Unlocking the Bacterial SecY Translocon

Highlights

- Validation of previously observed signal sequence-induced "unlocking" of SecYEG
- Conformational changes upon SecYEG unlocking are relayed to SecA binding site
- Unlocking the translocon perturbs the interaction between SecY and SecE
- Conformational changes distinct between secretion and membrane protein insertion

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In Brief

Corey et al. have used biochemical and computational techniques to analyze conformational changes in the bacterial SecYEG translocon upon signal sequence binding. These structural effects are likely key to understanding how the translocon is primed for translocation by the concerted action of the signal sequence and the ATPase SecA.

Corey et al., 2016, Structure 24, 1–10

April 5, 2016 ©2016 The Authors

http://dx.doi.org/10.1016/j.str.2016.02.001
Unlocking the Bacterial SecY Translocon

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http://dx.doi.org/10.1016/j.str.2016.02.001
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SUMMARY

The Sec translocon performs protein secretion and membrane protein insertion at the plasma membrane of bacteria and archaea (SecYEG/β), and the endoplasmic reticular membrane of eukaryotes (Sec61). Despite numerous structures of the complex, the mechanism underlying translocation of pre-proteins, driven by the ATPase SecA in bacteria, remains unresolved. Here we present a series of biochemical and computational analyses exploring the consequences of signal sequence binding to SecYEG. The data demonstrate that a signal sequence-induced movement of transmembrane helix 7 unlocks the translocon and that this conformational change is communicated to the cytoplasmic faces of SecY and SecE, involved in SecA binding. Our findings progress the current understanding of the dynamic action of the translocon during the translocation initiation process. The results suggest that the converging effects of the signal sequence and SecA at the cytoplasmic face of SecYEG are decisive for the intercalation and translocation of pre-protein through the SecY channel.

INTRODUCTION

An essential part of the biogenesis of many proteins is the crossing or insertion into a lipid membrane, a process known as protein translocation. The bulk of translocation occurs at the ubiquitous and essential Sec translocon: Sec61 in the eukaryotic ER membrane; SecYEG in the plasma membrane of bacteria, archaea (SecYEβ), and the thylakoid membrane of chloroplasts (Collinson et al., 2015). In bacteria, SecYEG associates with the cytoplasmic ATPase SecA (forming the transloconease) to carry out post-translational translocation/secreton of proteins across the membrane (Brundage et al., 1990), in a process driven by ATP synthesis and enhanced by the transmembrane proton motive force (PMF). Conversely, membrane proteins are inserted into the bilayer upon direct interaction of SecYEG with the ribosome nascent chain complex (Beck et al., 2000). Auxiliary proteins including SecD, SecF, and YajC assist in the secretion process (Duong and Wickner, 1997), while YidC cooperates in the insertion of transmembrane helices (TMs) of membrane proteins (Scotti et al., 2000); together with the SecYEG core-translocon they form a holotranslocon complex (Samuelson et al., 2000; Schulze et al., 2014).

The resting state structure of the Methanococcus jannaschii translocon (SecYEβ) reveals a channel through the center of the complex, formed between the two halves of SecY: TMs 1–5 and 6–10 (Figure 1A, top left panel; van den Berg et al., 2004). In the resting state, this channel is blocked at the center by a ring of conserved hydrophobic residues and a short “plug” helix (Figure 1A, red helix). SecY is braced at the back by SecE, with one TM tilted across the membrane and an amphipathic helix flat along the cytoplasmic face of the membrane (Figure 1A, beige helices). For channel opening, SecE presumably relaxes and allows separation of the two halves of SecY along with the displacement of the plug (Tam et al., 2005; van den Berg et al., 2004; Flower et al., 1995). This opening creates a pathway for proteins through the middle of the channel for secretion, as well as sideways into the bilayer via a “lateral gate” (LG) for membrane protein insertion (van den Berg et al., 2004).

In bacteria, secretory proteins with a cleavable N-terminal signal sequence (SS) engage SecYEG and SecA at the plasma membrane, with the secretory protein still in an unfolded state (Arkowitz et al., 1993; Hartl et al., 1990). Once engaged, the SS is inserted into the membrane at the interface between the bilayer and the LG of SecY (Gold et al., 2013; Briggs et al., 1986; Mcknight et al., 1991; Plath et al., 1998); we have previously proposed that this interaction unlocks the translocon prior to channel opening through a series of conformational changes within SecY (Hizlan et al., 2012). Notably, TM 7 straightens by ~40° toward the center of the channel to contact TM 10 (Figure 1A, right panels, turquoise helix). This in turn displaces the plug (Figure 1A, right panels, red helix), and likely primes the channel for translocation.

