Sequential Checkpoints Govern Substrate Selection During Cotranslational Protein Targeting

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Abstract

Proper protein localization is essential for all cells. However, the precise mechanism by which high fidelity is achieved is not well understood for any targeting pathway. To address this fundamental question we investigated the signal recognition particle (SRP) pathway in E. coli, which delivers proteins to the bacterial inner membrane through recognition of signal sequences on cargo proteins. Fidelity was thought to arise from the inability of SRP to bind strongly to incorrect cargos. Using biophysical assays, we found that incorrect cargos were also rejected through a series of checkpoints during subsequent steps of targeting. Thus high fidelity is achieved through the cumulative effect of multiple checkpoints; this principle may be generally applicable to other pathways involving selective signal recognition.
comparable to that of phoA-3A7L, it is not an SRP substrate due to an N-terminal extension (Fig. 1B) (14). Firefly luciferase, a cytosolic protein without signal sequences, was used as a negative control (Fig. 1B) (12). For all the experiments, homogeneous stalled RNCs were purified and used as cargos (8, 15).

We first tested the binding affinities of SRP for different cargos (Fig. 1A, step 1). RNC binding to SRP was detected as an increase in the fluorescence anisotropy of fluorescein-labeled SRP (C421). Cargos with the most hydrophobic signal sequences bound to SRP tightly (RNC1A9L and RNC2A8L), with equilibrium dissociation constants ($K_d$) of ~1 nM or less (Figs. 1C and S1). The next strongest cargo, RNC3A7L, also exhibited strong albeit attenuated binding to SRP, with $K_d$ ~ 10 nM (Fig. 1C). Nevertheless, the affinity of incorrect cargos or the empty ribosome for SRP was still substantial, with $K_d$'s of 80 – 100 nM (Fig. 1, D–E, and fig. S1; see also ref 16). At the cellular SRP concentration of ~400 nM (Fig. 1E) (17), a substantial amount of incorrect cargos could bind to SRP (18). Surprisingly, although EspP is not an SRP substrate, RNCEspP bound SRP as tightly as RNC3A7L (Fig. 1C). Thus the differences in cargo binding affinity do not provide sufficient discrimination against incorrect cargos, and additional factors in the bacterial cytosol did not increase the specificity of SRP-cargo binding (Fig. S2 and SOM text) (19). We therefore proposed that subsequent steps in the targeting pathway, including formation of the SRP-SR complex and GTP hydrolysis, provide additional checkpoints to reject the incorrect cargos (Fig. 1A, red arrows b–d and SOM text) (20).

We first tested whether the early SRP-SR complex is stabilized more strongly by the correct than the incorrect cargo (Fig. 1A, arrow b). We assembled cargo-SRP-SR early complexes in the absence of nucleotides; this blocks the rearrangement of the GTPase complex to subsequent conformations (7, 8). The equilibrium stabilities of the early complexes were measured using fluorescence resonance energy transfer (FRET) between donor- and acceptor-labeled SRP and SR (7). The early complex was significantly stabilized by RNC1A9L and RNC2A8L, with $K_d$ ~ 80 nM (Fig. 2A), and this stability was weakened up to 50-fold for the weaker cargos (Fig. 2, B–C, and fig. S3). With incorrect cargos such as RNCEspP and RNCluciferase, the FRET efficiency also plateaued at lower values, ~0.3 – 0.4 (Fig. 2, B and D, and fig. S3), compared to ~0.66 with the correct cargos (Fig. 2, A and D). This suggests that the SRP and SR are likely positioned differently in the early targeting complexes formed by the incorrect cargos.

A mispositioned early complex would lead to a slower rearrangement to form the closed complex (Fig. 1A, step 3). To test this hypothesis, we preformed the early targeting complex and directly measured its rearrangement using acrylodan-labeled SRP (C235), which specifically monitors the closed complex. With RNC1A9L, this rearrangement occurred at 0.3 s$^{-1}$ (Fig. 2E). RNC3A7L and RNCphoA mediated this rearrangement 40% slower (Fig. 2G and S4). Notably, RNCEspP and cargos weaker than RNC5A5L mediated this rearrangement 5–10-fold slower than RNC1A9L (Fig. 2, F–G, and fig. S4). Thus incorrect cargos do not induce the formation of a stable and productive early complex, and are more likely to exit the pathway (Fig. 1A, arrow b).

