Copyright © 2008, American Society for Microbiology and/or the Listed Authors/Institutions. All Rights Reserved.

1	Maximal Efficiency of Coupling between ATP Hydrolysis and Translocation
2	of Polypeptides Mediated by SecB Requires Two Protomers of SecA
3	
4	Running title: Maximal Efficiency Requires Two Protomers of SecA
5	
6	Chunfeng Mao, ¹ Simon J. S. Hardy, ^{1,2} and Linda L. Randall ^{1,*}
7	¹ Department of Biochemistry, University of Missouri, Columbia, MO 65211, USA
8	² Department of Biology, University of York, York YO10 5DD, UK
9	*Correspondence: 117 Schweitzer Hall, University of Missouri, Columbia, MO 65211, USA;
10	<u>craneje@missouri.edu;</u> Tel: 573-884-4160; Fax: 573-882-5635
11	
12	ABSTRACT
13	SecA is the ATPase that provides energy for translocation of precursor polypeptides through the
14	SecYEG translocon in Escherichia coli during protein export. We have previously shown that
15	when SecA receives the precursor from SecB the ternary complex is fully active only when two
16	protomers of SecA are bound. Here we have used variants of SecA and of SecB that populate
17	complexes containing two protomers of SecA to different degrees to examine both the hydrolysis
18	of ATP and the translocation of polypeptides. We conclude that the low activity of the
19	complexes with only one protomer is the result of a low efficiency of coupling between ATP
20	hydrolysis and translocation.

INTRODUCTION

2 The general secretory or Sec system in *E. coli* translocates precursors of proteins across the 3 cytoplasmic membrane into the periplasmic space (for review see (21)). Some proteins, such as 4 alkaline phosphatase and the binding proteins for amino acids and sugars, function as soluble 5 proteins in the periplasm; others, such as OmpA, are inserted into the outer membrane. The path 6 across the cytoplasmic membrane barrier is provided by a translocation channel comprising a 7 heterotrimeric complex, SecY, SecE, and SecG (SecYEG). The energy for the movement is 8 supplied by protonmotive force and the hydrolysis of ATP by SecA, which is a peripheral 9 component of the membrane-associated translocon. The Sec system can export polypeptides only 10 if they are devoid of stable tertiary structure. SecB, a small cytosolic chaperone, acts to capture 11 precursors before they acquire stable structure and introduces them into the secretory pathway by 12 delivering them to SecA. SecA can also bind precursors directly as evidenced by the viability of 13 strains of *E. coli* that lack SecB. However, efficiency of export is drastically reduced for many 14 proteins (13).

15

16 Crystal structures of both SecB and SecA have been solved. SecB is a tetramer (monomer, 17 17 kDa) organized as a dimer of dimers (6, 33). There are six dimeric forms of SecA (2, 11, 20, 27, 18 30, 34) that differ greatly in the contacts that stabilize the dimeric interface. However, the 19 structures of the protomers are all closely related and display only two different conformations, 20 an open state (18) and a closed state (11). In free solution SecA undergoes an equilibrium 21 between monomer and dimer characterized by an equilibrium constant of 0.1 μ M to 1 μ M 22 depending on the ionic strength and temperature of the solution (31). When SecA interacts with 2 (24).

3

4	Here we have asked why two protomers of SecA are required. We have examined the
5	translocation into inverted cytoplasmic membrane vesicles of two natural ligands, one a soluble
6	periplasmic protein, galactose-binding protein, and the other an outer membrane protein, OmpA.
7	We conclude that the need for two protomers of SecA in the complex is to achieve maximal
8	coupling efficiency between ATP hydrolysis and translocation.
9	
10	MATERIALS AND METHODS
11	Materials
12	$[\gamma^{-32}P]$ ATP was purchased from either PerkinElmer (Boston, MA) or GE Healthcare (Pittsburgh,
13	PA), [³⁵ S]- methionine from PerkinElmer, pre-coated polyethyleneimine cellulose thin layer
14	chromatography plates from Merck KGaA (Darmstadt, Germany), trypsin from Millipore
15	Corporation (Freehold, NJ), bovine pancreatic trypsin inhibitor (aprotinin), and
16	deoxyribonucleaseI (DNaseI) from Sigma-Aldrich (St.Louis, MO), NAP TM 10 and HiTrap TM
17	Blue HP columns from GE Healthcare Bio-sciences AB (Uppsala, Sweden), N-[(2-
18	pyridyldithio)ethyl]-4-azidosalicylamide (AET) from Toronto Research Chemicals Inc.
19	(Canada), Precision Plus Protein TM Standards (all blue) from Bio-Rad (Hercules, CA) and
20	Staphylococcus aureus micrococcal nuclease from Worthington (Lakewood, NJ).
21	
22	

Downloaded from jb.asm.org at University of Missouri-Columbia on October 27, 2009

