Competitive Binding of the SecA ATPase and Ribosomes to the SecYEG Translocon

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Background: Both SecA and the ribosome need to interact with the translocon during membrane protein insertion.

Results: SecA competes with ribosomes and ribosome-nascent chain complexes for binding to the translocon.

Conclusion: SecA and ribosome binding to the translocon is mutually exclusive, implying that during membrane protein insertion, both ligands bind the translocon in a sequential manner.

Significance: Insight in the mechanism of membrane protein insertion.

During co-translational membrane insertion of membrane proteins with large periplasmic domains, the bacterial SecYEG complex needs to interact both with the ribosome and the SecA ATPase. Although the binding sites for SecA and the ribosome overlap, it has been suggested that these ligands can interact simultaneously with SecYEG. We used surface plasmon resonance and fluorescence correlation spectroscopy to examine the interaction of SecA and ribosomes with the SecYEG complex present in membrane vesicles and the purified SecYEG complex present in a detergent-solubilized state or reconstituted into nanodiscs. Ribosome binding to the SecYEG complex is strongly stimulated when the ribosomes are charged with nascent chains of the monotopic membrane protein FtsQ. This binding is competed by an excess of SecA, indicating that binding of SecA and ribosomes to SecYEG is mutually exclusive.

The initiation of protein translocation or membrane protein insertion is dependent on the high affinity interactions between the SecYEG complex and SecA or the ribosome. In recent years, structural, biochemical, and computational approaches have provided detailed insights in the interaction between SecYEG and its cytosolic binding partners. Both SecA and the ribosome interact primarily with the largest subunit of the SecYEG complex, i.e. SecY. This interaction involves multiple contact points, and the main connections are formed by the ribosomal 23S rRNA, the ribosomal protein L23, and the fourth and fifth cytoplasmic loops (C4 and C5) of SecY (5–9). Substitution of the conserved arginine residues in these SecY loops severely reduces the interaction with the ribosome (7). In addition, the C-terminus of SecY interacts with the ribosomal protein L24, and the N terminus and amphipathic helix of SecE interact with proteins L23 and L29 (7–9). Intriguingly, SecA appears to bind to the same SecY loops that are important for ribosome interaction. Indeed, a recent crystal structure of SecYEG in complex with SecA (10) demonstrated contacts between the SecY C4 and C5 loops and SecA (11, 12). Mutational analysis identified the arginine at position 357 in the C5 loop as indispensable for SecA binding (11) but rather prevents the SecA-dependent initiation of protein translocation (14).

Although many substrates of the translocon only require the action of either SecA or the ribosome for translocation or membrane insertion, the biogenesis of membrane proteins with large periplasmic loops or domains, such as FtsQ and CyoA, requires both functions (15, 16). Insight in the timing and coordination of these activities is essential for the understanding of the mechanism of membrane protein insertion. It has been suggested that non-translating ribosomes and SecA do not compete for SecYEG binding and thus would be able to bind simultaneously (17). However, the overlapping binding sites (7, 8, 10) and the anticipated steric constraints upon association of these two large ligands make this difficult to envision. Here, we have monitored the binding of SecA, non-translating ribosomes and RNC complexes of the monotopic membrane protein FtsQ to the membrane-embedded and detergent-solubilized SecYEG complex using...
two different methods, i.e. surface plasmon resonance and fluorescence correlation spectroscopy. Our data demonstrate that SecA and the ribosome compete for binding to the SecYEG complex and that substrate loading is an important determining factor in this process.

**EXPERIMENTAL PROCEDURES**

**Strains and Plasmids**—Inner membrane vesicles (IMVs) with endogenous or overexpression levels of SecYEG were isolated from *Escherichia coli* SF100 transformed with the plasmids indicated in Table 1 as described previously (18). The R357E mutation and R255E,R256E double mutation were indicated in Table 1 as described previously (18). The mutations were introduced into Cys-less SecY by site-directed mutagenesis using a unique cysteine at position 148 of SecY was introduced into Cys-less SecY by site-directed mutagenesis using pEK20 (19) as template, yielding pNN260 (11) and pZW1, respectively. A unique cysteine at position 148 of SecY was introduced into pZW1 and pNN260 yielding pZW2 and pZW3, respectively.

Plasmid pUC19Strep3FtsQSecM (20) was used for the isolation of FtsQ108 RNCs. For plasmids encoding FtsQ RNCs of 77 or 87 residues and FtsQ with a deletion of the first transmembrane segment (TMS), the PstI-EcoRV fragment of pUC19Strep3FtsQSecM (20) was exchanged with fragments coding for FtsQ-(3–41), FtsQ-(3–51), and FtsQ-(49–122) yielding pEK765, pEK764 (21), and pEK767, respectively. All plasmids were verified by sequence analysis.