Mutations of SecY have been identified that allow in vivo transport of pre-proteins with a defective SS (prlA mutants) (Smith et al., 2005; Emr et al., 1981; Derman et al., 1993). These mutants also exhibit increased translocation activity and are not further stimulated by the PMF (Nouwen et al., 1996). They possibly act by stabilizing the unlocked state of SecY, otherwise induced by SS binding; consistent with this, most prlA mutations map to TMs 7, 10, and the plug of SecY (van den Berg et al., 2004; Osborne and Silhavy, 1993). For example, the potent prlA4 mutation (SecY F286,I408N in Escherichia coli; Figure 1B, red circles) (Emr et al., 1981; Duong and Wickner, 1999) might promote the relocation of TM 7, thereby unlocking the complex in the absence of a functional SS.
A structure of Thermotoga maritima SecYEG bound and activated by SecA reveals further substantial changes in the SecY channel (Zimmer et al., 2008). Upon SecA binding, the two halves of SecY separate—characterized by a widening of the SecY LG— and TMs 8 and 9 move outward, producing a shift in the SecE amphipathic helix (Zimmer et al., 2008). SecA binds SecY at two cytoplasmic loops (C4 and C5; Figure 1B). The latter includes the highly conserved “RPG” site (SecYR357,P358,G359 in *E. coli*), mutations of which perturb SecA interaction (Alami et al., 2007; Tam et al., 2005; de Keyzer et al., 2007) and thereby abolish translocation (Mori and Ito, 2001).

It is not yet known how the SS and SecA-driven conformational changes prime SecYEG for transport, nor is it clear how they compare with those required for membrane protein insertion. Here we present a series of biochemical and biophysical analyses, supported by all-atom molecular dynamics (MD) simulations, aimed at investigating the mobility of key regions of the SecY protein channel during the initiation process. The data verify the physiological significance of the SS-induced displacement of TM 7, visualized in the absence of SecA (Hizlan et al., 2012), and provide further insights into the dynamic action of the SecY translocon during the early stages of protein secretion.

### RESULTS

#### Production of Different States of SecYEG

A number of variants of *E. coli* SecYEG were produced (Table 1) to characterize the unlocking process: (1) the hyperactive SecYprlA4EG variant (SecYF286Y,I408N; red circles in Figure 1B), which allows translocation of substrates with a defective SS (Emr et al., 1981; Duong and Wickner, 1999); (2) the secretion-defective RPG variant, which perturbs SecA binding (SecYR357E,P358D,G359P; hereafter referred to as SecYEDPEG; “RPG” in Figure 1B) (Mori and Ito, 2001; Tam et al., 2005); and (3) a hitherto uncharacterized combination of the two (SecYprlA4-EDPEG).

To trap the complex in the reported unlocked state (Hizlan et al., 2012), we engineered a unique cysteine pair into an otherwise cysteine-free mutant of SecYEG to crosslink TMs 7 and 10 of SecY (SecYR357E,P358D,G359P; hereafter called SecYEDPEG; Figures 1A and 1B, orange and green circles; Figure S1A, sticks). Membranes from *E. coli* cells expressing this construct were either reduced (1 mM tris(2-carboxyethyl)phosphine [TCEP]) or oxidized (1 mM Cu-phenanthroline) prior to purification of SecYEG in detergent. The concentration of free thiols in the purified protein demonstrates that the oxidized form is >85% crosslinked in detergent solution (Figure S1B).

The anomalous behavior of SecY during SDS-PAGE is shared by many other membrane proteins and probably arises due to its partially folded state in SDS and, hence, faster migration through the gel and significantly reduced apparent molecular weight.

### Table 1. Description of the SecYEG Variants Employed in the Study

<table>
<thead>
<tr>
<th>SecYEG Variant</th>
<th>Substitutions</th>
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<tbody>
<tr>
<td>SecYprlA4EG</td>
<td>SecYF286Y,I408N</td>
</tr>
<tr>
<td>SecYEDPEG</td>
<td>SecYR357E,P358D,G359P</td>
</tr>
<tr>
<td>SecYprlA4-EDPEG</td>
<td>SecYF286Y,I408N,R357E,P358D,G359P</td>
</tr>
<tr>
<td>SecYprlA4EG</td>
<td>SecYF286Y,I408N</td>
</tr>
</tbody>
</table>

The name of the variant is shown in the left column and the corresponding point mutations in the right column.
Interestingly, SecY prlA4 has a slight, but consistent, increase in apparent MW (34 kDa; Figure S2 A); this presumably correlates with a less compact, or more flexible, state. Note that oxidized SecY7–10 also has reduced mobility, albeit less pronounced (Figure S2 A; 33 kDa), suggesting that the loosening observed in SecY prlA4 can be partially achieved by displacing TM 7 with a crosslink to TM 10.