The more favorable pre-equilibrium to form the early intermediate combined with the faster early-to-closed rearrangement would allow the correct cargos to mediate faster GTP-dependent assembly of a stable closed complex (Fig. 1A, steps 2–3). We characterized this cumulative effect using both FRET (Fig. 3, A–C, and fig. S5, F–G) and acrylodan-labeled SRP (C235) (Fig. S5). Both probes showed that the correct cargos mediated rapid assembly of the closed complex (Fig. 3A and S5A), and this rate decreased significantly with weaker signal sequences (Fig. 3, B–C, and fig. S5). Overall, there is a $\sim10^3$-fold kinetic advantage for correct cargos over incorrect ones.
discrimination between the strongest and weakest cargos in stable SRP-SR complex assembly, which delivers the cargo to the membrane (Fig. 3C and S5E).

If GTP hydrolyzed too quickly in the SRP-SR complex, this would abort the targeting reaction before the cargo is productively unloaded (8, 21). To test whether the correct cargos prevent premature GTP hydrolysis better than the incorrect cargos (Fig. 1A, step 4), we determined the GTPase rates from the cargo-SRP-SR complex. RNC1A9L and RNC2A8L reproducibly delayed GTP hydrolysis 6–8 fold (Fig. 3D and S6). RNC3A7L had a 3–4 fold inhibitory effect on the GTPase reaction (Fig. S6). In contrast, incorrect cargos such as RNC_EspP did not substantially affect the GTPase rate (Fig. 3, E–F, and fig. S6). Thus the fidelity of protein targeting can be further improved through kinetic proofreading mechanisms by using the energy of GTP hydrolysis (Fig. 1A, arrow d).

Our data suggested a model in which the incorrect cargos are rejected not only through binding affinity, but also through differences in the kinetics of SRP-SR complex assembly and GTP hydrolysis (Fig. 1A and 4A, top). Based on this model, we calculated the amount of substrates retained in the SRP pathway after each checkpoint (SOM Methods). The cargo binding step was not sufficient to discriminate against incorrect cargos, allowing over 75% of them to enter the SRP pathway (Fig. 4A, light grey). During cargo delivery through SRP-SR complex assembly, a large fraction of substrates weaker than phoA were rejected (Fig. 4A, dark grey). Finally, kinetic competition between GTP hydrolysis and cargo unloading further improved the discrimination between correct and incorrect substrates (Fig. 4A, black). To validate the model, we determined the targeting efficiency of proteins with various signal sequences using a well-established assay that tests the ability of E. coli SRP and SR to mediate the co-translational targeting of preproteins to microsomal membranes (SOM Methods) (22, 23). Substrates with signal sequences stronger than 3A7L were efficiently targeted and translocated (Fig. 4B and S7). In contrast, EspP and substrates with signal sequences weaker than phoA showed severe defects in translocation, and almost no translocation was detected for phoA-8A2L (Fig. 4B). The experimentally determined protein targeting efficiencies agreed well with predictions based on the kinetic and thermodynamic measurements (Fig. 4C), suggesting that our model (Fig. 1A and 4A) faithfully represents how SRP handles its substrates.

Thus fidelity during cotranslational protein targeting is achieved through the cumulative effect of multiple checkpoints, by using a combination of binding, induced fit, and kinetic proofreading mechanisms (SOM text). Although the incorrect cargos are not completely rejected during the initial binding step, they are discriminated repeatedly during subsequent steps possibly because they bind the SRP in a less productive mode (6). In addition, the translocation machinery provides another important checkpoint (24). Similar strategies of using multiple checkpoints to ensure fidelity have been demonstrated in tRNA synthetases (25), protein synthesis (26), and DNA and RNA polymerases (27, 28), and possibly represent a general principle for complex cellular pathways that need to recognize degenerate signals or to discriminate between correct and incorrect substrates based on minor differences.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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References and Notes