**Ribosome and RNC Complex Purification**—Non-translating ribosomes were purified from *E. coli* MRE600 (22). Cells were cultured in LB medium at 37 °C to an *A*₆₀₀ of 0.6 and harvested by centrifugation (9500 × *g*, 4 °C, 20 min). The pellet was resuspended in buffer A (20 mM Tris-HCl, pH 7.5, 100 mM NaCl, 10.5 mM MgOAc, 0.5 mM EDTA, 1 mM Tris(2-carboxyethyl)phosphine) and French-pressed twice at 700 p.s.i. Cell debris was removed by centrifugation (30,000 × *g*, 4 °C, 30 min), and the cleared lysate was laid on a 1 M sucrose cushion prepared in buffer B. After centrifugation (112,000 × *g*, 14 °C, 17 h), the ribosomal pellet was dissolved in buffer R and loaded on a StrepTactin column (IBA). The column was washed with two column volumes of buffer R containing 0.5 mM KCl and five column volumes of buffer R. RNCs were eluted with two column volumes of buffer R containing 2.5 mM desfihiotin. The eluate was concentrated using Millipore Amicon Ultra-4 or Ultra-15 centrifugal tubes (cut-off 50 kDa), and the ribosome concentration was determined. The presence of stalled nascent chains was confirmed by SDS-PAGE followed by Western blotting using an antibody against the STREP-tag (IBA).

**Surface Plasmon Resonance (SPR)**—SPR measurements were performed on a Biacore 2000 system (GE Healthcare) as described (26). In short, IMVs containing endogenous or overexpressed levels of SecYEG (mutants) were immobilized in the separate channels of a L1 sensor chip (GE Healthcare). Buffer B (50 mM Tris-HCl, pH 7.5, 150 mM KCl, 10 mM MgCl₂, 1 mM DTT, and 0.5 mg/ml BSA) containing SecA, ribosomes, or RNCs was injected into the channels at a flow rate of 20 μl/min, and SecYEG binding was probed at 25 °C. The binding surface of IMVs was regenerated by the injection of 100 mM Na₂CO₃, pH 10, following by system equilibration using buffer B. Data were corrected for background binding to the IMVs containing endogenous levels of SecYEG. For SecA-ribosome competition experiments, ribosome binding was measured in the presence or absence of a saturating concentration (96 nM or as indicated) of SecA in the running buffer B. Data were fitted by nonlinear regression analysis of the response levels at equilibrium using SigmaPlot (Systat Software Inc.). For the single site (A + B ↔ AB) interaction model including a non-saturable linear component the following equation was used,

\[
Y = \frac{B_{\text{max}} \times X}{K_D + X} + Ns \times X
\]  

(Eq. 1)

where *Y* is the SPR binding response at analyte (SecA or ribosome) concentration *X*, *B* max is the maximal binding response,
$K_D$ is the dissociation constant, and $N_s$ is the nonspecific binding coefficient.

For the two-site saturation model the following equation was used.

$$ Y = \frac{B_{max1} \times X}{K_{D1} + X} + \frac{B_{max2} \times X}{K_{D2} + X} \quad \text{(Eq. 2)} $$

**Fluorescence Correlation Spectroscopy (FCS)**—For fluorescence microscopy, SecY(L148C)EG was overexpressed, purified, and labeled with AlexaFluor 488-C5-maleimide (Invitrogen) as described (27). Reconstitution of the fluorescently labeled SecY(L148C)EG into MSP1D1 nanodiscs (28) was carried out as described (29, 30) with the following modifications. A synthetic lipid composition consisting of 25 mol % dioleoylphosphatidylglycerol, 5 mol % cardiolipin, 30 mol % dioleoylphosphatidylethanolamine, and 40 mol % dioleoylphosphatidylcholine was used for nanodiscs formation, and lipids were destabilized by 0.5% (v/v) Triton X-100 before SecYEG reconstitution. SecYEG-containing nanodiscs were isolated using a Tricorn Superdex 200 10/300 column (GE Healthcare). The concentration of reconstituted SecYEG-AlexaFluor 488 in the collected fractions was determined based on the specific fluorophore absorbance.

FCS experiments were performed using the inverted confocal microscope LSM 710 equipped with the Confocor 3 module (Carl Zeiss GmbH). A solution of 50 nM AlexaFluor 488 dye with a known diffusion coefficient of $300 \times 10^{-8}\text{ cm}^2\text{s}^{-1}$ (31) was used to adjust the laser intensity at 488 nm and to calibrate the observation volume. Before the experiment SecYEG complexes were diluted to a concentration of $\sim 100\text{ nM}$ in 50 mM Tris-HCl, 100 mM NaCl, 10 mM MgCl$_2$, and 5% (v/v) glycerol. For detergent-solubilized SecYEG the buffer was also supplied with 0.05% (w/v) n-dodecyl β-D-maltoside (DDM). Diffusion-driven fluctuations in the fluorescence intensity of SecYEG-conjugated AlexaFluor 488 were recorded over 10 s, and the measurements were repeated 10 times for each selected area. Recorded traces were averaged. For each experimental condition at least 10 measurements were done at different positions within the sample volume.

