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Published in:
The Journal of Biological Chemistry

DOI:
10.1074/jbc.M116.743831

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Document Version
Publisher's PDF, also known as Version of record

Publication date:
2016

Link to publication in University of Groningen/UMCG research database

Citation for published version (APA):

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Lipids Activate SecA for High Affinity Binding to the SecYEG Complex*

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Received for publication, June 17, 2016, and in revised form, August 30, 2016 Published, JBC Papers in Press, September 9, 2016, DOI 10.1074/jbc.M116.743831

Protein translocation across the bacterial cytoplasmic membrane is an essential process catalyzed predominantly by the Sec translocase. This system consists of the membrane-embedded protein-conducting channel SecYEG, the motor ATPase SecA, and the heterotrimeric SecDFyxJC membrane protein complex. Previous studies suggest that anionic lipids are essential for SecA activity and that the N terminus of SecA is capable of penetrating the lipid bilayer. The role of lipid binding, however, has remained elusive. By employing differently sized nanodiscs reconstituted with single SecYEG complexes and comprising varying amounts of lipids, we establish that SecA gains access to the SecYEG complex via a lipid-bound intermediate state, whereas acidic phospholipids allosterically activate SecA for ATP-dependent protein translocation.

The exact targeting mechanism of SecA to the membrane and the dynamics of its interaction with the SecYEG channel are poorly understood. Studies using cell fractions have shown that SecA cycles between the cytosol and the cytoplasmic membrane (5), which was suggested to be ATP-dependent (6). As shown with liposomes, SecA binds with low affinity to lipids, a process that is enhanced by the presence of negatively charged lipids (7, 8). In contrast, no binding was found to inner membrane vesicles (IMVs)3 that lack the negatively charged lipid phosphatidylglycerol (7). In the free soluble state, SecA is inactive for ATP hydrolysis and exhibits only poor peptide binding (7). In the lipid-bound state, SecA is thermolabile but is stabilized by the presence of unfolded secretory proteins, an activity that is termed SecA lipid ATPase. SecA binds with high affinity to the membrane-embedded SecYEG complex (K_D = 4.5 nM) (9), but it shows only low affinity binding to the detergent-solubilized SecYEG (3.9 μM) (10). Important, acidic phospholipids such as phosphatidylglycerol are essential for protein translocation. In vitro, the signal sequence of secretory proteins have been shown to bind, fold, and penetrate membranes containing acidic phospholipids. These experiments indicate that not only the presence but also the type of lipid might play a role in the targeting and/or functioning of SecA to the membrane, but an exact role for lipid binding has never been demonstrated.

The crystal structure of the SecA-SecY complex in solution has provided new insights into the binding mechanism of SecA (4). Binding mostly occurs through cytosolic loops 6–7 and 8–9 of SecY via electrostatic interactions to the polypeptide-cross-linking and helical scaffold domains of SecA. However, there are no distinct interactions with phospholipids that emerge from the structure. The SecA N terminus was shown earlier to be involved in lipid binding (11). This N terminus is not conserved, but its highly amphipathic nature is omnipresent. Because of its net positive charge, this region of SecA is predicted to be membrane surface seeking interacting with...

* This work was supported by the Stichting voor Fundamenteel Onderzoek der Materie (FOM). The authors declare that they have no conflicts of interest with the contents of this article.

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3 The abbreviations used are: IMV, inner membrane vesicle; OmpA, outer membrane protein A; DHFR, dihydrofolate reductase; Apo, apolipoprotein; MSP, major scaffold protein; DOPC, 1,2-dioleoyl-sn-glycerol-3-phosphocholine; DOPG, 1,2-dioleoyl-sn-glycerol-3-phosphoglycerol; DOPE, 1,2-dioleoyl-sn-glycerol-3-phosphoethanolamine; SEC, size exclusion chromatography; AFM, atomic force microscopy; MST, microscale thermophoresis; FCCS, fluorescence cross-correlation spectroscopy; NTA, nitritoltriacetic acid.
acidic phospholipids (12). Deletion of the N terminus results in the inactivation of SecA, but activity can be restored by replacing the N terminus with a His tag and supplementing SecYEG proteoliposomes with Ni⁺-NTA lipids, suggesting that membrane tethering is important for functioning (11). In the SecA-SecY structure, however, the helical amphipathic N terminus of SecA is positioned away from where the membrane would be located, and a major conformational change involving a 30 Å translational movement would be required to allow this region to deeply penetrate the membrane, which could potentially impact the SecY binding mode and SecA function. This N-terminal displacement of SecA suggests not only a tethering function of the N terminus but also a key role function in conformational activation of SecA upon lipid binding.

Earlier studies have shown that the presence of negatively charged lipids is essential for the activity of the Sec translocone. However, the actual role of the lipid bilayer in the translocation process remained to be elucidated. Here, we have used two different sizes of nanodiscs harboring single SecYEG complexes surrounded by different quantities of lipids to study the functional interaction between SecYEG and SecA. Our data suggest that high affinity binding of SecA to SecYEG is dependent on the presence of bulk acidic phospholipids. We further show not only that the SecA N terminus that interacts with acidic phospholipids is important to tether SecA to the membrane but also that this binding event induces a conformational change of SecA that promotes its interactions with SecYEG. Our data suggest that the lipid bound SecA is a true intermediate in the catalytic cycle and provides an explanation why SecA is primed for high affinity SecYEG binding upon its interaction with acidic phospholipids. We propose a new mechanism of protein translocation, whereby SecA first binds acidic phospholipids in the membrane whereupon the lipid bound SecA intermediate interacts with SecYEG with high affinity.