Similar effects were observed on the whole SecYEG complex following analysis by size-exclusion chromatography (Figure S2B). Evidently, the loosening of SecYEG brought about by SecY prlA4, and to a lesser extent SecY7–10 (and hence the movement of TM 7), may be correlated with the unlocking and activation process.

The Inactivation Caused by C5 Loop Variants Are Reversed by Substitutions inside the SecY Channel

Exposure of SecYEG to trypsin results in proteolysis at the cytoplasmic face of SecY, with cleavage occurring predominantly on the C4 loop (Figure 1B, scissors). Using low and high concentrations of trypsin (Figure 2 A + and ++), standard SecY is either partially or completely cleaved, respectively (Figure 2 A, lanes 1–3). Trypsin sensitivity is largely unaffected by the SecY prlA4 substitutions (Figure 2 A, PrlA4 in lanes 4–6), but considerably reduced with the SecY EDPEG substitutions, which are on the adjacent loop (C5) (Figure 2 A, EDP in lanes 7–9). The reduced trypsin sensitivity suggests that SecY EDPEG brings about a structure that is not cleaved by trypsin.

Figure 2. Competing Effects of the Activating Variant prlA4 and Inactivating Variant SecY EDPEG

(A) SecYEG, SecY prlA4EG (PrlA4), SecY EDPEG (EDP), and SecY prlA4-EDPEG (PrlA4-EDP) were subjected to trypsin digestion followed by Coomassie-stained SDS-PAGE to examine the effect of different amino acid substitutions on the conformation of the C4 loop of SecY (the site of a prominent tryptic cleavage site). Samples are either untreated (−) or treated with 0.075 μg/ml (+) or 0.75 μg/ml (+++) trypsin. The major bands correspond to full-length SecY, the N-terminal cleavage fragment, the C-terminal fragment, SecE, and a lower band for both SecE breakdown products and secondary SecY cleavage products (asterisk).

(B) The rate of ATP turnover (k cat) is shown for SecA bound to different variants of SecYEG reconstituted into PL (purple), and following addition of a translocation substrate pOA (blue). ATP turnover is diminished by the C5 substitutions (SecY EDPEG), but is then restored when combined with the prlA4 mutations. Error bars denote SEM of three repeats.

(C) Translocation of pOA into PL reconstituted with different SecYEG variants. Translocation efficiencies were determined by western blotting against pOA following a protease K protection assay (transported pOA is in the PL interior and protected from proteolysis); for a representative blot see Figure S3A. Results were quantified against a non-protease K-treated control and normalized to standard SecYEG, shown with SEM of three repeats. As in (B), the abolished activity of SecY EDPEG (EDP) is rescued by the prlA4 mutations.

(D) Affinities of SecYEG for SecA in detergent solution, determined through quenching of a fluorescent marker on SecA (SecA*). The decrease in fluorescence was plotted and fitted to tight binding equations (raw data are shown in Figure S3B). The calculated K D values are shown with SEM of three runs. **p = 0.004.

(E) Representative structures of the cytoplasmic face of SecY (cartoon representation) following MD simulations of SecYEG, SecY EDPEG, and SecY prlA4-EDPEG. Structures are viewed from the side, with the cytosolic face indicated by the black line. The C4 loops are shown in light blue, the C5 loops green, and the rest of SecY white. In each panel, the position of the Cα of the conserved arginine in the RPG motif (mutated to glutamic acid in the middle and right-hand panels) is shown as a green sphere and labeled “R” or “E” accordingly. The approximate position of the tryptic cleavage sites is indicated by a pair of scissors. These structural changes are quantified in Figure S3D.
broad conformational change affecting both loops C5 (containing the RPG motif) and C4 (containing the trypsin site). This effect is reversed when SecYEDPEG is combined with the prlA4 substitutions (SecYprlA4-EDPEG) (Figure 2A). The activating prlA4 mutations (Figures 2B and 2C, SecY prlA4-EDPEG) were used to reconstitute monitor ATP-driven protein translocation. The results show that, as expected (Mori and Ito, 2001), SecY EDPEG displays a dramatic loss in both SecA activation and translocation activity (Figures 2B, 2C and 3A). Interestingly, as with trypsin sensitivity, this effect is partially ameliorated by the prlA4 mutations (Figures 2B and 2C, SecY prlA4-EDPEG). Thus, the effects of prlA4 are dominant to the inactivating SecY EDPEG with respect to transport activity.