Autocorrelation curves were built and analyzed using ZEN 2010 software package (Carl Zeiss GmbH, Germany). Triplet state population of the fluorophore was below 20%, and it was omitted in the following analysis as the autocorrelation traces were fitted within 10 μs to a 10-s time range. The data were analyzed assuming free three-dimensional diffusion of SecYEG, and the auto-correlation curves were fitted according to the equation,

$$ G(t) = \frac{1}{N_i} \left(1 + \frac{t}{\tau_D} \right)^{-1} \left(1 + \frac{1}{\frac{\omega_0}{Z_0} t \frac{1}{\tau_D}} \right) \quad \text{(Eq. 3)} $$

where $G(t)$ is the amplitude of the auto-correlation function, $N_i$ is the average number of fluorescent particles in the laser focus, and $\tau_D$ is the diffusion time through the focus. To analyze SecYEG diffusion in the presence of ribosomes and to estimate fractions of free and ribosome-bound SecYEG, a two-component fitting model was applied,

$$ G(t) = \frac{1}{N_1} \left(1 + \frac{t}{\tau_{D1}} \right)^{-1} \left(1 + \frac{1}{\frac{\omega_0}{Z_0} t \frac{1}{\tau_{D1}}} \right) \times \frac{1}{N_2} \left(1 + \frac{t}{\tau_{D2}} \right)^{-1} \quad \text{(Eq. 4)} $$

where $N_i$ and $\tau_{Di}$ are average number of fluorescent particles and diffusion time of $i$th species, respectively. The diffusion times of SecYEG and SecYEG-ribosome complexes were preset for the fitting procedure. For the free SecYEG the value was directly measured, whereas the diffusion time of the SecYEG-ribosome complex was approximated with that of fluorescently labeled FtsQ108-RNC, which was experimentally determined in our setup ($\sim 900 \mu$s).

**SecA Co-sedimentation Assay**—SecA (92 nM) was incubated with non-translating ribosomes (60 nM), FtsQ108 (50 nM), and FtsQ108ΔTMS (50 nM) in buffer B for 20 min at room temperature. Samples were loaded on a 35% (w/v) sucrose cushion and centrifuged for 2 h at 350,000 $\times$ g, 4°C. Pellets were dissolved in SDS-PAGE sample buffer and analyzed by SDS-PAGE and silver staining.

**In Vitro Protein Translocation Assay**—In vitro translocation of fluorescent-labeled proOmpA(C290S) was done as described (32). Translocated, protease-resistant proOmpA was separated by SDS-PAGE and visualized with a LAS-4000 imager (FujiFilm) using the SyBr Blue Y515 Di filter.

**Miscellaneous**—SDS-PAGE, Western blotting, and silver staining were performed according to standard protocols. IMVs concentrations were determined using the Bio-Rad DC protein assay kit using BSA as standard. ProOmpA(C290S) and SecA were purified as described (33, 34). Quantification of the SDS-PAGE bands was done using Aida/2D densitometry (Raytest).

**RESULTS**

**Detection of SecYEG-Ribosome Interaction by Surface Plasmon Resonance**—We employed SPR to follow the binding of ribosomes to the membrane-embedded SecYEG complex in real time. This method was previously shown to accurately detect the high affinity interaction between SecA and SecYEG (11, 26). IMVs containing overexpressed levels of SecYEG were immobilized on a Biacore L1 chip, and the association and dissociation of SecA (Fig. 1A) and ribosomes (Fig. 1B) was followed in time. To correct for “bulk” contributions to the SPR signal and nonspecific binding to the membrane and chip surface, all measurements were corrected for binding to IMVs bearing endogenous SecYEG levels that were immobilized in a reference channel (26) (supplemental Fig. S1A). Both SecA and ribosome injection resulted in a SecYEG-dependent SPR response (Fig. 1B).

To validate that the observed SPR response reflected the binding of ribosomes to the SecYEG complex, we analyzed the interaction between ribosomes and two SecY mutant complexes, i.e. SecY(R255E,R256E)EG and SecY(R357E)EG. These mutations are located in the SecY cytoplasmic loops C4 and C5, respectively, and have been shown to disturb the SecYEG-ribo-
some interaction in detergent solution (7). Both SecYEG mutants were overexpressed to similar levels as Cys-less SecYEG (supplemental Fig. S1A). The alteration in the charge distribution in these cytosolic loops, however, resulted in a reduced mobility of SecYEG on SDS-PAGE (supplemental Fig. S1A). The SecY(R357E) mutation, which has been reported to affect the initiation of SecA-dependent protein translocation (13, 14), inhibited proOmpA translocation completely, whereas the SecY(R255E,R256E)EG mutant was normally active (supplemental Fig. S1B). Importantly, both mutants allowed for significant SecA binding (Fig. 1A), indicating that the mutations did not result in a major disturbance of the loop conformation. However, the SPR responses upon injection of ribosomes over immobilized SecY(R255E,R256E)EG or SecY(R357E)EG IMVs were dramatically reduced as compared with Cys-less SecYEG IMVs (Fig. 1B). These observations are consistent with the notion that the mutated loop residues are important for the SecYEG-ribosome interaction (7) and demonstrate that the SPR method genuinely monitors the interaction between ribosomes and the membrane-embedded SecYEG complex.