Results

Formation of SecYEG-containing Small and Large Nanodiscs—To examine the influence of phospholipids on the reconstituted SecA-SecYEG complex, two different nanodisc systems were employed. Nanodiscs are lipid patches that are formed by a protein belt, i.e. a scaffold protein. By changing the scaffold protein, the size and therefore the amount of lipids in the nanodisc can be changed (Fig. 1). To generate small nanodiscs, the membrane scaffold protein MSP1E3D1 (13) was used. This scaffold protein is derived from the apolipoprotein A-1 and forms a two-copy helical belt, yielding discs of ~13 nm (13). In contrast, for large nanodiscs, the scaffold protein ApoE422k, which is the 22-kDa fragment of apolipoprotein E-4, was employed (14). For this scaffold protein, nanodiscs sizing from 14.5 to 28 nm have been reported (15). Importantly, with ApoE422k, the ratio of scaffold protein to lipid determines the disc size. Computer simulations have shown that each additional copy of ApoE422k increases the disc diameter by ~4.5 nm (15). By using both nanodiscs systems, it is possible to generate compartments where the SecYEG channel is embedded by low or high lipid quantities, respectively (See below). In the nanodiscs, the copy number of reconstituted SecYEG per disc can be determined via Poisson distribution. Because single

SecYEG complexes are sufficient for protein translocation (16), a SecYEG to lipid molar ratio (0.25:1800) was chosen that favors the formation of discs with single SecYEG complexes. This method was used previously for the small nanodiscs (17), but the increased size of the large nanodiscs caused us to ascertain the above assumption experimentally. According to the molar ratio and ~1100 lipid molecules/31-nm nanodisc (assuming 8 copies of ApoE422K (15)), 16% of the discs are expected to contain a single copy, whereas 80% will be empty, and less than 3% will contain multiple copies of SecYEG, as shown previously (18). Nanodiscs were formed using sodium cholate as described (19), with a lipid composition of DOPC:DOPA:DOPG (40:30:30 molar ratio) (16). Small and large nanodiscs were subjected to size exclusion chromatography (SEC) and analyzed by SDS-PAGE. Peak fractions (fractions 19–26 for large discs and fractions 29–34 for small discs) were pooled (Fig. 2A) and further analyzed by atomic force microscopy (AFM) (Fig. 2B). The diameters of the discs (n = 200 for both large and small discs) were analyzed, plotted in a histogram, and fitted to a Gaussian model (Fig. 2C). The mean diameter of the small nanodiscs was found to be ~12.7 nm (σ = 4 nm), which is consistent with data from previous studies (13). The diameter of the large nanodiscs was found to be 31 nm (σ = 9 nm). To demonstrate that the discs consist of a lipid bilayer, the height of large discs was measured (n = 200) (Fig. 2D). On average, the discs had a thickness of 3.8 nm (σ = 0.3 nm). Considering that the lipids used in this study had a acyl chain length of 18 carbon atoms, the thickness value is in good agreement with the expected values for single lipid bilayers (20). Moreover by AFM, the presence of SecYEG complexes was detected being evident because of an increase in height, which could represent the periplasmic or cytoplasmic loop of SecYEG (21) (Fig. 2E). Examination of a set of large nanodiscs (n = 200) showed that 14.5% of the discs showed such elevations, which is in agreement with the theoretical expected distribution of single SecYEG complexes over the nanodiscs (Fig. 2D). With the experimentally determined average sizes of the small and large nanodiscs of 12.7 and 31 nm,
the number of lipids present in these discs amounts to ~160 and 1100, respectively, assuming 0.6 nm² as the lipid head surface area (22). To account for the space occupied by single SecYEG complexes, small and large SecYEG nanodiscs contain 120 and 1060 lipids, respectively, assuming a surface area of 20 nm² for the SecYEG channel.

To further validate the monomeric state of the SecYEG complexes in the large nanodiscs, Fluorescence cross-correlation spectroscopy (FCCS) was performed. Herein, SecY¹⁴₈EG was exposed in the label reaction to Atto647N and Alexa Fluor488 to ensure that each complex had either one or the other attached to it with equal probability. The labeled complexes were reconstituted into large nanodiscs as reported previously for small nanodiscs (17). The autocorrelation of the fluorescence of both fluorophores was recorded, and the cross-correlation (black) was determined. To ensure a monomeric state, SecYEG, ApoE₄₂₂k, and lipids were mixed in a molar ratio of 0.25:10:1800. The monomeric state of SecYEG in the discs was confirmed by a low cross-correlation. In contrast the oligomeric state, represented by high cross-correlation, was achieved when the amount of SecYEG was increased using SecYEG, ApoE₄₂₂k, and lipid molar ratio of 1:10:1800.
Efficient FRET.