Next, the affinity of SecYEG for SecA was measured using a fluorescent probe on SecA (hereafter SecA*) which, when bound by a non-hydrolyzable analog of ATP (AMP-PNP), is quenched upon SecY binding (Deville et al., 2011)(Figure S3B). SecA* binds tightly to standard SecYEG (K_D ~12 nM; Figures 2D and S3B), with the affinity slightly lower for SecY EDPEG (K_D ~36 nM; Figure 2D). This is perhaps surprising, as previous results have indicated a higher affinity for SecY prlA4EG (de Keyzer et al., 2002; van der Wolk et al., 1998). A possible explanation for this disparity is the use in the previous study of crude inner membrane vesicles (IMVs) rather than purified components; experiments conducted in IMVs may be influenced by auxiliary translocon components (e.g., SecDF) or unknown effector proteins.

The affinity of SecA for SecY EDPEG is more than 10-fold weaker than for standard SecYEG (K_D ~135 nM; Figure 2D); this is unsurprising, as the RPG site is located on the SecY-SecA interface (Figure S3C). Once again, however, this effect is reversed when the SecY EDPEG variant is paired with the prlA4 mutations (K_D ~42 nM; Figure 2D).

The RPG Substitutions Perturb the Adjacent C4 Loop

To investigate the structural changes associated with the SecY EDPEG substitutions, we ran all-atom MD simulations using the M. jannaschii SecYEβ crystal structure as starting coordinates (PDB: 1RHZ; van den Berg et al., 2004). Three simulations each of SecYEβ and the corresponding variants SecY prlA4Eβ, SecY EDPEGβ, and SecY prlA4-EDPEGβ were run for 300–400 ns to provide suitable sampling.

It is apparent from the MD data that the SecY EDPEGβ substitutions on loop C5 have little effect on the conformation of this loop (Figures 2E and S3D). Interestingly, they do produce considerable secondary structural change on the adjacent C4 loop—which contains the primary trypsin cleavage site—with a marked increase in β-sheet composition (Figures 2E and S3D, right panel). SecY prlA4Eβ and SecY prlA4-EDPEGEβ are largely unchanged when compared with the native SecYEβ. Therefore, the MD results are consistent with the trypsin sensitivity data (Figure 2A), showing that the SecY EDPEGβ substitutions on the C5 loop are transmitted to the adjacent C4 loop, and that these effects are reversed by the distant prlA4 mutations.

The activating prlA4 mutations are consistently dominant over the inactivating SecY EDPEG, demonstrating that the impaired function of SecY EDPEG is not simply a loss of a direct contact site for SecA. The data here reveal a functional complementation between regions deep within the protein channel (prlA4 sites; TM 7 and TM 10) and the cytosolic surface, typical of linked conformational changes over long range.

Straightening of TM 7 toward TM 10 Activates SecYEG

The SecY7–10EG variant (Figures 1 and S1A) comprises an engineered cysteine pair between TMs 7 and 10 with the two native cysteines in SecY replaced with serine (which has no effect on the function of SecYEG; Kaufmann et al., 1999). This strategically placed thiol pair was designed to trap the translocon in the un-locked state, as seen with SS binding (Hizlan et al., 2012), and is thought also to be favored by SecY prlA4EG.

SecY prlA4EG has an enhanced translocation activity but is unaffected by the PMF (Nouwen et al., 1996), so SecY7–10EG should exhibit similar properties. To this end, we carried out translocation assays on each variant in PL with and without PMF stimulation, achieved by co-reconstituting SecYEG with the light-driven proton pump bacteriorhodopsin (bR; Figure S4), as described previously (Schulze et al., 2014). As expected, SecY prlA4EG is activated with respect to translocation compared with the standard form of SecYEG (Figure 3A, “–Light” lanes), and this activity is not further enhanced by the PMF (Figure 3A, “+Light” lanes). Unexpectedly, the cysteine pair of SecY7–10EG alone has the same effect on translocation activity as SecY prlA4EG, whether oxidized or reduced. However, when TM 7 is permanently fixed to TM 10 by a disulfide bond the channel is found in the activated state (as it is with the SecY prlA4EG and SecY7–10EG reduced). The fact that the disulfide crosslink retains this activated state suggests that the proximity of TMs 7 and 10 is a feature of this activation process (although it can also be achieved without the permanent cross-link). Therefore, the movement of TM 7 toward TM 10 activates the SecYEG complex, presumably in the manner proposed for SecY prlA4EG, i.e., through the displacement of TM 7.

The Activation Pathways for Protein Secretion and Membrane Insertion Are Distinct

Experiments were conducted to establish whether the activation and unlocking process described for protein secretion also holds true for membrane protein insertion. The perturbation of TM 7 in SecY prlA4EG or SecY7–10EG did not result in increased insertion activity of a model substrate, F_O(a) (Figure 3B). Therefore, for this substrate the activation pathways for membrane protein insertion are distinct to those seen for protein secretion, at least with respect to the mobility of TM 7.