Previously, we observed that the SecA association and dissociation phases do not fit to a simple bimolecular interaction model (26, 35). Similarly, an accurate determination of the association and dissociation rates of ribosome binding could not be obtained by such simple data fitting models. Therefore, an apparent affinity of the interaction was determined by nonlinear regression analysis of the response levels at equilibrium (Fig. 2). For the SecYEG-SecA interaction, this resulted in an apparent dissociation constant ($K_D$) of 4.5 nM when fitted according to a simple single site ($A$/$H$) interaction model including a non-saturable linear component (Fig. 2A, Table 2).

### Table 2

<table>
<thead>
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<th>Analyte</th>
<th>$B_{max1}$</th>
<th>$B_{max2}$</th>
<th>$K_D$</th>
<th>$K_S$</th>
<th>$N_s$</th>
<th>$R^2$</th>
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<td>4.5</td>
<td>0.15</td>
<td>ns</td>
<td>ns</td>
<td>0.9931</td>
</tr>
<tr>
<td>Ribosomes</td>
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<td>70</td>
<td>0.03</td>
<td>8.8</td>
<td>0.12</td>
<td>7.9</td>
</tr>
<tr>
<td>FtsQ108 RNCs</td>
<td>117</td>
<td>114</td>
<td>0.12</td>
<td>7.9</td>
<td>0.12</td>
<td>7.9</td>
</tr>
</tbody>
</table>

FIGURE 1. Specific binding of SecA and ribosomes to the SecYEG complex monitored by SPR. Shown is an SPR sensogram of the binding of SecA (A) or non-translating ribosomes (B) to IMVs containing overexpressed levels of Cys-less SecYEG (solid), SecY(R357E)EG (spaced dashed), or SecY(R255E,R256E)EG (dashed). Binding was measured at 25 °C with a flow rate of 20 μl/min. Data were corrected for background binding to IMVs containing endogenous SecYEG levels. The ligand association and dissociation phases are represented above the sensograms by black and white bars, respectively. The concentration of SecA and ribosomes was 48 and 27 nm, respectively.

FIGURE 2. SecA, ribosomes, and RNCs interact with the SecYEG complex with nanomolar affinity. SecYEG binding at increasing concentrations of SecA (A), non-translating ribosomes (B), and FtsQ108 RNCs (C) was determined by SPR as described in Fig. 1. The equilibrium SPR responses were fitted by nonlinear regression analysis. The fitting for SecA was performed based on a model assuming single site with a nonspecific binding parameter. For fitting for non-translating ribosome and RNCs, a two-site saturation model was used. Binding data are summarized in Table 1.
cence microscopy, and biochemical methods (11, 26, 33, 36). In contrast, the ribosome binding responses fitted best to a two-site saturation model yielding apparent $K_D$ values of 0.03 and 8.8 nM, respectively (Fig. 2B, Table 2). Although the fit does not explain the mechanism of the interaction, the lower affinity value is in the same order as reported previously by equilibrium binding experiments, i.e. 14–17 nM with IMVs (17) and 6 nM with SecYEG proteoliposomes (37). These results demonstrate that both SecA and the ribosome interact with the SecYEG complex with nanomolar affinity.

**Increased Ribosome Binding in Presence of Nascent Chain**—During membrane protein insertion, the SecYEG complex interacts with translating ribosomes. To determine whether the presence of a nascent chain influences the SecYEG-ribosome interaction, ribosomes carrying chimeric nascent chains of the *E. coli* inner membrane protein FtsQ were isolated. To halt translation, constructs were used in which the stalling motif of the *E. coli* secretion monitor protein SecM (38) was fused behind the first 41, 51, or 72 amino acids of FtsQ, resulting in nascent chains of 77, 87, or 108 amino acids long (FtsQ77, FtsQ87, FtsQ108 (20), respectively). At these nascent chain lengths the FtsQ TMS is expected to be either half-exposed (FtsQ77), nearly completely exposed (FtsQ87), and completely exposed (FtsQ108) from the ribosomal exit site (4) (Fig. 3A).

Constructs were preceded by a triple STREP-tag to allow separation of RNCs from non-translating ribosomes (20). After isolation, the presence of the nascent chains was verified by immunoblotting against an STREP-tag antibody (Fig. 3B). Surprisingly, injection of the FtsQ87 and FtsQ108 RNCs resulted in an SPR response that was 3–5-fold higher compared with non-translating ribosomes (Fig. 3C), indicating that the SecYEG-ribosome interaction is strongly stimulated by the presence of a nascent chain. The shorter FtsQ77 RNC increased binding to a lesser extent (Fig. 3C), suggesting that the enhanced interaction of translating ribosomes with the SecYEG complex is dependent on the length of the nascent chain with an optimal interaction when the TMS of FtsQ is exposed from the ribosome tunnel. Indeed, injection of a FtsQ nascent chain of 108 residues without TMS (FtsQ108-/H9004TMS) yielded a significantly reduced binding response compared with FtsQ108 exposing the TMS (Fig. 3D), suggesting that TMS contributes to the enhanced binding of RNCs to the translocon.

