence of the bound SP (trajectory T3).

We now analyze the microsecond timescale MD trajectories in the context of a recent study by Shen et al. (16), which used single molecule FRET experiments to monitor the SRP conformational dynamics. This experimental study labeled the SRP RNA distal end and NG domain and found that the SRP samples both a low efficiency and a high efficiency FRET state, with differences in FRET efficiency that correspond to distance changes of ~10 nm. Fig. 8 presents results from the four microsecond timescale MD simulations of the SRP, plotting the distance between the G domain and RNA corresponding to the donor-acceptor distance measured in the single molecule FRET experiments. Trajectories T1 and T2, which are initialized from the syn conformation of the SRP, show 6–8 nm changes in donor-acceptor distance during the course of the simulation. Trajectory T2, which has a bound SP, has greater fluctuations in this coordinate than T1, which does not have a bound SP. Trajectories T3 and T4, which are initialized from the anti conformation of the SRP, exhibit little deviation from the initial distance. These results support the conclusion that nanometer-length scale changes in SRP conformation are intrinsic features of the SRP, although we do not suggest that the conformational states observed in these trajectories correspond directly to the low-FRET and high-FRET states reported by Shen et al. (16).

Interestingly, the trajectories reveal that nanometer-length scale changes in distance occur via the rigid-body movement of M and NG domains connected by the flexible linker region, although the conformation of each individual domain is unchanged. In Fig. 9A, for each residue pair $i$ and $j$ in the SRP, we plot $\delta_{ij}(t) = |d_{ij}(0) - d_{ij}(t)|$, the difference in the distance $d_{ij}$ between the two C$\alpha$ atoms of each residue at the initial configuration of the trajectory and at various subsequent times $t$ along trajectory T2. This metric reveals only small changes in distance between atom pairs that both belong to the M domain or that both belong to the NG domain, which correspond to the indicated diagonal blocks in Fig. 9A. Large distance changes are only observed in atom pairs for which one atom is in the M domain and the other is in the NG domain, which correspond to the off-diagonal blocks. This suggests that the observed changes in conformation and donor-acceptor distance are the result of changes in the relative orientation of the M and NG domains, whereas the M and NG domains individually evolve as rigid bodies. Fig. 9B further illustrates this point by plotting the average of $\delta_{ij}$ over NG-NG pairs, M-M pairs, and M-NG pairs along the trajectory. It is clear that only the M and NG domains undergo large changes in relative orientation in trajectory T2.
The same conclusion holds for the other microsecond trajectories that were performed in this study.

The microsecond timescale dynamics revealed in these simulations suggest a mechanism for coordinating the spatial and temporal organization between independent functional domains. In SRP, SP binding occurs in the M domain, and downstream targeting events, such as dimerization of the SRP with its receptor and GTPase activity of the SRP, involve the NG domain of the SRP (81). From the earlier FE simulations (Fig. 4 and Table 2), it is evident that SP binding leads to sequence-specific changes in the conformational distribution in the flexible linker domain. Fig. 9 further reveals that such changes in the linker domain...
manifest as large lengthscale changes in the relative orientation of the M and NG domains of the SRP, thus coordinating SP binding at the M domain with downstream steps involving the NG domain.

Conclusions—The fidelity of co-translational protein targeting and recognition is a remarkable feature of cellular signaling pathways that demands understanding from the perspective of molecular interactions. The central component of co-translational targeting, the SRP, is capable of binding a diverse set of SPs while differentiating between sequences with similar biochemical features. In this work, we have demonstrated allosteric coupling between SP binding and conformational changes of the linker region of the SRP. This coupling is sensitive to the sequence identity of the SP, thus providing a mechanism for conferring SP sequence specificity in the signaling pathway. Furthermore, microsecond timescale simulations reveal that the SRP undergoes large scale conformational changes that are characterized by rigid body motion of individual domains connected by a flexible linker, which enables small conformational changes of the linker upon SP binding to be conferred onto larger ~10-nm lengthscales.

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