Precursor protein proOmpA cannot be translocated via the SecYEG pore, which stalls translocation and brings the donor-acceptor pair in close proximity for efficient FRET. B, SecYEG proteoliposomes (black), SecYEG large discs (blue), or SecYEG small discs (gray) were incubated in the presence of Cy3-conjugated proOmpA-DHFR and 50 nM (left panel) or 1000 nM (right panel) SecA. Translocation was initiated by the addition of ATP, and the formation of a stable SecYEG-preprotein intermediate was recorded following the acceptor fluorescence. C, ProOmpA translocation as a function of SecA concentration using SecYEG proteoliposomes (black circles), SecYEG large discs (white squares), or SecYEG small discs (white circles). The acceptor fluorescence signal after addition of ATP (time window $t = 70–120$ s) was plotted against a logarithmic time scale, and data points were fitted linearly with the equation $y = a + bx$, where $a$ represents initial intensity with $b$ as translocation rate. The translocation rate was plotted against the SecA concentration. Large nanodiscs are highly active and support translocation at low SecA concentrations (50 nM), whereas small nanodiscs need a very high SecA concentration ($1–2 \mu M$).

**Lipid-dependent SecA Activity**

To study the influence of the available lipid surface on the translocation efficiency, the SecA-dependent translocation by SecYEG was measured using a FRET-based translocation assay (16). Herein, the preprotein proOmpA fused at its C terminus to a dihydrofolate reductase (DHFR) domain was used. This domain can be folded in the presence of methotrexate and NADPH, so that the translation of proOmpA–DHFR via SecYEG is stalled as the bulky, folded DHFR domain blocks further translocation. The unique cysteine at position 282 of proOmpA–DHFR and Cys148 at the periplasmic side of SecYEG were labeled with the FRET pair Cy3 and Atto647N, respectively. When both fluorophores get in close proximity, an increase of the FRET signal is detected reminiscent of translocation in the compartment-less system (Fig. 3A).

SecYEG proteoliposomes and SecYEG reconstituted in small or large nanodiscs were incubated with fluorophore-conjugated proOmpA–DHFR and SecA until a steady fluorescent signal was achieved. Translocation was initiated by addition of 2 mM ATP, which resulted in the formation of a SecYEG-preprotein translocation intermediate as evidenced by an increase of the acceptor fluorescence (Fig. 3B). As expected, FRET was strictly dependent on the presence of ATP and SecA (data not shown) (16). Both nanodiscs support translocation, but with SecYEG reconstituted in the large nanodiscs, already low SecA concentrations (~50 nM) sufficed to observe an efficient FRET signal. This concentration of SecA compares favorable with the SecA dependence of protein translocation in SecYEG proteoliposomes (23). In contrast, for SecYEG-containing small nanodiscs, very high SecA concentrations (~1 mM) were needed to obtain a FRET signal indicating very inefficient translocation (Fig. 3C). These data suggest that the presence of a larger available lipid surface in the large nanodiscs allows for efficient translocation.

**SecA Binding to SecYEG Is Dependent on the Lipid Surface**

To determine whether the translocation deficiency of SecYEG in small discs was due to an impeded interaction of SecA with SecYEG, the SecA-SecYEG binding was determined using microscale thermophoresis (MST). With this method, the movement of a fluorescently labeled molecule along a temperature gradient is traced. By applying heat to these molecules, their hydration entropy decreases, which results in an enhanced diffusion out of the heated spot. This effect can be monitored by a decrease of fluorescence in the heated area. When adding a binding partner, the hydration of the fluorescently labeled protein will change, resulting in a different movement along the temperature gradient, which can be detected by an altered, usually slower decrease of fluorescence. Such binding events can be transformed into a binding curve. To employ the MST method, fluorescently labeled SecYEG in small or large nanodiscs was titrated with increasing amounts of SecA, and the temperature gradient movement of the nanodiscs was traced (Fig. 4A). SecA binds with high affinity to SecYEG present in large nanodiscs ($K_D \sim 300$ nM) with a Hill coefficient of 1, indicating a noncooperative binding between two SecA dimers and one SecYEG complex. In contrast, SecA binding to SecYEG in small nanodiscs occurred with a very low affinity ($K_D \sim 3 \mu M$). Instead of the synthetic lipid mixture, SecYEG was reconstituted in large nanodiscs comprising native *Escherichia coli* lipids. Binding of SecA to SecYEG in native lipids was slightly less efficient compared with the synthetic lipid mixture DOPC:DOPG (40:30:30 molar ratio) (Fig. 4A). No SecA binding was detected, when SecYEG large discs were used that lacked the anionic lipid DOPG, DOPC:DOPG (40:60, molar ratio). This is in good agreement with earlier studies showing that anionic lipids are essential for protein translocation (7, 8).

To investigate the SecA binding in the absence of SecYEG, the ApoE422k was fluorescently labeled. Now a linear increase of the binding to large nanodiscs was observed, which did not saturate (Fig. 4B). The nonsaturating binding behavior of SecA suggests nonspecific lipid binding. Taken together, these results indicate that the available lipid surface is an important factor in high affinity SecA-SecYEG binding.