The interaction between RNCs and SecYEG was further studied using the FtsQ108 construct that has previously also been used for cross-linking and structural studies (4, 6). As observed for non-translating ribosomes, the R255E,R256E mutations in the C4 loop of SecY abrogated the interaction with FtsQ108 RNCs (Fig. 3E, dashed line). However, these RNCs showed substantial interaction with SecY(R357E)EG (Fig. 3E, spaced dashed line), which is consistent with our previous data that indicated the R357E mutation still supports membrane protein insertion (14). These data demonstrate that translating and non-translating ribosomes interact differently with the SecYEG complex.

**Interaction of Purified SecYEG Complex with SecA and Ribosomes**—To verify the effect of a nascent chain on the SecYEG-ribosome interactions, we monitored the binding reaction between the purified components. Binding was probed...
using FCS, a sensitive confocal microscopy technique that allows for the analysis of molecular interactions in equilibrium at nanomolar substrate concentrations (39). In FCS, fluctuations of fluorescence resulting from fluorescently labeled proteins diffusing through the femtoliter-sized confocal excitation volume are monitored and auto-correlated over the measurement time. Temporal decay in the correlation function provides a precise estimate of the diffusional mobility of the fluorescently labeled biomolecule. Binding of the fluorescent protein to large non-labeled components can be detected as alterations of its mobility (Fig. 4A). First, the interaction between the solubilized SecYEG complex and ribosomes was determined in a solution containing 0.05% of the detergent DDM. Purified and fluorescently labeled SecY(C148)EG-AlexaFluor 488 (~100 nm) was illuminated when diffusing through the laser focal volume, and temporary fluctuations in the fluorescence intensity were recorded and used to build an autocorrelation curve (Fig. 4A). The FCS data were analyzed assuming unrestricted three-dimensional diffusion of the labeled SecYEG in solution, and diffusion coefficient \( (D) \) of \( 26 \pm 3 \times 10^{-8} \) cm\(^2\)/s was obtained. The addition of an excess SecA (1.8 \( \mu \)M) decreased the diffusion coefficient of solubilized SecYEG only slightly by ~10% to 23.9 \( \pm \) 1.5 \( \times \) 10\(^{-8} \) cm\(^2\)/s (data not shown). However, upon the addition of an excess FtsQ108 RNCs (250 nM), the mobility of SecY(L148C)EG-AlexaFluor 488 reduced substantially, resulting in a pronounced shift of the autocorrelation trace (Fig. 4A). The average diffusion time decreased ~3-fold, and the \( D \) reduced to 10.0 \( \pm \) 0.9 cm\(^2\)/s, suggesting that SecYEG was bound to RNC and diffused slowly as a part of the large complex. In agreement with the Einstein-Stokes equation, the observed reduction in the diffusion coefficient correlated with the 3-fold difference in dimensions of SecYEG protein enlarged by a belt of detergent molecules (radius \( R \approx 3 \) nm (40, 41) and the ribosome \( R \approx 10 \) nm (42)).

The addition of an excess (250 nm) of non-translating ribosomes to SecYEG reduced the translocon mobility to a lesser extent (Fig. 4A), suggesting that a large fraction of the SecYEG was not bound to these ribosomes and retained their high diffusional mobility. To quantify the binding efficiency, the autocorrelation traces of SecYEG were fitted assuming two SecYEG fractions in the ensemble: uncomplexed and ribosome/RNC-bound. The data suggested that more than 98% of SecYEG was bound with FtsQ108 RNCs, whereas only 67% was bound to non-translating ribosomes (Fig. 4B). As was reported previously (7), the R255E,R256E and R357E mutations in SecYEG reduced binding of non-translating ribosomes in the detergent environment (supplemental Fig. S2). However, in agreement with the SPR data presented above, the binding defect of the R357E mutant could be partially restored by the presence of a nascent chain (supplemental Fig. S2).

Because the above studies were performed in detergent solution, we also investigated the role of the lipids in SecA and RNC binding to SecYEG. Herein, fluorescently labeled SecY(C148)EG-AlexaFluor 488 was reconstituted into small lipid patches, known as nanodiscs (28). These provide a physiologically relevant lipid environment for the embedded membrane proteins (supplemental Fig. S3). Because nanodiscs are monodisperse in aqueous solution, they are well suited for fluorescence microscopy applications, including FCS. The diffusion coefficient of SecYEG reconstituted into nanodiscs (SecYEG-Nd) was similar to that of detergent-solubilized SecYEG \( (D = 27 \pm 3 \times 10^{-8} \) cm\(^2\)/s). In agreement with previous reports (8), no binding was observed for non-translating ribosomes. In contrast, the mobility of SecYEG-Nd decreased substantially in the presence of FtsQ108 RNCs, indicating that the presence of a nascent chain triggered ribosome binding to the membrane-embedded translocon. Fitting the autocorrelation curves with the two-component model suggested that 82%...
of the SecYEG formed complexes with RNCs (Fig. 4B). As observed with the native membrane-embedded and solubilized SecYEG complexes, the interaction of FtsQ108 with SecYEG-Nd was partially inhibited by the R357E mutation and more strongly blocked by the R255E,R256E mutation of SecYEG (supplemental Fig. S2). Together, these FCS results corroborate the observation made by SPR that the presence of the nascent chain strongly promotes the SecYEG-ribosome interaction the SPR data but also indicate that the interaction between SecYEG and the ribosome is influenced by the environment of the SecYEG complex (solubilized versus membrane-embedded) showing a greater interaction between empty ribosomes and SecYEG when present in detergent.