1 **Protein Purification**

2 SecA, SecAC4, SecAdN7, SecAdN10, SecB, and SecBL75Q were purified from strains of *E.coli* 3 harboring plasmids that express the proteins as previously described (24, 25) except that cells 4 expressing wild-type SecA and those expressing SecAdN10 were suspended at 0.5 gram wet 5 weight of cell pellet per mL of buffer and disrupted using a French Press at 8000 psi. The P11 6 column was omitted from the SecAdN10 purification. For cross-linking experiments, 7 SecAC4S350C and SecAC4I641C were purified from strains harboring plasmids derived from 8 plasmid pT7secAC4 that carries a gene for SecA in which the four native cysteine residues were 9 substituted by serine (23). For each SecA species a single cysteine was introduced at the site of 10 interest by site-directed mutagenesis (Quickchange, Stratagene). Proteins were purified as 11 described (24) with minor changes. Micrococcal nuclease (314 units/mL final) was included 12 along with DNaseI (5000 units/mL final) to degrade nucleic acids and decrease the viscosity of 13 the lysate. SecAC4S350C was purified by chromatography on a HiTrap QAE column followed by a HiTrap BlueTM HP instead of P11. For purification of SecAC4I641C only a HiTrap BlueTM 14 15 HP column was used. Precursor galactose-binding protein was purified as described (29). The precursor of OmpA labeled with [³⁵S] –methionine was produced from strains harboring 16 17 plasmid pET503, which encodes proOmpA with the substitution C290S, and was purified as 18 described (7). The culture was grown in the M9 minimal medium supplemented with glycerol 19 (0.4% w/v) as carbon source, thiamine $(4 \mu \text{g/mL})$ and ampicillin (0.1 mg/mL). When the culture 20 reached an optical density of 0.6 at 560 nm, IPTG was added to 0.1 mM to induce proOmpA. Eighty minutes after induction, 1 mCi $[^{35}S]$ – methionine (90 µL) mixed with 900 µL of 30 µM 21 22 nonradioactive methionine was added to the culture. For use in the cross-linking experiments a 23 species of proOmpA with both native cysteines changed to serine, C290S and C302S, was

constructed by site-directed mutagenesis (Quickchange, Stratagene) and purified as described
 (7). The precursors were stored in buffers containing denaturant: 1 N guanidine hydrochloride
 (GuHCl), 10 mM HEPES-KOH, pH 7.6, 0.3 M KOAc, 5 mM Mg(OAc)₂, 1 mM EGTA for
 precursor galactose-binding protein and 4 M urea, 10 mM HEPES-KOH, pH 7.6, 0.1 M KOAc
 for proOmpA. All proteins were stored at -80°C.

6

7 Cytoplasmic Membrane Vesicles

Inverted cytoplasmic membrane vesicles were prepared as described (32) from E.coli strain 8 9 HB3616, harboring plasmids that express SecE and SecG (pMAN809 with tac-SecG insertion 10 (16)) and SecY (pMAN510 (16)) under the control of the tac promoter, except that to induce the 11 proteins IPTG was added to 1 mM. In order to remove endogenous SecA the isolated vesicles were exposed to 5 M urea in 50 mM TrisCl, pH 8, 2 mM dithiothreitol (DTT) on ice for 30 min, 12 centrifuged (65000 rpm, 30 min, 4°C, type 65 rotor, Beckman Coulter, Fullerton, CA), and 13 14 suspended in 10 mM HEPES-KOH, pH 7.6, 0.3 M KOAc, 5 mM Mg(OAc)₂, 2 mM DTT. The vesicles were tested for translocation of precursors and for ATPase activity in the absence of 15 16 added SecA. There was no protection of either precursor ligand and no detectable hydrolysis of 17 ATP within the first 2 minutes. All calculations of efficiency of coupling were done within this 18 time frame.

19

20 Translocation and ATPase Assays

Both assays were done in the same reaction mixture, which was made up in glass tubes (12 x 75
mm) so that temperature equilibration would occur rapidly. All mixtures contained 13 mM
HEPES-KOH, pH 7.6, 250 mM KOAc, 5 mM Mg(OAc)₂, 2 mM DTT, 3.3 mM [γ-³²P] ATP

1	(specific activity: 1.8 Ci/mole), and 2 μM SecA dimer. When specified, SecB was added to 2 μM
2	tetramer and urea-treated inverted membrane vesicles to a final concentration of 0.6 mM lipid.
3	The reactions were initiated by dilution of precursor (precursor galactose-binding protein or
4	$[^{35}S]$ -methionine proOmpA) to 2 μ M from denaturant and the glass tubes were immediately
5	transferred to a water bath at 30°C. For reaction mixtures containing precursor galactose-binding
6	protein the final concentration of GuHCl was 12 mM and for those containing proOmpA the
7	final urea concentration was 36 mM. For the experiments in which we determine the ratio of
8	ATP hydrolysis to translocation we do not provide NADH or an ATP regenerating system
9	because we are measuring hydrolysis of ATP using $[\gamma^{-32}P]$ ATP.
10	

For assessment of both translocation and ATPase activity from the same reaction mixture, 11 12 samples of 10 μ L (from a total mixture of 100 μ L) were taken into tubes held on ice at times as 13 indicated, from which samples of 2 μ L were immediately removed to tubes with 2 μ L 0.1 M 14 EDTA on ice for the ATPase assay. To assess translocation, trypsin (5 mg/mL in 1 mM HCl) 15 was added to a final concentration of 0.5 mg/mL to the tubes containing 8 µL reaction mixture 16 and the samples were incubated for 15 min on ice at which time proteolysis was stopped by 17 addition of bovine pancreatic trypsin inhibitor (44 mg/mL in H₂O) to a final concentration of 5 18 mg/mL. For determination of the total amount of precursor added to the assay, 10 µL samples 19 were taken at time points 1 min and 6 min and no trypsin was added. Non-reducing SDS gel 20 sample buffer containing N-ethylmaleimide (8 mM) was added and the samples were boiled 21 immediately for 3 min and analyzed by SDS polyacrylamide gel electrophoresis on the same day 22 to avoid sample degradation.