**Lipids Induce a Conformational Change to SecA to Prime It for High Affinity SecYEG Binding**

Previously, it was shown that the N terminus of SecA binds acidic phospholipids (11, 12) and is capable of penetrating the lipid bilayer (24) to function solely to tether the SecA to the lipid bilayer (11). However, in the *Thermotoga maritima* SecA-SecYEG complex structure in detergent, the N terminus of SecA is positioned such that it

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**FIGURE 3.** Large lipid surface enhances SecA-dependent translocation. A, principle of real time FRET-based assay. The prefolded DHFR domain fused to the precursor protein proOmpA cannot be translocated via the SecYEG pore, which stalls translocation and brings the donor-acceptor pair in close proximity for efficient FRET. B, SecYEG proteoliposomes (black), SecYEG large discs (blue), or SecYEG small discs (gray) were incubated in the presence of Cy3-conjugated proOmpA-DHFR and 50 nM (left panel) or 1000 nM (right panel) SecA. Translocation was initiated by the addition of ATP, and the formation of a stable SecYEG-preprotein intermediate was recorded following the acceptor fluorescence. C, ProOmpA translocation as a function of SecA concentration using SecYEG proteoliposomes (black circles), SecYEG large discs (white squares), or SecYEG small discs (white circles). The acceptor fluorescence signal after addition of ATP (time window $t = 70–120$ s) was plotted against a logarithmic time scale, and data points were fitted linearly with the equation $y = a + bx$, where $a$ represents initial intensity with $b$ as translocation rate. The translocation rate was plotted against the SecA concentration. Large nanodiscs are highly active and support translocation at low SecA concentrations (50 nM), whereas small nanodiscs need a very high SecA concentration (~1–2 $\mu M$).
would not contact the lipid bilayer (Fig. 5A). A relocation of the N terminus to the lipid bilayer can occur only when SecA undergoes a large conformational change. To discriminate between a tethering function and a conformational change, we designed a SecA mutant harboring a 10-amino acid linker after the first 20 amino acids of the N terminus (SecALinker). If the N terminus is solely responsible for membrane tethering, the linker insertion should not affect the activity. In contrast, the linker would disrupt a lipid induced conformational function of the N terminus. Plasmids bearing wild type SecA, SecALinker, and the N-terminal truncated SecAΔN20, which was previously shown to be inactive (11), were transformed into E. coli BL21.19 to test for complementation of the SecA function (Table 1). This strain harbors a secATs mutation and is not viable at non-permissive temperatures (42 °C), whereas growth can be restored by a plasmid-based SecA expression. Although the SecALinker mutant was able to complement the SecA deficiency, it did so with a much lower efficiency than the wild type SecA.

**FIGURE 4.** SecA-SecYEG binding affinity is dependent on lipid surface area. MST was performed to measure the binding of SecA to nanodiscs. A, SecA binding curves of SecYEG reconstituted in small (black squares) or large (black and white circles and white squares) nanodiscs as a function of the SecA concentration. With the large nanodiscs, the lipid composition was changed showing SecYEG in native E. coli lipids (white squares), DOPG:DOPC:DOPE, 30:40:30 (molar ratio) (black circles), and DOPC:DOPE, 40:60 (molar ratio) (white circles). B, SecA binding curves of lipid-filled (white circles) and SecYEG-reconstituted (black circles) large discs as a function of SecA concentration. Although SecA binds to empty discs in a nonsaturable manner (linear) up to 10 μM SecA, the interaction with SecYEG discs remains specific (sigmoidal). The experiments were performed in triplicate.

**FIGURE 5.** The SecA N terminus allosterically activates SecA for high affinity binding to SecYEG. A, side view of the T. maritima SecA-SecYEG complex (4). The trimeric SecYEG complex is highlighted in red, yellow, and blue, respectively. The N terminus of SecA (gray) is highlighted in magenta and demonstrates the remote location from the membrane. To penetrate the membrane, the N terminus would have to perform an ∼30 Å translational movement (arrow). B, E. coli BL21.19 temperature-sensitive secA strain, expressing plasmid-borne SecA mutants, was grown at permissive (30 °C) or nonpermissive (42 °C) temperature. When overexpressed (lower panel), SecA and a SecALinker could rescue the phenotype at the nonpermissive temperature. SecAΔN20 is inactive but also complements when overexpressed at a low level. C, SecA (variant) sample was supplemented with SecYEG proteoliposomes, and the ATPase activity was measured in the absence and the presence of proOmpA. SecAΔLinker (gray bars) shows low enzymatic activity at low concentrations (50 nM) but close to wild type (black bars) activity at high concentrations (500 nM). D, the SecA-dependent translocation of proOmpA into SecYEG proteoliposomes was plotted against the SecA concentration. SecAΔLinker (gray bars) shows a low translocation activity at low concentrations but wild type (black bars) activity at 2000 nM. The N-terminal deletion (white bar) of SecA disrupts its function. The asterisk represents the SecYEG proteoliposome sample that lacked the polar lipid DOPG, which caused a translocation deficiency for SecA (mutants). E, MST-based SecA binding curves to SecYEG large discs for the wild type SecA (black squares), SecAΔLinker (dark gray diamonds), and SecAΔN20 (light gray circles). Although SecAΔLinker and SecAΔN20 show a decrease in binding affinity, they showed saturation of binding at very high concentrations. The experiments were performed in triplicate.
In contrast, the deletion of the N terminus was lethal (Fig. 5B). When the SecA (variants) were overexpressed in the same strain, the SecA linker could largely restore growth, whereas SecAΔN20 hardly complemented.