**Competitive Binding of SecA and Ribosomes**—It has been proposed that SecA and ribosomes interact with the translocon non-competitively (17). This would imply that ribosomes and SecA bind to different and independent sites on SecYEG. To investigate this phenomenon, we designed an SPR competition experiment to determine whether the SecYEG complex can accommodate both SecA and the ribosome simultaneously. First, the SecYEG complexes in the immobilized IMVs were saturated with SecA using a running buffer containing excess SecA (96 nM). Subsequent injection of ribosomes in the SecA-containing running buffer resulted in a binding response that was 2–3-fold lower than in the absence of SecA (Fig. 5, A and B). Stabilizing the SecYEG-SecA interaction by the addition of AMP-PNP, which prevents the dissociation of SecA from the SecYEG complex (26, 43), or a combination of ATP and azide, which prevents the release of SecA from SecYEG (44), further decreased the ribosome binding up to 8-fold (Fig. 5C). As a control, AMP-PNP or ATP-azide alone only slightly affected the SecYEG-ribosome interaction (Fig. 5C). When the ribosomes were loaded with the FtsQ108 nascent chain, the SecYEG interaction was affected even more severely by the presence of SecA (Fig. 5D). SecA has been reported to interact with ribosomes and RNCs (45, 46). However, the reduced interaction was not caused by sequestration of ribosomes or RNCs by the SecA present in the running buffer, as co-sedimentation assays showed that under the conditions used for SPR less than 3% of the RNCs were SecA-bound (Fig. 5E). This is consistent with the observation that the SecA-ribosome interaction is of low affinity (0.9 μM) (45). The results, therefore, indicate that SecA competes with both translating and non-translating ribosomes for SecYEG binding.

**Detergents Affect Interaction between SecYEG and Ligands**—Because the diffusion coefficient of SecYEG was strongly affected by binding of a ribosome, but not by SecA (see above), the RNC- and SecA-bound SecYEG pop-

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**FIGURE 5.** SecA competes with the ribosome for SecYEG binding. A, equilibrium SPR responses at increasing concentrations of non-translating ribosomes were measured in the presence (white) or absence (black) of SecA (96 nM) in the SPR running buffer. B, binding of non-translating ribosomes (6.75 nM) in the presence or absence of SecA (96 nM) is shown. C, normalized ribosome binding in the presence (white) or absence (black) of 96 nM SecA with or without AMP-PNP (1 mM) or ATP (1 mM) and azide (5 mM) is shown. Binding responses were normalized to the equilibrium response in the absence of SecA. D, binding of FtsQ108 RNCs (3.38 nM) in the presence or absence of a SecA (96 nM) is shown. E, co-sedimentation of SecA (92 nM) with ribosomes (60 nM), FtsQ108 (50 nM), and FtsQ108; ΔTMS (50 nM) is shown. The SecA input represents amount of SecA added to the reaction mixture. RU, response units.
the presence of 1 mM AMP-PNP, a clear competition for SecA and FtsQ108 RNCs were added to SecY(L148C)EG-Nd in SecYEG-Nd based on the translocon mobility only. When both inactivation between SecA and RNC FtsQ108 binding to
which is similar to SecYEG-Nd alone. This allows for a discrimi-
nated by AMP-PNP manifested a
alyzed SecYEG complex. The SecYEG-Nd-SecA complex stabil-
mine whether the detergent environment of the SecYEG com-
plex caused this apparent discrepancy with the SPR results, the
same experiment was performed using the nanodisc-reconsti-
tuted into nanodiscs. Although the molecular environment
influences the observed interactions, the interactions are spe-
cific and inhibited by mutations in SecY that were previously
reported to result in ribosome binding defects. Interestingly,
ribosome binding to the SecYEG complex was highly stimu-
lated by the presence of a nascent chain, in particular when this
nascent chain exposed a hydrophobic polypeptide sequence.
The presence of a nascent chain also partially relieved the ribo-
some binding defect of the SecY(R357E) mutant. Taken
together this suggests that translating and non-translating ribo-
somes interact differently with the SecYEG complex and indica-
tes an interplay, possibly a cooperative effect, of the nascent
chain in the SecYEG-ribosome interaction. A recent cryo-EM
structure of an FtsQ RNC complex with the translocon indi-
cated that the nascent chain engages in interactions between
the ribosomal proteins L23 and L24 and the loops of SecY,
resulting in conformational changes in the participating loops
(8). These interactions and conformational changes might
tighten the interaction between the RNC complex and the
translocon, and this may be promoted by hydrophobic stretches in the nascent polypeptide chain.