1	Analyses by SDS Gel Electrophoresis, Immunoblotting and Thin Layer Chromatography
2	Polyacrylamide (10%, w/w) gels for experiments with $[^{35}S]$ -methionine labeled proOmpA were
3	dried and exposed to an imaging plate (Fuji Film, Stamford, CT) overnight, scanned using a
4	Phosphor Imager (Fuji Film, Stamford, CT), and analyzed with ImageGauge 4.0 (Fuji Film,
5	Stamford, CT). Immunoblots of 10% w/w gels were used to analyze all precursor galactose-
6	binding protein experiments and the experiments presented in Fig. 3. Blots were processed by
7	incubation with a rabbit antiserum raised to appropriate purified protein, then with goat
8	antibodies raised to rabbit IgGs and conjugated with horseradish peroxidase (Bio-Rad), followed
9	by staining with a 4-chloro-1-naphthol/hydrogen peroxide solution. All samples for precursor
10	galactose-binding protein were electrophoresed on the same gel as the purified precursor applied
11	in quantities of 5 ng, 10 ng, and 20 ng to generate a standard curve. The amount of protein at
12	each time point was determined using only those intensities that were within the linear range of
13	the standard. A Kodak EDAS 290 digital camera was used to capture images of the immunoblots
14	and TotalLab software (version 2.01; Nonlinear Dynamics Ltd.) was used to quantify the band
15	intensities. The concentration of the precursor added to the assay in combination with the
16	percentage of protein protected from trypsin digestion was used to determine the concentration
17	of precursor protected for both ligands.

19 Thin layer chromatography was used to analyze the hydrolysis of ATP. One microliter of the 20 samples taken into EDTA for the ATPase assay as described above was applied to a pre-coated 21 thin layer chromatography plate and dried. After application of all samples the plate was 22 developed in 125 mM KH₂PO₄ (14). After drying, the plates were exposed and scanned using the

- 1 Phosphor Imager. The ATPase activity was estimated from the proportion of total radioactivity 2 that migrated as inorganic phosphate ($R_f \sim 0.59$).
- 3

4 Calculation of the Efficiency of Coupling ATP Hydrolysis to Translocation

5 The coupling of the hydrolysis of ATP to the translocation of precursor polypeptides was

- 6 calculated using time points taken within the first 2 minutes. Early time points were used for two
- 7 reasons. Firstly, ADP has higher affinity for SecA than does ATP (9). Therefore, the
- 8 accumulation of ADP at later times would be expected to suppress activity. We did not include
- 9 an ATP regenerating system since we use the appearance of 32 P-phosphate as the assay for
- 10 hydrolysis. Secondly, as the process of translocation approaches a plateau the translocation
- 11 ATPase activity will be replaced by membrane ATPase activity since SecA without precursor
- 12 bound will still bind SecYEG. This would result in false values for the coupling.
- 13

14 Cross-linking with a Photoactivatable Reagent

15 Each of the SecA variants, SecAC4S350C and SecAC4I641C, has a single cysteine as specified.

- 16 Each variant was labeled with the sulfhydryl specific photoactivable reagent, N-[(2-
- 17 pyridyldithio)ethyl]-4-azidosalicylamide (AET) (1). The protein to be labeled was exchanged

18 into 100 mM Na₂B₄O₇, 100 mM KOAc, pH 8.3 using a NAPTM10 column. All subsequent steps

19 were done in the dark. AET (stored at 45 mM in DMSO at -80°C) was added at a 10-fold molar

- 20 excess over cysteine in the SecA and the mixture was incubated for 2 hours at room temperature
- 21 followed by 1 hour on ice. Free AET was removed by exchange of the protein into 10 mM
- 22 HEPES-HOAc, pH 6.7, 300 mM KOAc, 5 mM Mg(OAc)₂ using a NAPTM10 column. The AET
- 23 modification was confirmed by MALDI mass spectrometry. Mixtures (~ 30μ L) of the AET-

labeled SecA (12 µM dimer) and precursor (either precursor galactose-binding protein or proOmpA at 12 µM) in 10 mM HEPES-HOAc, pH 6.7, 300 mM KOAc, 5 mM Mg(OAc)₂ were prepared and placed in the shallow spots of a porcelain spot plate held on ice and irradiated with a mercury lamp for 1 min. Samples were analyzed by SDS polyacrylamide (10%, w/w) gel electrophoresis using both reducing and non-reducing sample buffers. Gels (both reduced and non-reduced samples) were run in duplicate and subjected to immunoblotting using antisera to

ion reduced samples) were run in duprede und subjected to minimuliorotating using

7 purified SecA as well as to the relevant precursor.

8

Characterization of Protein Preparations and Determination of Lipid Concentration 9 10 Protein concentrations were determined spectrophotometrically at 280 nm using extinction coefficients of 47,600 M⁻¹cm⁻¹ for SecB tetramer, 157,800 M⁻¹cm⁻¹ for SecA dimer, 52,955 11 $M^{-1}cm^{-1}$ for proOmpA and 37,410 $M^{-1}cm^{-1}$ for precursor galactose-binding protein. Since 12 13 modification by AET changes the absorbance of the SecA variants, those concentrations were 14 determined by electrophoresis of at least three samples of the protein in increasing quantity on 15 the same SDS polyacrylamide (14%, w/w) gel as three quantities of pure SecA of known 16 concentration to generate a standard curve. The amount of protein in the standards and the 17 variants was determined as described for quantification of protein on immunoblots. 18 Concentration of lipid in the vesicle preparation was determined as described on the Avanti Polar 19 Lipids website (www.avantilipids.com) using an average molar mass for *E.coli* lipids of 741 Da. 20 21 All protein preparations used in this study were subjected to rigorous characterization. They were 22 analyzed by mass spectrometry. The only degradation detected was in SecAdN10 where 23 approximately 10% of the protein was cleaved to generate two fragments of 67,000 Da and