To investigate the SecA (variant) activity in vitro, an ATPase activity assay in the presence of SecYEG proteoliposomes was performed (Fig. 5C). Thereby, the ATPase hydrolysis activity of SecA in the absence (basal ATPase) and the presence of proOmpA (translocation ATPase) was determined by measuring the free phosphate concentration using a malachite green reagent. At low concentrations (50 nM), SecA (variants) barely showed ATPase activity. In the presence of proOmpA the translocation ATPase activity of wild type SecA increased, whereas the mutants were still barely active. Interestingly, at high SecA concentrations (500 nM), SecA linker performed ATPase activity close to wild type level in the absence and presence of proOmpA.

To further examine the concentration-dependent SecA linker activity, proOmpA translocation assays were performed with SecYEG proteoliposomes. Translocation of proOmpA into the SecYEG proteoliposomes results in the appearance of proOmpA that is protected against externally added proteinase K. The formation of proteinase K-protected proOmpA is dependent on ATP, whereas the rate of translocation saturates at ~200 nM SecA (Fig. 5D). As expected, SecAΔN20 did not show any translocation activity. However, the SecA with the linker insertion was barely active at nominal SecA concentrations (200 nM), but at very high concentration (~500 nM) was supported proOmpA translocation at rates comparable with those observed with wild type SecA at much lower concentrations.

To investigate the reason for the remarkable concentration-dependent activity of SecA linker, the binding of SecA to SecYEG reconstituted into large nanodiscs was tested (Fig. 5E). In comparison with wild type SecA, the binding affinity of SecA linker is strongly reduced. However, at very high concentration (2 μM), saturation of binding was detected, demonstrating that the linker insertion impacts translocation by reducing the SecA-SecYEG binding affinity. Interestingly, SecAΔN20 shows a similar reduced binding affinity, but this mutant is also inactive at high concentrations. These data therefore suggest that the lipid interaction of the N terminus of SecA, in addition to tethering the SecA to the membrane, functions by allosterically activating the SecA for high affinity SecYEG binding and ATP hydrolysis.

### Discussion

During protein translocation, SecA and SecYEG form a functional interaction unit. A crucial step in this process is the targeting of SecA to the cytoplasmic membrane. Although it has been shown that anionic lipids are crucial for the SecA function (7, 8), the exact role of the lipids has remained elusive. Here, we designed small and large nanodiscs, containing a single copy of SecYEG surrounded by low or high lipid quantities, respectively. The formation of nanodiscs was confirmed by size exclusion chromatography and AFM. It demonstrated that SecYEG and the scaffold protein MSP1E3D1 (for small discs) or ApoE422k (for large discs) eluted in one fraction during the purification. Small discs showed an average diameter of 12.7 nm, which is in good agreement with the 13 nm reported before (13). The large nanodiscs had a size of 31 nm. However, in comparison with the small discs, the size distribution of the large discs was much broader. Considering that MSP1E3D1 always forms a two-copy belt around the lipids, whereas the copy number of ApoE422k can vary, the broader size distribution is not surprising. It has been suggested that the ApoE422k to lipid ratio determines the particle size (15). A previous study using the same reconstitution ratio as described in this work (ApoE422k/lipid of 10:1300) reported a monodisperse disc size of 23 nm. However, the disc sizes increased and a broader size distribution was achieved when the lipid composition was changed from POPC to a more complex lipid composition (19). Therefore the disc sizes reported here are in good agreement with early studies.

The SecYEG complex was reconstituted such that according to Poisson distribution a monomeric state was achieved in 16% of the discs, whereas 80% remained empty. It has been reported earlier that the disc size increased when the membrane protein bacteriorhodopsin was reconstituted (25). With bacteriorhodopsin, not only monomeric but also trimeric states were achieved. Here, the presence of reconstituted single SecYEG complexes did not have an effect on the disc size. When measuring the height of the large discs, some discs showed a local increase in height. These height increases have been shown in earlier studies to correspond to the periplasmic and cytoplasmic loop of SecYEG (21). Therefore, the local height increases allowed us to determine the number of discs containing...
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SecYEG. Approximately 14.5% of the discs showed an increased height representing reconstituted SecYEG complexes, which is in good agreement with the Poisson distribution calculation to determine the reconstitution efficiency. To further assess the oligomeric state of SecYEG in the large discs, a FCCS experiment was performed. Thereby the cross-correlation between differently labeled and reconstituted SecYEG complexes was determined. For large discs the cross-correlation was determined to be less than 10%, which is due to excitation cross-talk and unspecific double-labeling as reported earlier (17). When the SecYEG to ApoE422k ratio was changed from 0.25:10 to 1:10, statistically resulting in 14% of the discs containing multiple copies of SecYEG, a cross-correlation of 50% was detected. Overall, these data are in good agreement with the FCCS experiments performed by Taufik et al. (17) demonstrating a monomeric SecYEG state in small discs.