The interaction between translating or non-translating ribo-
somes and the membrane-embedded SecYEG complex was
competed by SecA, demonstrating that the two ligands
bind the translocon simultaneously. Surprisingly, even though
the presence of a nascent chain enhances the ribosome-translo-
con interaction, SecA competed stronger with translating than
with non-translating ribosomes. Because our current under-
standing of the exact mechanism of the translocon-RNC inter-
action is still limited, this observation is not readily explained.
However, it is clear from the SPR data that the interaction is not
a simple one-to-one binding event. Using nonlinear regression
analysis of the response levels at equilibrium, ribosome binding
data could only be fitted to a “two-site saturation” model
according to which SecYEG partly interacted with the ribo-
somes with an apparent affinity in the low nanomolar scale
consistent with previously reported values (17, 37) and partly
with a very high affinity that was particularly prominent upon
binding of RNCs. This two-site saturation model suggests that
the dramatically increased binding response in the presence of
a nascent chain is caused by an increase in the number of bind-
ing sites rather than an improved affinity. Possibly, RNCs
recruit binding sites that were previously masked. Alterna-
tively, binding of a RNC to a SecYEG complex causes an intrin-
sically higher SPR signal than binding of a non-translating ribo-
some, whereas the number of binding sites remains constant.
This could for instance occur when the nascent chain, by inter-
acting with SecYEG, pulls the ribosome closer to the membrane
surface, which in turn would result in an increased SPR signal.

SPR and FCS readily detected the association of ribosomes
with the SecYEG complex present in membrane vesicles and
the purified SecYEG present in detergent solution or recon-
stituted into nanodiscs. During the biogenesis of membrane proteins with large
periplasmic domains, both the ribosome and SecA need to
associate with the SecYEG complex to drive the membrane par-
titioning of hydrophobic segments and the translocation of hydrophilic domains, respectively. How the translocon is able
to coordinate the binding of these two ligands is not well under-
stood. Here we have used two distinct spectroscopic techniques
to follow the binding of SecA and of translating and non-trans-
lating ribosomes to the SecYEG complex.

Discussion

During the biogenesis of membrane proteins with large
periplasmic domains, both the ribosome and SecA need to
associate with the SecYEG complex to drive the membrane par-

ulations can be discriminated based on their diffusional mobi-
ity. SecA and FtsQ108 RNC were added to DDM-solubilized
SecY(L148C)EG-AlexaFluor 488 together, and the translocon
diffusion was monitored by FCS. AMP-PNP was added to sta-
bilize the SecYEG-SecA complex. Assuming that both SecYEG-
SecA and SecYEG-RNC complexes are formed within the
ensemble, a two-component model can be used for analysis.
Surprisingly, even in the presence of a large excess of SecA (1.8
µM versus 250 nM RNCs), 87% of the translocons were com-
plicated with the RNCs in detergent solution (Fig. 6). To deter-
mine whether the detergent environment of the SecYEG com-
plex caused this apparent discrepancy with the SPR results, the
same experiment was performed using the nanodisc-reconsti-
tuted SecYEG complex. The SecYEG-Nd-SecA complex stabi-
lized by AMP-PNP manifested a D of 23 ± 3 × 10^{-8} cm^2/s,
which is similar to SecYEG-Nd alone. This allows for a discrimi-
nation between SecA and RNC FtsQ108 binding to
SecYEG-Nd based on the translocon mobility only. When both
SecA and FtsQ108 RNCs were added to SecY(L148C)EG-Nd in
the presence of 1 mM AMP-PNP, a clear competition for
SecYEG binding was observed, and the propensity to form a
complex with the RNCs decreased with the SecA-RNC ratio
(Fig. 6). In the presence of an elevated AMP-PNP concentration
(5 mM), RNC binding was further reduced, up to 3-fold (supple-
mental Fig. S4). These results indicate that the interaction
between SecYEG, translating ribosomes, and SecA is strongly
dependent on the molecular environment; although detergents
promote ribosome binding and weaken the competitive inter-
action with SecA, SecA and translating ribosomes bind com-
petitively to the membrane-embedded SecYEG. Importantly,
the FCS data corroborate with the SPR data and demonstrate that SecA and (non)translating ribosomes compete for binding
to the membrane-embedded SecYEG complex.

**Discussion**

During the biogenesis of membrane proteins with large
periplasmic domains, both the ribosome and SecA need to
associate with the SecYEG complex to drive the membrane par-

![FIGURE 6. SecA competes with ribosomes for binding to SecYEG reconstituted into nanodiscs. Binding of RNCs to purified SecYEG in nanodiscs in the presence of increasing concentrations of SecA with AMP-PNP. The fraction of RNC-bound SecYEG complex was determined by FCS as described under “Results.” Detergent-solubilized SecYEG primarily interacted with RNC even in a large excess of SecA (white bar).](http://www.jbc.org)
In this scenario the interaction with RNCs would direct the SecYEG complex toward the high affinity state (see Table 2: 50% high affinity sites for RNCs versus 25% for non-translating ribosomes). This very tight binding of translating ribosomes to the translocon could be the result of multiple steps, so that inhibition of RNC binding by SecA could occur if the initial association is of relatively low affinity. Further studies will be required to elucidate the molecular basis of the ribosome translocon interaction.