1	33,000 Da. The ratio of absorbance at 280 nm to 260 nm for all preparations was 2.0 except for
2	SecAC4, which was 1.8, thus there is no significant contamination by nucleic acid or
3	nucleotides. Column chromatography used with a static light scatter detector and analytical
4	centrifugation showed that the protein preparations contained no aggregation and formed the
5	expected complexes with stoichiometry of either A1:B4 or A2:B4 complex consistent with our
6	previously published results (24). The purity of all preparations was between 80% and 90%.
7	
8	RESULTS
9	Translocation of Precursors and ATP Hydrolysis Mediated by SecA Variants with
10	Different Dimer Equilibria
11	Complexes between wild-type SecA and SecB as well as between variants were assayed for both
12	translocation of precursors and the associated translocation ATPase activity to determine the
13	effect of varying the number of protomers of SecA in the complex. The species of SecA studied
14	were wild-type SecA and three variants of SecA that have been characterized previously (5, 24),
15	SecAdN10 (amino acids 2 through 11 deleted), SecAdN7 (amino acids 2 through 8 deleted) and
16	SecAC4, which lacks zinc. In addition to the complex between wild-type SecA and SecB we
17	studied a complex containing a SecB variant, SecBL75Q, which carries a substitution that
18	interferes with binding the zinc site on SecA. In earlier work (24) we used size-exclusion
19	chromatography coupled with static light scatter to demonstrate that the complex between SecB
20	and SecA has a mass of 272 kDa, indicating that the stoichiometry of the wild-type complex is
21	two protomers of SecA (mass, 204 kDa for dimer) bound to a tetramer of SecB (mass, 69 kDa),
22	referred to hereafter as A2:B4. It is possible to populate complexes containing only one protomer
23	of SecA because although SecA and SecB both display two-fold symmetry the A2:B4 complex is

1 stabilized by contacts that are distributed asymmetrically. Only one protomer of SecA can bind if 2 the area of contact that is between residues in the β -sheets that form the flat sides of SecB and 3 the C-terminal zinc domain of SecA is eliminated. Here we have examined complexes that have 4 that contact disrupted in two different ways: wild-type SecA bound to a mutant of SecB that has 5 a residue located on the flat β -sheet changed, SecBL75Q, and wild-type SecB bound to a SecA 6 variant, SecAC4, that lacks zinc because the cysteines that coordinate the zinc are replaced by 7 serines. Both complexes were shown to have a stoichiometry of A1:B4 (24), i.e. a molar mass of 8 171 kDa.

9

Complexes with a stoichiometry of A1:B4 can also be populated by using species of SecA that 10 11 have the monomer - dimer equilibrium altered by truncation at the N-terminus. Deletion of amino acids 2 through 8, SecAdN7, shifts the equilibrium toward monomer ($K_d \sim 24 \mu M$) (5), 12 whereas deletion of amino acids 2 through 11 results in a species, SecAdN10, that exists only as 13 a monomer ($K_d > 230 \mu M$) (5). SecAdN10 was shown to form a complex with SecB of mass 169 14 15 kDa corresponding to a stoichiometry of A1:B4 (24), whereas the mass observed for a mixture of 16 SecAdN7 and SecB was 235 kDa indicating an equilibrating population that contains both 17 A1:B4 and A2:B4. It is not possible to estimate the amount of any one species in such a mixture 18 because it would contain not only complexes with stoichiometry A1:B4 and A2:B4 but also free 19 SecA which would itself be in equilibrium between monomer and dimer.

20

The various combinations of SecA and SecB displaying different equilibrating mixtures of
complexes with A1:B4 and A2:B4 stoichiometries were tested for the translocation of precursor
into inverted membrane vesicles and for the coupled hydrolysis of ATP. Translocation of the

1	precursor of the outer membrane protein OmpA, proOmpA, mediated by SecAdN10:SecB
2	complexes (A1:B4) is 22% of that mediated by complexes containing wild-type SecA (A2:B4),
3	$0.19 \mu\text{M}$ versus $0.86 \mu\text{M}$, respectively at the 6 min time point; whereas ATP hydrolysis is 52%
4	of the activity of the wild-type complex (Figs. 1A and 1B). Similar results are seen for
5	translocation of precursor galactose-binding protein (Figs. 2A and 2B); translocation of precursor
6	by the SecAdN10:SecB complex is approximately 24% of that mediated by a wild-type complex,
7	whereas ATP hydrolysis is 37% of wild-type activity. These data indicate that the efficiency of
8	coupling of hydrolysis of ATP to movement of the precursor through the translocon is much
9	poorer for the SecA:SecB complex which has a stoichiometry of A1:B4 than it is for complexes
10	having two protomers bound to SecB as in the wild-type complexes. The translocation activity of
11	complexes containing SecB and SecAdN7 (Figs. 1A and 2A) lies between the activity of wild-
12	type complexes and that of complexes containing SecAdN10. As described above, the
13	equilibrating population of complexes between SecAdN7 and SecB contains both A1:B4 and
14	A2:B4; therefore, we conclude that shifting the equilibrium to complexes containing two
15	protomers of SecA results in more robust translocation. In addition the efficiency of coupling of
16	ATP hydrolysis to translocation is improved. The data in Figures 1 and 2 were used to calculate
17	the efficiency of coupling of hydrolysis of ATP to translocation of a precursor polypeptide
18	(Table 1, See Materials and Methods for calculations). The efficiency of coupling for proOmpA
19	was the highest for the A2:B4 wild-type complex which hydrolyzed approximately 2500 moles
20	of ATP per mole of precursor protected. Complexes with SecAdN7, which contain a mixture of
21	A1:B4 and A2:B4, displayed a coupling of 3600 moles ATP hydrolyzed per mole precursor
22	protected and the lowest efficiency was observed with A1:B4 complexes: SecAdN10 and SecB,
23	zincless SecAC4 and SecB, and SecA and SecBL75Q. The coupling efficiency for precursor