The Movement within SecY of TM 7 toward TM 10 Is Relayed to the Amphipathic Helix of SecE

SecYEG activation has previously been linked to destabilization of the complex (Duong and Wickner, 1999), potentially by promoting a separation of the two halves of SecY. We reasoned that this effect could also be induced by low levels of a
denaturant, such as SDS. To monitor the effect of denaturation, we exploited the sensitivity of intrinsic tryptophan fluorescence to conformational changes within proteins: increasing concentrations of SDS were titrated into SecYEG, and tryptophan emission spectra measured.

The general effect of SDS addition is a red shift in emission, probably due to increasing solvent exposure of tryptophan residues as the complex unfolds (Figure S5A). This shift can be expressed as a ratio of the amplitude of the emission at 330 nm and 350 nm ($A_{330}/A_{350}$) and plotted against increasing concentrations of SDS (Figure 4A). The profiles show that the reduced SecYEG and the reduced Sec$Y_{7-10}$EG (Figure 4A, blue and green traces, respectively) reveal a distinct transition at $\sim0.01\%$ SDS. The unfolded forms of the complex, i.e., SecYE$\text{prlA4}$EG and oxidized Sec$Y_{7-10}$EG (Figure 4A, red and orange traces, respectively), exhibit a significant reduction in $A_{330}/A_{350}$ without SDS, and a diminished transition at $\sim0.01\%$ SDS. This effect is also observed when substituting SDS for a $\beta$-octyl glucoside (Figure S5B), for which SecYEG destabilization and dissociation has previously been characterized (Duong and Wickner, 1999). Therefore, the data indicate that the activated complexes are already in a partly destabilized form.

*E. coli* SecYEG has eight native tryptophan residues (four in SecY, three in SecE, and one in SecG; see Figure 4B). To localize the conformational change seen in Figure 4A, we created variants in which only one native tryptophan was retained, with the others substituted with phenylalanine, the product of which was still fully functional (Figure S5C). Each variant was individually examined and the observed transition was located to W84 (Figure 4C, blue line), positioned on the amphipathic helix of SecE (Figure 5A, W84). In a $\text{prlA4}$ background no transition is apparent: W84 already appears to be exposed to solvent even in the absence of denaturant (Figure 4C, pink line). When W84 was substituted with phenylalanine but the other native tryptophan residues retained, as expected the amplitude of the denaturation curve was significantly diminished (Figure 4D, black line). This effect mirrors SecYE$\text{prlA4}$EG retaining all eight native tryptophan residues, whereas W84 is unresponsive to denaturant. While the remaining seven tryptophan residues play a small role in this transition, the W84 effect is clearly dominant.

It is evident from the data that the decreased transition of SecY$_{7-10}$EG, caused by the ready exposure of W84, is also apparent when TM 7 is crosslinked to TM 10 (Figure 4A). Therefore, the movement of TM 7 to TM 10 brings about a conformational change involving the amphipathic helix of SecE (containing W84), which is also apparent upon complex destabilization.

Mobility of the Amphipathic Helix of SecE in Response to the Perturbation of TM 7

To help understand the conformational effects arising from TM 7 perturbation, we ran MD simulations, as above, using *M. jannaschii* SecYE$\text{prlA4}$ (PDB: 1RHZ; van den Berg et al., 2004), and the corresponding variants SecYE$\text{prlA4}$E$\text{b}$, oxidized SecY$_{7-10}$E$\text{b}$, and reduced SecY$_{7-10}$E$\text{b}$.

Post-simulation structural alignment of the different complexes reveals that the region with highest variability is the amphipathic helix of SecE (Figure 5A), near the position of SecYE$_{\text{prlA4}}$G (Figure 5A, cyan asterisk). The helices in the unchanged SecYE$\text{b}$ simulations display an upward tilt (i.e., toward the periplasm) compared with the input structure, whereas the SecY$_{\text{prlA4}}$E$\text{b}$ simulations display a downward tilt toward the cytoplasm (Figure 5A, blue and red, respectively). When TM 7 is fixed by crosslinking to TM 10, the position of the amphipathic helix resembles that of SecY$_{\text{prlA4}}$E$\text{b}$ (Figure 5A, orange), whereas with no crosslink the helix location resembles the standard SecYE$\text{b}$ form (Figure 5A, green). Note that the SecYE$\text{b}$ and SecY$_{\text{prlA4}}$E$\text{b}$ simulations in Figure 5A are the same as those presented in Figure 2E.