SecA-ribosome competition only occurred when SecYEG was in its native, membrane-embedded state, as the presence of SecA did not affect the SecYEG-ribosome interaction in detergent solution. Because FCS indicated that solubilized SecYEG has a higher propensity to bind non-translating ribosomes compared with the membrane-embedded SecYEG (this study) and detergents have been shown to affect the activity and oligomeric state of SecA (47–49), it is likely that the interaction between SecA, the ribosome, and the solubilized SecYEG complex is affected by the presence of the detergent. The observation that the molecular environment is a critical factor in the interaction between the translocon and its soluble interactions partners should be considered in future studies.

The observation that SecA and the ribosome compete for binding to the SecYEG complex even when the ribosome is charged with a nascent chain of a SecA-dependent membrane protein implies that during membrane protein insertion, SecA and the ribosome do not interact with the translocon simultaneously. Because many membrane proteins contain large periplasmic domains, the competitive interaction of SecA and the ribosome has implications for the membrane insertion mechanism. Although for single-spanning membrane proteins like FtsQ it is possible to envision a scenario where the ribosome dissociates from the SecYEG complex after membrane partitioning of the TMS to allow SecA binding, the sequence of events will be more complicated when a large periplasmic domain is followed by another TMS. For these proteins, SecA and the ribosomes might need to cycle on and off from the translocon in a sequential manner, a process for which the timing needs to be carefully orchestrated. It is plausible that the SecYEG affinity to SecA and translating ribosomes depends on the hydrophobicity of the emerging polypeptide chain. The ribosome might be released from the SecYEG complex once it encounters a large enough polar periplasmic polypeptide domain that would allow SecA to bind to this domain followed or concomitantly with SecYEG binding (Fig. 7). The micromolar SecA concentrations in the cytoplasm would ensure efficient cycling of SecA on SecYEG to translocate these polar domains (33). Likewise, SecA may dissociate from the SecYEG complex once it encounters a newly emerging TMS. In this respect, SecA has been reported to dissociate from the SecYEG complex when it encounters a potential TMS during the translocation of a secretory protein (50, 51). Early recognition of a TMS by the translating ribosome (52) may restore the high affinity interaction between SecYEG and the ribosome for the insertion of the next TMS. Future studies should be directed to unravel the interplay between SecA, RNCs, and the SecYEG complex including targeting factors such as signal recognition particle and FtsY (53).

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**Fig S1. Overexpression of SecYEG mutants.** (A) IMVs containing endogenous SecYEG levels or overexpressed Cys-less SecYEG, SecY(R255E,R256E)EG or SecY(R357E)EG was analyzed by SDS-PAGE and Coomassie brilliant blue staining. (B) Translocation of fluorescein labeled proOmpA(C290S) into IMVs containing the indicated overexpressed SecYEG mutants.
Fig. S2. Interaction between ribosomes and RNCs and purified SecYEG mutants. Fluorescence correlation spectroscopy (FCS) was used to quantify binding of ribosomes and RNCs to SecY(L148C)EG-AlexaFluor 488 (black bars), and its derivatives bearing the SecY mutations R357E (grey bars) or R255E,R256E (white bars). Binding was analyzed both for detergent-solubilized and nanodisc-reconstituted SecYEG. SecYEG was reconstituted into nanodiscs in parallel preparations using SecYEG:MSP:lipid ratio of 1:8:200, and fraction 15 of size-exclusion chromatography was used for the binding assay. Analysis of FCS data is described in the main text.
**Fig. S3. Reconstitution of SecYEG in nanodiscs.** SecY(L148C)EG conjugated with AlexaFluor 488 fluorophore (AF488) was reconstituted with phospholipids in presence of MSP1D1 scaffold protein (MSP). MSP-encapsulated nanodiscs were isolated using size-exclusion chromatography, and collected in 1 mL fractions. SDS-PAGE confirmed that SecYEG and MSP co-eluted in fractions 12 to 16, as visualized by (A) fluorescence and (B) Coomassie staining. (C) Individual nanodiscs (Fraction 15) were imaged using cryo-electron microscopy, and showed as circular entities of ~10 nm diameter (encircled in white).
Fig. S4. AMP-PNP-stimulated SecA binding abolishes SecYEG:RNC interactions. FCS was used to quantify RNC binding to nanodisc-reconstituted SecY(L148C)EG-AlexaFluor 488 in the absence and presence of increasing amounts of SecA. AMP-PNP at 5 mM was used to promote SecA binding to the translocon. In presence of SecA an up to 3-fold reduction in SecYEG:RNC binding was observed.