2	intermediate for complexes formed with SecAdN/ and lowest for those complexes which
3	populate only A1:B4 complexes (Table 1).
4	
5	Since the stoichiometry between SecA and SecB has been varied using both full length SecA as
6	well as a truncated species and wild-type SecB as well as a variant of SecB, it is safe to conclude
7	that the observed decrease in efficiency of the coupling of ATP hydrolysis to translocation
8	results from a decrease in occupancy of complexes having a stoichiometry of A2:B4 and not
9	from either the truncations of SecA or the mutation of SecB.
10	
11	Stoichiometry of Complexes between SecA and Precursors
12	Although SecA mediates a higher level of translocation when it receives precursors via a
13	SecB:precursor complex as opposed to binding precursor directly, it does function <i>in vitro</i> in the
14	absence of SecB (Fig. 3). SecAdN10, which is monomeric shows no detectable processing in the
15	absence of SecB; whereas SecAdN7, which does populate dimer but to a lesser extent than does
16	wild-type (24), shows a low level of translocation which is greatly enhanced by addition of
17	SecB. SecA can mediate translocation in the absence of SecB in vivo as evidenced by the
18	viability of SecB null strains (13). Therefore, it is of interest to determine the stoichiometry of a
19	complex between SecA and precursors in the absence of SecB. We have not been able to
20	demonstrate a complex between SecA and precursors using size exclusion chromatography;
21	therefore, we can not use the approaches we have applied previously to reveal stoichiometry by
22	determination of molar mass using static light scatter (24). As an alternative we have used cross-

galactose binding protein showed the same trend, most efficient for wild-type complexes,

1

23 linking. We chose a photoactivatable cross-linking reagent, N-[(2-pyridyldithio)ethyl]-4-

1	azidosalicylamide (AET) to determine whether one protomer of SecA or two interact with one
2	precursor polypeptide. The reagent was attached via a disulfide exchange reaction to two SecA
3	variants, each having a single cysteine, one at position 350 and the other at position 641. These
4	sites were identified as contact sites between SecA and its precursor ligands by site-directed spin
5	labeling and electron paramagnetic resonance (EPR) spectroscopy (3). Irradiation of solutions
6	that contained complexes between the derivatized SecA variants and either proOmpA or
7	precursor galactose-binding protein generated covalent linkages between the proteins. Analyses
8	of the irradiated samples by SDS polyacrylamide gel electrophoresis followed by
9	immunoblotting using antisera to SecA and to each of the precursors revealed that the complex
10	formed between SecA and either proOmpA or precursor galactose-binding protein had a molar
11	mass of approximately 250 kDa (Fig. 4). This was true whether the reagent was linked to SecA
12	at amino acid position 350 or at position 641. Addition of reducing agent to the SDS gel sample
13	buffer caused disappearance of the high molecular weight cross-linked species (data not shown)
14	as expected since the AET moiety is linked to SecA via a disulfide bond. The ability of the AET-
15	labeled SecAC4I641 to cross-link to SecB was tested as a control for specificity. EPR studies (3)
16	show that SecAC4I641 is not a site of contact within the SecA-SecB complex. As expected no
17	cross-linked species was observed (data not shown). Bands migrating with an apparent mass of
18	approximately 150 kDa that appeared upon irradiation even in the absence of a precursor ligand
19	are likely to be cross-linked dimers of the two SecA species, SecAC4S350AET and
20	SecAC4I641AET. The aberrant migration of the dimers can be explained because the pairs of
21	SecA polypeptides are tethered at two internal positions thereby preventing the chain from
22	becoming fully extended in SDS.

We conclude that each precursor species binds two protomers of SecA to give the observed mass, approximately 250 kDa (204 kDa for SecA dimer and 37 kDa for proOmpA or 36 kDa for precursor galactose-binding protein). Since chemical cross-linking results in an irreversible reaction we have no information relating to the affinity of SecA for the precursors or whether the protomers of SecA bind to the precursor sequentially or as a dimer.

- 6
- 7

DISCUSSION

We have previously shown that formation of a complex between SecA and SecB that is capable 8 of mediating a high level of translocation of precursors requires two protomers of SecA bound to 9 10 a tetramer of SecB. Within the complex there are three areas of contact. One site involves the C-11 terminal zinc containing region of SecA which interacts with negatively charged region on the 12 flat β -sheets that form the sides of the SecB dimer of dimers (4). A second interaction is between 13 the extreme C-terminal flexible tail of SecB and the amino-terminal 11 amino acids of SecA. A 14 third area of contact that provides energy of stabilization is less defined but involves residues 15 lying on the β -sheets of SecB as well as along the interface of the dimer of dimers (22). Even 16 though each of the binding partners displays two-fold symmetry, the contacts between them are 17 distributed asymmetrically. When the contact between the zinc domain on SecA and the side of 18 SecB is disrupted or when 10 amino acyl residues are deleted from the amino terminus of SecA, 19 only one protomer of SecA is bound yielding a complex of stoichiometry A1:B4 that displays a 20 very low activity both in vivo (12) and in vitro (10, 24, 25, 32).