Distance analysis between the N-terminal tip of the SecE amphipathic helix and a rigid region of SecY (Figure 5A, yellow box) confirms that the effect is consistent across all three repeats, or in four of five repeats in the case of oxidized SecY$_{7-10}$EG (Figure 5B), and is apparent even in the early (20 ns) stages of the simulations (Figure S6A). Of the five repeats for the oxidized SecY$_{7-10}$EG trajectories (with the relocated TM7), one behaves like the reduced states (marked with a light orange arrow); thus, the analysis predicts that the position of the amphipathic helix of SecE has a strong, but not absolute,
dependence on the location of TM 7. To ensure this was not an archaea-specific artifact, we also ran simulations using SecYE from the Gram-negative Thermus thermophilus (PDB: 2ZJS; Tsukazaki et al., 2008). For simulations of standard SecYE and a SecYprlA4E equivalent, the SecE amphipathic helix behaves in a remarkably similar way to the corresponding feature of M. jannaschii (Figure S6B). Indeed, the SecE region most perturbed in these simulations is the region containing the SecYEW84G equivalent.

Note that, with respect to the SecE amphipathic helix, the MD simulations all diverge from the input structure (Figure 5B, black dashed line), despite having generally low root-mean-square deviations (Figure S7). This is possibly due to the presence of lipid bilayer in the MD simulations, absent in the crystal structure.

The simulation data are consistent with the tryptophan fluorescence analysis that localizes the primary effect of TM 7 movement to W84—on the amphipathic helix of SecE. Together, the results suggest that the change in conformation of TM 7 at the center of the channel, normally induced by SS binding, is coupled to the cytosolic surface of SecE. These SecE conformational changes differ from those reported in the SecA-bound SecYEG crystal structure (PDB: 3DIN, Figure 5C; Zimmer et al., 2008), suggesting a distinct role for the SecE amphipathic helix during SS-induced unlocking and SecA-mediated activation of the translocon.

DISCUSSION

This study explores the dynamic action of SecYEG during the initial stages of protein translocation, and builds on previous structural data of the E. coli SecYEG complex bound by a preprotein mimic (Hizlan et al., 2012). In this previous study, we proposed that the association of the SS with SecYEG causes a conformational change involving TM 7, TM 10, and the plug of the membrane bilayer (Gold et al., 2010; Lill et al., 1990; Robson et al., 2009) are sufficient to tip the balance toward the unlocked state, whereas in the absence of a biological membrane (i.e., in detergent solution) the translocon in this unlocked state with use of an engineered disulfide crosslink between TMs 7 and 10 of SecY (SecY7–10EG; W84) and all other tryptophans changed to phenylalanine, in either standard SecYEG or prlA4 background. Error bars are the SEM of three repeats.

Curiously, when SecYEG is in a membrane environment, the engineered cysteines in TMs 7 and 10 alone (uncrosslinked) stimulate protein translocation, similar to SecYprlA4EG and the crosslinked variant SecY7–10EG (Figure 4A). However, when in a non-membrane environment they fail to replicate the unlocked state characteristic of the activated SecYprlA4EG (Figure 4A). Most likely, TM 7 is delicately poised between the two states, and for the uncrosslinked variant the known activating properties of the membrane bilayer (Gold et al., 2010; Lill et al., 1990; Robson et al., 2009) are sufficient to tip the balance toward the unlocked state, whereas in the absence of a biological membrane (i.e., in detergent solution) the crosslink is required for activation. This is not without precedent, as clear conformational differences have been observed in the transmembrane region of SecYEG in structures of the complex determined in detergent compared with the native lipid environments (Bostina et al., 2005).

Our analyses indicate that the unlocking mechanism, involving TM 7 at the center of SecY, is transmitted to the cytosolic face of...
SecYEG, with the most prominent structural change in the SecE amphipathic helix (Figures 4 and 5), which is well conserved between archaea and bacteria (Enno et al., 1994). In addition, through analysis of SecYEG with and without prlA4, it is clear that channel unlocking leads to a restructuring of the cytoplasmic loops of SecY (Figure 2). These loops are universally important in the interaction of SecY with translocation partners, such as the ribosome or SecA, with the R357 position completely conserved across all three domains of life. It is conceivable that the region containing the RPG motif acts as a conserved coupling switch for the activation of the translocon, either by SecA or the ribosome, or even analogous translocation partners in organisms lacking SecA. Indeed, the functional complementation of bacterial SecY by the archaeal counterpart (Auer et al., 1991) supports the existence of a common mechanism for activation.

The reported conformational coupling between SS and SecA binding is consistent with the observed transactivation of SecA by the SS (Gouridis et al., 2009), which occurs via SecYEG (Hizlan et al., 2012). Indeed, structural studies have shown that both SS and SecA binding have major effects on SecYEG (Hizlan et al., 2012; Zimmer et al., 2008); in the LG, plug, and C4 and C5 loops of SecY, and the amphipathic helix of SecE (Collinson et al., 2015). The perturbation of the amphipathic helix of SecE shown here, induced by the SS via TM7, might release the SecE brace around SecY and promote channel opening, as originally proposed on the basis of the SecYEβ structure (van den Berg et al., 2004). This concerted action of SS and SecA presumably primes SecYEG for the introduction of pre-protein, and acts as a prelude to ATP- and PMF-driven translocation (Figure 6).