Based on our findings and the resulting average size distribution of the small and large nanodiscs, the single reconstituted SecYEG complexes will be surrounded by ~120 and 1060 phospholipids, respectively. Strikingly, SecYEG complexes present in the small nanodiscs are barely active for protein translocation using a FRET assay reported previously (17). Only at very high SecA concentration, activity is detected. In contrast, the single SecYEG present in the large nanodiscs is active already at the SecA concentrations needed to induce protein translocation in SecYEG proteoliposomes or IMVs. The translocation activity of SecA in the nanodiscs as addressed with a FRET based assay correlates with the ability to bind SecYEG. Although in the large nanodiscs, protein translocation already was detected at 50 nM SecA, very high SecA concentrations were needed to support protein translocation in the small nanodiscs (i.e. up to 1 μM). SecYEG present in large nanodiscs showed a KD for SecA binding of ~300 nM, whereas with the small nanodiscs a KD of ~3 μM was obtained. The latter is in the same order of magnitude as binding as the KD of SecA to detergent-solubilized SecYEG (~3.9 μM) (10). This poor binding affinity might be explained by several aspects. According to the dimensions of SecYEG, ~20% of the small discs will be occupied by the translocation complex, leaving only a low lipid surface area unoccupied (26). Therefore, the binding of SecA to the disc might be hindered because of spatial interference with SecYEG. This idea is supported by the observation that SecA-empty disc binding curves show a linear increase and, as expected, no saturation. The data imply that SecA binds unspecifically to lipids but specifically to SecYEG. Compared with proteoliposomes, SecYEG large discs still show a lower binding affinity for SecA (1–3 nM versus 300 nM, respectively). This difference could suggest that although the disc size was increased, the lipid area is still not large enough to support the most efficient binding. Further, as shown by AFM, only 14.5% of the discs contained a copy of SecYEG. Although not labeled, the remaining 85.5% empty discs can be bound unspecifically by SecA. Even though unspecific binding of SecA to empty discs was only observed at high SecA concentrations, we cannot exclude that lipid binding interfered with the KD calculations from the MST data. Therefore the calculation yields an apparent KD. Further, a binding defect could be due to the planar organization of the bilayer in the discs, possibly providing a different lateral lipid pressure as compared with curved liposomes. It is well established that SecA binds acidic phospholipids through its amphipathic N terminus. This region of SecA is known to penetrate the lipid bilayer as shown with Langmuir planar lipid monolayers (6) and membrane vesicles (27). Possibly, a spherical shape of the liposomes favors membrane insertion of the N terminus as compared with the planar nanodiscs bilayers.

To investigate the importance of the lipid composition, SecA binding to SecYEG large discs comprising native E. coli lipids was also measured. E. coli cytoplasmic membranes contain ~25% PG (28), which is similar to the DOPG concentration of the synthetic lipid mixture used in this study. SecA binding was slightly less efficient when SecYEG was reconstituted into native E. coli lipids. No SecA binding was detected when the SecYEG large discs lacked the anionic lipid DOPG, which is consistent with earlier studies showing that this lipid mixture also does not support translocation (7, 8, 23).

Our data support the notion that SecA first needs to bind to acidic phospholipids via ionic interactions with its N terminus before it can bind SecYEG with high affinity. Previously, we have shown that acidic phospholipids form a annulus around the SecYEG channel (29), and we hypothesize that this phenomenon contributes to the SecA binding and activity. The N terminus may have two distinct functions: a membrane tethering and/or an allosteric function, whereby lipid binding induces a conformational change on SecA. In this respect, in the T. maritima SecA-SecYEG complex structure, the N terminus of SecA is positioned such that it would not contact the lipid bilayer. It is important to note here that the crystal structure was produced with detergent-solubilized protein in the absence of a membrane. However, as predicted from this structure, at least a 30 Å translational movement of the N terminus of SecA is needed to penetrate the membrane. Given the proximity of the N terminus of SecA to nucleotide binding fold 1, such reallocation is predicted to evoke a conformational change to SecA that may directly affect the ATPase activity and the ability of SecA to binding SecYEG. To discriminate between a sole tethering function and the proposed conformational change or a combination of both, a SecA linker mutant was constructed that contained a flexible 10 amino acid linker after the first 20 N-terminal residues. This linker should not disrupt the N-terminal tethering function but should no longer or less efficiently be able to inflict the lipid binding-dependent proposed conformational change. Indeed, the activity of SecA linker was substantially reduced both in vitro and in vivo, but activity was fully restored when high levels of SecA linker were used. In contrast, removal of the N-terminal 20 amino acid (SecAΔN20) rendered SecA essentially inactive even when tested at high concentrations, consistent with previous studies (11). Both the SecA linker and SecAΔN20 showed a reduced ability to bind SecYEG as compared with the wild type SecA. The residual lipid binding of SecAΔN20 might relate to the C terminus that has been shown to also bind to lipids (30). Importantly, the observation that high levels of SecA linker restore the activity is consistent with our proposed allosteric binding mechanism in which SecA is initially recruited to the membrane via a lipid-bound intermediate whereupon it changes its conformation thereby becoming
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TABLE 2

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Experimental Procedures

Cloning Procedure—Wild type secA and secA mutants (Table 1) were cloned into pTrc99A. Standard cloning techniques were used to generate the N-terminally truncated SecA (SecAΔN20) that lacks amino acids 1–20. The generation of secAΔ linker, with an (SAG)2(SAAG) linker inserted after the first 20 N-terminal amino acids, was carried out by overlap PCR using the primers indicated in Table 2.