21

Here we have asked why two protomers of SecA must be present for full activity in the reaction
cycle of SecA and SecB during translocation. We have made use of variants of both SecA and

1 SecB and the complexes they form, all of which are well characterized. Assays of the extent of 2 translocation of precursors and the associated translocation ATPase activity have allowed us to 3 establish a correlation between an increase in the efficiency of coupling of ATP hydrolysis to 4 translocation of polypeptides and an increase in the population of complexes containing two 5 protomers of SecA bound to SecB. We have established this correlation by varying the 6 population of complexes that have the stoichiometry of A1:B4 and A2:B4 by using species of 7 SecA truncated at the extreme amino terminus with wild-type and a truncated SecB, wild-type 8 SecA with a mutant of SecB, and wild-type SecB with full-length SecA containing no zinc. In 9 every case the higher efficiency of coupling occurs when the A2:B4 complex is more populated. 10 The construct with amino acids 2 through 8 deleted (SecAdN7) was previously studied by Mori 11 et al (17) under the name of SecA N-8. The two preparations differ in that our purified protein 12 retains the N terminal methionine whereas theirs does not. It was shown that SecA N-8 was 13 defective in protein translocation, the translocation ATPase activity and the topological inversion 14 of SecG. The investigators concluded that the amino-terminal region of SecA is involved in 15 functional interaction with SecG. In our work described here we have eliminated the possibility 16 that the effects we observe are specific for deletions at the amino terminus. It seems possible that 17 the effects observed by Mori et al (17) also result from the oligomeric state of SecA and not 18 directly from the lack of seven amino acyl residues. Perhaps the inversion of SecG involves two 19 protomers of SecA.

20

By what mechanism might two protomers of SecA increase the coupling of ATP hydrolysis to
movement? Lill *et al.* (15) showed that in a system using inverted membrane vesicles, such as
that we use here, SecA demonstrates a high rate of non-productive hydrolysis, hydrolyzing more

than 1000 moles of ATP per mole of precursor translocated, a level similar to that observed here 1 2 for wild-type SecA (Table 1) as well as that observed by others (28). It was subsequently shown 3 by Schiebel *et al.* (26) that the poor coupling is the result of a backward slippage of the precursor 4 undergoing translocation. It might be that the two protomers of SecA are required to work 5 together to prevent the backward movement. One protomer might insert a segment of the bound 6 precursor polypeptide into the translocon channel and release it upon hydrolysis of ATP; the 7 second protomer would remain bound to the next more distal segment of the precursor 8 polypeptide and, if also bound to SecYEG, would prevent backward movement of the 9 polypeptide chain. This idea is consistent with observations by others (8, 19). Osborne and Rapoport (19), based on cross-linking of SecA and proOmpA to a covalent dimeric form of 10 SecYEG, proposed that one copy of SecYEG serves as the translocation channel and the other 11 provides a static binding site for SecA. These authors proposed that only a single protomer of 12 13 SecA is involved, but a slight modification adapts this model to the idea we have put forth here, 14 two protomers of SecA act together to translocate precursors, one staying bound while the other 15 releases the polypeptide into the channel and dissociates from SecYEG. Duong (8) observed 16 association of both monomeric and dimeric SecA to the translocon using the same covalent 17 dimeric SecYEG, but upon addition of ATP only SecA monomers remained associated.

18

SecA can mediate translocation of precursors through the Sec secretory pathway in the absence of SecB both *in vivo* (13) and *in vitro*. It is currently debated whether when acting in the absence of SecB, SecA functions as a monomer or dimer. Although the *in vitro* translocation activity is very low without SecB, the trend is the same as seen with SecB complexes: SecAdN7 shows very low but detectable activity whereas SecAdN10 shows none. This correlation suggests that although monomers do function the dimers are more efficient. We have shown here that even
when SecA interacts directly with precursors two protomers can associate with each precursor
polypeptide. Since it is unlikely that SecA has a different mechanism of coupling hydrolysis to
movement of the polypeptide through the translocon in the presence and absence of SecB, the
presence of two protomers might act together as proposed here to prevent backward slippage.

6

We conclude that maximal efficiency of translocation is achieved when two protomers of SecA in complex with SecB act together. A single protomer of SecA can perform the necessary conversion of chemical energy (hydrolysis of ATP) to mechanical work (movement of the precursor through the translocon), but tight coupling of this conversion requires two protomers. An exciting direction for future research is elucidation of the movement within SecA at the resolution of aminoacyl sidechains that transduces the chemical energy of hydrolysis of ATP to insertion of precursors into and through the translocon.

- 14
- 15

ACKNOWLEDGEMENTS

We thank Donald Oliver (Wesleyan University) for supplying the plasmid pT7*secAC4* that
carries the *secA* gene with the four native cysteine residues changed to serine residues, Angela A.
Lilly for construction of the SecA variants, Beverly DaGue (The Proteomics Center, University
of Missouri) for MALDI mass spectrometry, Hilary Roth for assistance with the *in vitro* activity
assays and Jennine M. Crane for critically reading the manuscript. This work was supported by
NIH research grant GM29798 to L.L.R..