The events described here relate to SecA-dependent bacterial secretion. Post-translational secretion through the eukaryotic secretory pathway occurs via the mammalian Sec61 complex (Collinson et al., 2008), and SecY bound to SecA, T. maritima SecYEG-A (3DIN) (Zimmer et al., 2008) (SecA not shown). The amphipathic helix of SecE is shown in both structures, respectively in blue and red, and the equivalent key tryptophan residue of the E. coli SecY-W84 (W20 in T. maritima) in blue spheres. The recent structure of the mammalian Sec61 complex associated with the ribosome and nascent secretory protein (Voorhees and Hegde, 2016) is most likely captured beyond the initiation phase, showing the channel in the open state with widened LG and shifted TM 7, equivalent to that shown in Figure S6A. The distance for the input structure is highlighted by the black dotted line. In the right panel, the single oxidized SecY WT EG simulation, which resembles the reduced states, is marked with a light orange arrow.

Figure 5. Molecular Dynamics Simulations of the SecY Complex
(A) Post-simulation snapshots of the M. jannaschii SecYEβ variants with the whole complex shown in gray except the amphipathic helix of SecE, which is colored according to variant: standard SecYEβ in blue; PrlA4 in red; oxidized SecY WT EG in orange and reduced SecY WT EG in green. The immobile region of SecY used for distance analysis in (B) is highlighted with a yellow box. The equivalent position of W84 is marked with a cyan asterisk.
(B) Distance analysis between the SecE amphipathic helix and a rigid region of SecY (yellow box in A). The first 20 ns of simulation are shown in detail in Figure S6A. The distance for the input structure is shown by the black dotted line. In the right panel, the single oxidized SecY WT EG simulation, which resembles the reduced states, is marked with a light orange arrow.
(C) Crystal structures of the SecY complex alone, M. jannaschii SecYEβ (1RH2) (van den Berg et al., 2004), and SecY bound to SecA, T. maritima SecYEG-A (3DIN) (Zimmer et al., 2008) (SecA not shown). The amphipathic helix of SecE is shown in both structures, respectively in blue and red, and the equivalent key tryptophan residue of the E. coli SecY-W84 (W20 in T. maritima) is highlighted by cyan spheres.

Experimental Procedures

Chromatography media were purchased from GE Healthcare. Detergents were obtained from Glycon and lipids from Avanti Polar Lipids. NuPAGE gels were bought from Life Technologies. Unless stated, all other materials were supplied by Sigma-Aldrich.

Protein Production

Point mutations were introduced using the QuickChange protocol (Stratagene) and confirmed by sequencing. The SecYEG variants were expressed and purified by procedures developed for the standard complex (Collinson et al., 2001). In brief, SecYEG-pBAD was expressed in C43 E. coli cells, which were lysed and the membrane fraction isolated by centrifugation; the SecYEG protein was solubilized from the membrane fraction using 1% DDM (n-dodecyl β-D-maltoside) and then purified using nickel chromatography.
Limited proteolysis was performed by mixing SecYEG in detergent with porcine trypsin at 0.75 mg/ml or 0.075 mM or reduced with 1 mM TCEP for 1 hr at 4°C, before reverting to the standard protocol. In the case of the reduced samples, 0.5 mM TCEP was added to all buffers downstream of the reduction step.

Reconstitution into PL was carried out as previously described (Gold et al., 2016), using E. coli polar lipids and removing excess polar lipids and removing excess detergent with Bio-Beads (Bio-Rad) by dialysis. Note that the presence of the lipids does not alter the oxidation state of SecY7–10EG (Figure S4).

Size-Exclusion Chromatography

SecYEG samples or protein standards of equal concentrations were run at a constant flow of 0.5 ml/min through a Superose 6 10/300 column in 20 mM Tris (pH 8), 130 mM NaCl, 10% glycerol, and 0.02% DDM in a Hellma ultracentrifuge. SecA bound samples or protein standards of equal concentrations were run at a constant flow of 0.5 ml/min through a Superose 6 10/300 column in 20 mM Tris (pH 8), 130 mM NaCl, 10% glycerol, and 0.02% DDM, while monitoring the absorbance at 280 nm.

Biochemical Assays

Limited proteolysis was performed by mixing SecYEG in detergent with porcine trypsin at 0.75 μg/ml or 0.075 μg/ml, incubating at room temperature for 20 min, then mixing with lithium dodecyl sulphate gel loading buffer to quench the reaction. Results were analyzed with SDS-PAGE.