In Vivo Complementation Assay—Plasmids encoding for wild type SecA, SecAΔN20, and SecAΔ linker were transformed into the secATs mutant strain E. coli BL21.19 and tested for the ability to complement the SecA deficiency by growing on LB plates at nonpermissive temperatures as described previously (33). For overexpression, plates were supplemented with 10 μM isopropyl 1-thio-β-D-galactopyranoside.

Protein Production and Purification—E. coli BL21 (DE3) harboring wild type SecA or SecA mutants, designed in this study, was grown at 37 °C to an A500 of 0.6, whereupon protein expression was induced by addition of 0.5 mM isopropyl 1-thio-β-D-galactopyranoside. After 2 h of growth, the cells were harvested at 6000 x g for 15 min at 4 °C, resuspended in 25 mM HEPES/KOH, pH 6.5, and stored at −80 °C. SecA was purified as described before (34).

The proOmpA derivative fused to dihydrofolate reductase (proOmpA-DHFR) was overexpressed from pET504 in E. coli DH5α, purified from inclusion bodies, and stored in 8 M urea as described previously (35). ProOmpA-DHFR harboring a cysteine mutation at position 282 in the OmpA domain was labeled with Cy3. Free dye was removed by TCA precipitation.

SecYEG was overexpressed in E. coli SF100 and purified from IMVs as described before (35). Briefly, IMVs were solubilized with 2% n-dodecyl β-D-maltoside for 30 min in the presence of 1 Complete Protease inhibitor tablet (Roche). The solubilized membranes were incubated with Ni²⁺-NTA beads (Qiagen) for 2 h and transferred to a Bio-spin micro column (Bio-Rad). The column was washed with 6 column volumes of a washing buffer containing 50 mM phosphate buffer, pH 7, 100 mM KCl, 0.1% n-dodecyl β-D-maltoside, 10 mM imidazole, and 20% glycerol. The Ni²⁺-NTA-bound SecYEG was labeled with 600 μM of either Atto647N, Alexa 488 or Cy5 for 2 h at 4 °C. The labeling procedure was performed at pH 7 to ensure a higher labeling specificity and efficiency. Free dye was removed by extensive washing with washing buffer. SecYEG was eluted with 300 mM imidazole. The purity and concentration of SecYEG and the fluorophores were estimated by SDS-PAGE and spectrophotometrically. The extinction coefficient used for SecYEG at 280 nm was 71,000 M⁻¹ cm⁻¹. The extinction coefficients for the fluorophores were used as provided by the manufacturers.

The expression clone to produce the scaffold protein ApoE422k representing an N-terminal 22-kDa fragment of the human apolipoprotein E4, harboring a His6 and thioredoxin tag was kindly provided by Prof. James Rothman (Yale University, New Haven, CT). ApoE422k was produced and purified as described (19). MSP1E3D1 was kindly provided by Prof. Stephan Sligar (University of Illinois, Urbana, IL) and produced and purified as previously reported (36).

Reconstitution of SecYEG into Proteoliposomes—A lipid mixture containing DOPC:DOPG:DOPE (molar ratio 40:30:30) or DOPC:DOP (molar ratio 40:60) (Avanti Biochemicals, Birmingham, AL) (100 μg; 4 mg/ml) was solubilized with 0.5% Triton X-100 and mixed with 2.5 nmol of purified SecYEG. Reconstitution was performed as described before (37).

NanoDisc Reconstitution of SecYEG—For nanodisc formation, a lipid mixture containing DOPC:DOPG:DOPE (molar ratio 40:30:30), DOPC:DOP (molar ratio 40:60), or E. coli phospholipids (Avanti Biochemicals) was dried in a vacuum evaporator. Remaining traces of chloroform were removed by further drying of the lipid film in a desiccator overnight. Lipids were resuspended in a buffer containing 23 mM sodium cholate,
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25 mM HEPES/KOH, pH 7.4, 140 mM KCl, 0.17 mM DTT. For small nanodisc formation, SecYEG, MSP1E3D1, and lipids were mixed in a molar ratio of 1:10:250. Large nanodiscs were produced by mixing SecYEG, ApoE422k, and lipids at a molar ratio of 0.25:10:1800. The reconstitution mixtures were incubated at 4 °C for 1 h. Detergent was removed using Bio-Beads SM2 sorbent (Bio-Rad) in an overnight step. Minor amounts of formed proteoliposomes were removed by a centrifugation at 250,000 × g for 30 min. Nanodiscs were subjected to size exclusion chromatography by fast protein liquid chromatography using a Superose 6 column (GE Healthcare), and 0.5-ml fractions were collected in 50 mM HEPES/KOH (pH 7.4), 100 mM KCl, and 5% glycerol. Nanodisc containing fractions were analyzed by SDS-PAGE.