1		REFERENCES
2	1.	Cai, K., Y. Itoh, and H. G. Khorana. 2001. From the Cover: Mapping of contact sites in
3		complex formation between transducin and light-activated rhodopsin by covalent crosslinking:
4		Use of a photoactivatable reagent. Proc. Natl. Acad. Sci. U. S. A. 98:4877-4882.
5	2.	Chen, Y., X. Pan, Y. Tang, S. Quan, P. C. Tai, and SF. Sui. 2008. Full-length Escherichia
6		coli SecA dimerizes in a closed conformation in solution as determined by cryo-electron
7		microscopy. J. Biol. Chem.:C800160200.
8	3.	Cooper, D. B., V. F. Smith, J. M. Crane, H. C. Roth, A. A. Lilly, and L. L. Randall. 2008.
9		SecA, the motor of the secretion machine, binds diverse partners on one interactive surface. J Mol
10		Biol 382: 74-87.
11	4.	Crane, J. M., C. Mao, A. A. Lilly, V. F. Smith, Y. Suo, W. L. Hubbell, and L. L. Randall.
12		2005. Mapping of the docking of SecA onto the chaperone SecB by site-directed spin labeling:
13		insight into the mechanism of ligand transfer during protein export. J Mol Biol 353:295-307.
14	5.	Das, S., E. Stivison, E. Folta-Stogniew, and D. Oliver. 2008. Re-examination of the Role of the
15		Amino-Terminus of SecA in Promoting Its Dimerization and Functional State. J.
16		Bacteriol :JB.00593-08.
17	6.	Dekker, C., B. de Kruijff, and P. Gros. 2003. Crystal structure of SecB from Escherichia coli. J.
18		Struct. Biol. 144:313-319.
19	7.	Does, C. v. d., J. d. Keyzer, M. v. d. Laan, and A. J. M. Driessen. 2003. Reconstitution of
20		Purified Bacterial Preprotein Translocase in Liposomes, p. 86-98. In N. Duzgunes (ed.),
21		Liposomes, Part B, vol. 372. Academic Press.
22	8.	Duong, F. 2003. Binding, activation and dissociation of the dimeric SecA ATPase at the dimeric
23		SecYEG translocase. EMBO J. 22:4375-4384.
24	9.	Fak, J. J., A. Itkin, D. D. Ciobanu, E. C. Lin, X. J. Song, Y. T. Chou, L. M. Gierasch, and J.
25		F. Hunt. 2004. Nucleotide exchange from the high-affinity ATP-binding site in SecA is the rate-

1		limiting step in the ATPase cycle of the soluble enzyme and occurs through a specialized
2		conformational state. Biochemistry 43: 7307-27.
3	10.	Fekkes, P., J. G. de Wit, A. Boorsma, R. H. Friesen, and A. J. Driessen. 1999. Zinc stabilizes
4		the SecB binding site of SecA. Biochemistry 38: 5111-5116.
5	11.	Hunt, J. F., S. Weinkauf, L. Henry, J. J. Fak, P. McNicholas, D. B. Oliver, and J.
6		Deisenhofer. 2002. Nucleotide control of interdomain interactions in the conformational reaction
7		cycle of SecA. Science 297: 2018-2026.
8	12.	Kimsey, H. H., M. D. Dagarag, and C. A. Kumamoto. 1995. Diverse effects of mutation on the
9		activity of the Escherichia coli export chaperone SecB. J. Biol. Chem. 270:22831-22835.
10	13.	Kumamoto, C. A., and J. Beckwith. 1985. Evidence for specificity at an early step in protein
11		export in Escherichia coli. J. Bacteriol. 163:267-274.
12	14.	Levit, M. N., T. W. Grebe, and J. B. Stock. 2002. Organization of the receptor-kinase signaling
13		array that regulates Escherichia coli chemotaxis. J. Biol. Chem. 277:36748-36754.
14	15.	Lill, R., K. Cunningham, L. A. Brundage, K. Ito, D. Oliver, and W. Wickner. 1989. SecA
15		protein hydrolyzes ATP and is an essential component of the protein translocation ATPase of
16		Escherichia coli. EMBO J. 8:961-966.
17	16.	Matsuyama, S., J. Akimaru, and S. Mizushima. 1990. SecE-dependent overproduction of
18		SecY in Escherichia coli. Evidence for interaction between two components of the secretory
19		machinery. FEBS Letters 269:96-100.
20	17.	Mori, H., H. Sugiyama, M. Yamanaka, K. Sato, M. Tagaya, and S. Mizushima. 1998.
21		Amino-Terminal Region of SecA Is Involved in the Function of SecG for Protein Translocation
22		into Escherichia coli Membrane Vesicles. J Biochem 124:122-129.
23	18.	Osborne, A. R., W. M. Clemons, Jr., and T. A. Rapoport. 2004. A large conformational
24		change of the translocation ATPase SecA. Proc. Natl. Acad. Sci. U. S. A. 101:10937-42.
25	19.	Osborne, A. R., and T. A. Rapoport. 2007. Protein Translocation Is Mediated by Oligomers of
26		the SecY Complex with One SecY Copy Forming the Channel. Cell 129: 97-110.

1	20.	Papanikolau, Y., M. Papadovasilaki, R. B. G. Ravelli, A. A. McCarthy, S. Cusack, A.
2		Economou, and K. Petratos. 2007. Structure of Dimeric SecA, the Escherichia coli Preprotein
3		Translocase Motor. J. Mol. Biol. 366:1545-1557.
4	21.	Papanikou, E., S. Karamanou, and A. Economou. 2007. Bacterial protein secretion through
5		the translocase nanomachine. Nat. Rev. Microbiol. 5:839-851.
6	22.	Patel, C. N., V. F. Smith, and L. L. Randall. 2006. Characterization of three areas of
7		interactions stabilizing complexes between SecA and SecB, two proteins involved in protein
8		export. Protein Sci 15:1379-86.
9	23.	Ramamurthy, V., and D. Oliver. 1997. Topology of the integral membrane form of Escherichia
10		coli SecA protein reveals multiple periplasmically exposed regions and modulation by ATP
11		binding. J. Biol. Chem. 272:23239-23246.
12	24.	Randall, L. L., J. M. Crane, A. A. Lilly, G. Liu, C. Mao, C. N. Patel, and S. J. Hardy. 2005.
13		Asymmetric binding between SecA and SecB two symmetric proteins: implications for function
14		in export. J. Mol. Biol. 348: 479-489.
15	25.	Randall, L. L., J. M. Crane, G. Liu, and S. J. Hardy. 2004. Sites of interaction between SecA
16		and the chaperone SecB, two proteins involved in export. Protein Sci 13:1124-1133.
17	26.	Schiebel, E., A. J. Driessen, F. U. Hartl, and W. Wickner. 1991. Delta mu H+ and ATP
18		function at different steps of the catalytic cycle of preprotein translocase. Cell 64:927-939.
19	27.	Sharma, V., A. Arockiasamy, D. R. Ronning, C. G. Savva, A. Holzenburg, M. Braunstein,
20		W. R. Jacobs, Jr., and J. C. Sacchettini. 2003. Crystal structure of Mycobacterium tuberculosis
21		SecA, a preprotein translocating ATPase. Proc. Natl. Acad. Sci. U. S. A. 100:2243-2248.
22	28.	Tomkiewicz, D., N. Nouwen, R. van Leeuwen, S. Tans, and A. J. M. Driessen. 2006. SecA
23		Supports a Constant Rate of Preprotein Translocation. J. Biol. Chem. 281:15709-15713.
24	29.	Topping, T. B., and L. L. Randall. 1997. Chaperone SecB from Escherichia coli mediates
25		kinetic partitioning via a dynamic equilibrium with its ligands. J. Biol. Chem. 272:19314-19318.