Figure 1. Potential fidelity checkpoints in the SRP pathway

(A) Model for potential checkpoints during co-translational protein targeting. A cargo (RNC) with a signal sequence (magenta) enters the pathway upon binding SRP, and is either retained (black arrows) or rejected (red arrows) at each step (numbered 1–5). T and D denote GTP and GDP, respectively. (B) Signal sequence variants used in this study. Bold highlights the hydrophobic core. Blue highlights the N-terminal signal sequence extension of EspP. (C, D) Equilibrium titrations of SRP-RNC binding. Nonlinear fits of data gave $K_d$ values of 0.55±0.20, 8.4±2.0, 13.6±3.0, 108±11 and 130±12 nM for RNC$_{1A9L}$ (C, ●), RNC$_{3A7L}$ (C, ■), RNC$_{EspP}$ (C, ◆), RNC$_{phoA}$ (D, ■) and RNC$_{luciferase}$ (D, ●), respectively. Error bars are SDs from three independent experiments. (E) Summary of the binding
affinities of SRP for different cargos. The dashed line denotes the cellular SRP concentration of 400±58 nM. Error bars are SEs of the fits.
Figure 2. Correct cargos stabilize the early intermediate and mediate faster rearrangement to the closed complex

(A, B) Equilibrium titrations of the early intermediate. Nonlinear fits of data gave $K_d$ values of 78±5, 110±8, 311±21 and 2060±201 nM and FRET endpoints of 0.68±0.02, 0.64±0.02, 0.41±0.03, and 0.34±0.02 for RNC1A9L (A, ●), RNC2A8L (A, ■), RNCEspP (B, ■), and RNCLuciferase (B, ●), respectively. Error bars are SDs from three independent experiments.

(C, D) Summary of the $K_d$ values (C) and FRET end points (D) of the early intermediates formed by different cargos. Error bars are SEs of the fits in C and SDs from three independent experiments in D.

(E, F) Measurements of the early to closed rearrangement. Nonlinear fits of data gave rate constants of 0.31±0.02 s$^{-1}$ with RNC1A9L (E) and 0.31±0.02 s$^{-1}$ with RNC2A8L (F).

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0.039±0.003 s$^{-1}$ with RNC$_{\text{luciferase}}$ (F). Error bars are SDs from three independent experiments. (G) Summary of the rate constants for the early to closed rearrangement with different cargos. Error bars are SEs of the fits.
Figure 3. Correct cargos accelerate GTP-dependent complex formation but delay GTP hydrolysis

(A, B) Rate constants of SRP-SR complex assembly in GMPPNP measured by FRET. Linear fits of data gave $k_{on}$ values of 9.9±1.3×10^6, 8.8±1.6×10^6, 2.0±0.2×10^7, 6.3±0.4×10^6, 1.1±0.2×10^6 and 1.8±0.3×10^3 M^{-1}s^{-1} for RNC_{1A9L} (A, ●), RNC_{2A8L} (A, ■), RNC_{3A7L} (B, ◆), RNC_{phoA} (B, ■), RNC_{5A5L} (B, ◆) and RNC_{luciferase} (B, ▲), respectively. Error bars are SDs from three independent experiments. (C) Summary of GTP-dependent complex assembly rate constants with different cargos. Error bars are SEs of the fits. (D, E) Effects of cargo on GTP hydrolysis from the SRP•SR complex. Nonlinear fits of the data gave maximal GTPase rate constants ($k_{cat}$) of 0.72±0.03 s^{-1} without cargo (●), and 0.11±0.01.
0.38±0.02, 0.51±0.08, and 0.65±0.22 s⁻¹ with RNC₁₉₉₁ (D, ■), RNC₅₆₅₁ (D, ◆), RNCₑₛₚ (E, ■) and RNC_#{luciferase} (E, ◆), respectively. Error bars are SDs from three independent experiments. (F) Summary of GTPase rate constants in the presence of different cargos. Error bars are SEs of the fits.
Figure 4. Stepwise rejection of incorrect cargos from the SRP pathway
(A) Top panel, cargos are either retained (black arrow) or rejected (red arrow) during each checkpoint. Lower panel, predicted fraction of cargos retained in the SRP pathway during each checkpoint (Supplementary text). (B) SRP-dependent protein targeting and translocation of the model substrates. pPL and PL denote the precursor and processed forms of the substrate, respectively. (C) Predicted protein targeting efficiencies (● and ○) agree well with the experimentally determined values (■), quantified from the data in (B). Translation elongation rates of 20 (●) and 10 amino acids/s (○) were used for the E. coli and eukaryotic ribosomes, respectively, to calculate the targeting efficiencies.