The concentration of free thiols in purified SecYEG samples was measured by assaying for binding to a thiol-reactive fluorescence dye, CPM (7-diethylamino-3-(4’-maleimidophenyl)-4-methylcoumarin). Assays were performed using 1 μM SecYEG in detergent incubated with 10 μM CPM. CPM was excited at 389 nm and emission scans were read at 420–520 nm on a Jobin Yvon Fluorolog (Horiba Scientific).

In vitro ATPase and translocation assays involving the model substrate proOmpA (pOAp) were performed essentially as described previously (Gold et al., 2007; Robson et al., 2009). In brief, the ATP turnover rate of 0.3 mM SecA was determined alone, with 2.4 μM SecYEG in PL and in the presence of 0.7 μM pOAp. Reactions were followed by coupling ATP hydrolysis to NAD+ reduction via pyruvate kinase and lactate dehydrogenase, using absorbance at 340 nm. After 30 min, untranslocated material was digested with protease K and translocation yields were determined by western blotting for 20 min, then mixing with lithium dodecyl sulphate gel loading buffer to translocate material.

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were excited at 288 nm, and an emission scan was recorded between 320 and 370 nm. SDS or j-ocetyl glucoside was titrated into each sample at a concentra-
tion range of 0.01%–3%. Three separate emission scans were run for each
concentration, and ratios of the fluorescence at 330 and 350 nm were plotted as a function of SDS or j-ocetyl glucoside concentration.

Molecular Dynamics Simulations

Models for the simulations were built using the crystal structures M. jannaschii
SecYE (1RHZ; van den Berg et al., 2004) and T. thermophilus SecYE (2JJS; Tsukazaki et al., 2008). The missing loops in SecY were added using
MODELLER (Fiser et al., 2003) and amino acid substitutions were made using
SCWRL4 (Krivoy et al., 2009). Simulations were run in Gromacs 4.6.4 (Berendsen
et al., 1995) using the OPLS all-atom force field (Jorgensen et al., 1996). The pro-
tein was embedded in a 512 united-atom palmitoyl-oleoyl phosphatidylcholine
membrane (Ullschneider and Ullschneider, 2009) using g_memb (Wolf et al.,
2010). The protein-membrane structures were solvated with explicit (Sim-
ple Point Charge) water and sodium and chloride ions to a neutral charge and
concentration of 0.15 M. The systems were energy minimized using the steepest
descents method, and equilibrated for 1 ns using the NPT (constant temperature
and constant pressure) ensemble at 300 K with the Bussi-Donadio-Parrinello
thermostat and semi-isotropic Parrinello-Rahman pressure coupling. Bond
lengths were constrained using the LINCS method. Non-bonded interactions
were dealt with by the Verlet cutoff, and the neighbor search list was updated
every 20 steps. Long-range electrostatic interactions were calculated using the
particle mesh Ewald method, and a cutoff of 1.0 nm was applied for van
der Waals and short-range electrostatic interactions.

Secondary structure analyses of the loops involved residues 229–252
(C4) and 351–362 (C5) (M. jannaschii numbering) and used the DSSP program
(Joosten et al., 2011). Distance analyses were done using the Gromacs
utility g_dist.

Homology Modeling

For illustrative purposes only (Figure 4B), a model was built for E. coli SecYEG
using MODELLER (Sali and Blundell, 1993), with 1RHZ as a template (van den
Berg et al., 2004). This model was only used to demonstrate the positioning of
the tryptophan residues in the E. coli system and was not energy minimized or
subjected to MD simulation.

All molecular models are represented using PyMOL (Schrödinger).

SUPPLEMENTAL INFORMATION

Supplemental Information includes seven figures and can be found with this article online at http://dx.doi.org/10.1016/j.str.2016.02.001.

AUTHOR CONTRIBUTIONS

Experiments were designed by R.A.C., W.J.A., and I.C. Experiments
were performed by R.A.C., W.J.A., J.K., A.R., S.M., and I.C. Data were
analyzed by R.A.C., W.J.A., J.K., and I.C. The manuscript was written by

ACKNOWLEDGMENTS

This work was funded by the BBSRC: R.A.C. (BBSRC South West Bioscience
Doctoral Training Partnership), I.C. and W.J.A. (BB/I008675/1), and J.K.
(BBSRC doctoral training grant). We are very grateful to our dedicated tech-
nical support team Jeannette Hobbs, Kathleen Moreton, and Deborah Carter.
Thanks also to R. B. Sessions for continued guidance and advice. This work
was carried out using the computational facilities of the Advanced Computing
Research Center, University of Bristol (http://www.bris.ac.uk/acrc/). Additional
simulations were carried out using computer time on ARCHER provided by
HECBSim, the UK High End Computing Consortium for Biomolecular
Simulation (hecosim.ac.uk), supported by EPSRC.

Received: November 27, 2015
Revised: January 26, 2016
Accepted: February 9, 2016
Published: March 10, 2016

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