Downloaded from
<u> </u>
.asm.org
a
Ē
Jniversity
0
f Missouri-Columbia on
October 27,
, 2009

1	30.	Vassylyev, D. G., H. Mori, M. N. Vassylyeva, T. Tsukazaki, Y. Kimura, T. H. Tahirov, and
2		K. Ito. 2006. Crystal Structure of the Translocation ATPase SecA from Thermus thermophilus
3		Reveals a Parallel, Head-to-Head Dimer. J. Mol. Biol. 364:248-258.
4	31.	Woodbury, R. L., S. J. Hardy, and L. L. Randall. 2002. Complex behavior in solution of
5		homodimeric SecA. Protein Sci. 11:875-882.
6	32.	Woodbury, R. L., T. B. Topping, D. L. Diamond, D. Suciu, C. A. Kumamoto, S. J. Hardy,
7		and L. L. Randall. 2000. Complexes between protein export chaperone SecB and SecA.
8		Evidence for separate sites on SecA providing binding energy and regulatory interactions. J. Biol.
9		Chem. 275:24191-24198.
10	33.	Xu, Z., J. D. Knafels, and K. Yoshino. 2000. Crystal structure of the bacterial protein export
11		chaperone secB. Nat. Struct. Biol. 7:1172-1177.
12	34.	Zimmer, J., W. Li, and T. A. Rapoport. 2006. A Novel Dimer Interface and Conformational
13	ſ	Changes Revealed by an X-ray Structure of B. subtilis SecA. J. Mol. Biol. 364: 259-265.

FIGURE LEGENDS

2	Figure 1. Translocation of proOmpA mediated by A1:B4 and A2:B4 complexes. A. Protection of	
3	proOmpA. B. Associated ATPase activity of the translocation. In vitro assays contained	
4	proOmpA, SecA species and SecB species at 2 μ M each expressed as dimeric SecA and	
5	tetrameric SecB: wild-type SecA (circles), SecAdN7 (down triangles), and SecAdN10 (squares)	
6	with wild-type SecB; SecAC4 with wild-type SecB (up triangles); wild-type SecA with	
7	SecBL75Q (diamonds). The data here and in Figures 2 and 3 were fitted to a hyperbola using	
8	SigmaPlot 2001 software. The error bars shown are standard deviations. All assays were done at	
9	least three times. When error bars are not obvious, they fall within the symbols. For all fits, 0.45	
10	min was used as the intercept on the x-axis.	
11		
12	Figure 2. Translocation of precursor galactose-binding protein mediated by A1:B4 and A2:B4	
13	complexes. A. Protection of precursor galactose-binding protein (pGBP). B. Associated ATPase	
14	activity of the translocation. Concentrations of proteins and symbols in the figures are the same	
15	as in Figure 1. The samples were analyzed by immunoblotting.	
16		
17	Figure 3. Translocation of proOmpA in the presence and absence of SecB. In vitro translocation	
18	of 2 μ M proOmpA by SecA wild-type (circles) or SecAdN7 (triangles) at 2 μ M SecA expressed	
19	as dimer was carried out in the presence (filled symbols) or absence (open symbols) of 2 μM	
20	SecB expressed as tetramer. The efficiency of translocation without SecB is extremely low;	
21	therefore, to maximize activity for these experiments 1.7 mM NADH, 7.5 mM phosphocreatine,	
22	and 37 mg/mL creatine phosphokinase were included in the <i>in vitro</i> system to regenerate ATP.	
23	The samples were analyzed by immunoblotting.	

Figure 4. Cross-linking of AET-SecA to precursors. A. Cross-linking to proOmpA. B. Crosslinking to precursor galactose-binding protein (pGBP). Immunoblots using antisera to SecA are
shown in the upper panels and those using antisera to the precursors in the lower panels. Crosslinked complexes are indicated by dots and irradiation by hv. Molecular mass markers are shown
to the right.

SecA and SecB complexes tested Precursor SecA WT SecAC4 SecA WT SecAdN7 SecAdN10 SecBL75Q SecB WT SecB WT SecB WT SecB WT 5600 proOmpA 2500 3600 5800 5800 N.D.^b 11500 11500 pGBP 4200 6800

Table 1. Coupling of ATP hydrolysis to translocation^a

^a Mole ATP hydrolyzed per mole precursor protected

5 6

7

8

9 10

11 12 13

14 15

16

17 18 ^b Not determined, reliable results could not be generated because of the low level of precursor protected