In Vitro proOmpA Translocation Assay—The activities of wild type and mutated SecA were analyzed by a standard proOmpA translocation and protease protection assay (38).

ATPase Activity Assay—ATPase activity assay of SecA was performed with minor modifications as described below (7, 39). A reaction mixture containing 25 mM HEPES/KOH, pH 7.4, 25 mM KCl, 5 mM MgCl₂, 0.1 mg/ml BSA, 2 mM DTT, 0.04 mg/ml SecB, 5 μM SecYEG proteoliposomes, and 50 or 500 nM wild type or mutated SecA was prepared. To measure the proOmpA-dependent SecA activity, proOmpA was added to a final concentration of 0.04 mg/ml. The basal SecA activity was measured by adding an 8 μM urea buffer instead of proOmpA. Reactions were initiated by addition of 5 mM ATP and performed at 37 °C for 30 min. Following, samples were diluted to reach an ATP concentrations below 0.25 mM. Free phosphate was quantified using a Malachite Green Phosphate assay kit (Gentaur).

Atomic Force Microscopy—Nanodiscs were diluted in a buffer containing 50 mM HEPES/KOH, pH 7.4, 50 mM KCl, 5% glycerol, and 100 mM MgCl₂ to a final concentration of ~1 nm. By incubation of this solution for 10 min with freshly cleaved mica, the nanodiscs were immobilized on the surface. Nonimmobilized material was rinsed off. AFM images were recorded in tapping mode in ScanAsyst-fluid regime by a Multimode 8 instrument, Controller V (Bruker). Images were taken using SNL-A silicon probes with a reflective Au coating on the back side and a tip radius of 2 nm. All images were obtained in buffer at room temperature using a spring constant of 0.35 N/m. A 2 kHz tapping frequency was used with a scan size of 3 μm, a scan speed of 0.2 Hz, and a 1024 lines/sample resolution capability. Analysis of height and diameter of the recorded images was performed manually using NanoScopeAnalysis 1.2 software.

Microscale Thermophoresis—Microscale thermophoresis experiments were performed using a Monolith NT.115 from Nanotemper Technologies (Munich, Germany) to test the binding of SecA to the SecYEG containing nanodiscs. A serial dilution of unlabeled SecA or a SecA mutant was prepared using a buffer containing 50 mM HEPES/KOH, pH 7.4, 50 mM KCl, 5% glycerol, and 0.5 mg/ml BSA. Cy 5-labeled SecYEG reconstituted in either small or large nanodiscs, and ATP was added to a final concentration of 50 nM and 5 μM, respectively. The samples were loaded into Monolith NT.115 series MST premium coated capillaries, and MST measurements were performed using 80% LED power and 80% IR-laser power. The data were fitted using the Hill equation.

FRET Measurements—FRET assays to examine protein translocation into SecYEG reconstituted nanodiscs were performed using SLM2 spectrofluorometer (Aminco Bowmann), as described previously (16). Briefly, proOmpA-DHFR was labeled with Cy3-maleimide (donor) (λ_ex = 550, λ_em = 570), and the DHFR domain was folded in the presence of methotrexate and NADPH. SecYC148EG was labeled with Atto647N (acceptor) (λ_ex = 650, λ_em = 670). FRET-based real time translocation of 200 nM prefolded proOmpA-DHFR-Cy3 was performed in the presence of 200 nM SecYC148EG-Atto647N reconstituted small or large nanodiscs, 50 mM HEPES/KOH, pH 7.4, 30 mM KCl, 5 mM MgCl₂, and 10 mM DTT. Translocation was initiated with 5 μM ATP, whereby the donor fluorophore was excited at 525 nm, and FRET efficiency was measured as an increase in acceptor fluorescence at 670 nm.

Fluorescence Cross-correlation Spectroscopy—FCCS experiments were performed on a dual color laser scanning LSM710 inverted confocal microscope (Zeiss). A helium–neon laser at 488 nm and an argon laser at 633 nm were used to excite the fluorophore-conjugated SecYC148EG. Fluorescence was recorded in blue (505–570 nm) and red (640–700 nm) channels. SecYC148EG was labeled with Atto647N and AF488 simultaneously (17, 18). The labeling efficiency for each fluorophore reached ~55%, resulting in a 110% overall labeling efficiency. This slight overlabeling suggests a small amount of unspecific labeling. Fluorescently labeled SecYC148EG was reconstituted into large nanodiscs. Autocorrelation in fluorescence for both fluorophores was recorded and analyzed as described before (40).

Author Contributions—S. K. designed and performed most of the experiments, analyzed the results, and wrote the paper. J. D. W. conducted the cloning and in vivo activity assays. I. V. and J. P. B. designed and conducted experiments that provided the basis of the work. P. G. performed atomic force microscopy experiments and was supervised by A. H. A. J. M. D. and A. M. V. O. conceived the idea for the project, designed the experiments, supervised the work, and wrote the paper. All authors contributed to the editing of the manuscript and approved the final version.

Acknowledgments—We thank M. Exterkate, I. Kusters, and A. B. Seinen for technical support and many valuable comments and discussions on the project.

References
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