

# ATPase Active-Site Electrostatic Interactions Control the Global Conformation of the 100 kDa SecA Translocase

Dorothy M. Kim,<sup> $\dagger, \ddagger, \perp$ </sup> Haiyan Zheng,<sup> $\$, \parallel$ </sup> Yuanpeng J. Huang,<sup>\$</sup> Gaetano T. Montelione,<sup> $\$, \parallel$ </sup> and John F. Hunt<sup>\*,<sup>†</sup></sup>

<sup>†</sup>Department of Biological Sciences and Northeast Structural Genomics Consortium, 702A Fairchild Center, MC2434, Columbia University, New York, New York 10027, United States

<sup>‡</sup>Department of Biochemistry and Molecular Biophysics, Columbia University, 650 West 168th Street, New York, New York 10032, United States

<sup>§</sup>Center for Advanced Biotechnology and Medicine, Department of Molecular Biology and Biochemistry, and Northeast Structural Genomics Consortium, Rutgers, The State University of New Jersey, Piscataway, New Jersey 08854, United States

<sup>II</sup>Department of Biochemistry and Molecular Biology, Robert Wood Johnson Medical School, University of Medicine and Dentistry of New Jersey, Piscataway, New Jersey 08854, United States

# **Supporting Information**

**ABSTRACT:** SecA is an intensively studied mechanoenzyme that uses ATP hydrolysis to drive processive extrusion of secreted proteins through a protein-conducting channel in the cytoplasmic membrane of eubacteria. The ATPase motor of SecA is strongly homologous to that in DEAD-box RNA helicases. It remains unclear how local chemical events in its ATPase active site control the overall conformation of an ~100 kDa multidomain enzyme and drive protein transport. In this paper, we use biophysical methods to establish that a single electrostatic charge in the ATPase active site controls the global conformation of SecA. The enzyme undergoes an ATP-modulated endothermic conformational transition (ECT) believed to involve similar structural mechanics to the protein transport reaction. We have characterized the effects of an isosteric glutamate-to-glutamine mutation in the catalytic base, a mutation which mimics the immediate electrostatic consequences of ATP hydrolysis in the active site.



Calorimetric studies demonstrate that this mutation facilitates the ECT in *Escherichia coli* SecA and triggers it completely in *Bacillus subtilis* SecA. Consistent with the substantial increase in entropy observed in the course of the ECT, hydrogen– deuterium exchange mass spectrometry demonstrates that it increases protein backbone dynamics in domain–domain interfaces at remote locations from the ATPase active site. The catalytic glutamate is one of ~250 charged amino acids in SecA, and yet neutralization of its side chain charge is sufficient to trigger a global order–disorder transition in this 100 kDa enzyme. The intricate network of structural interactions mediating this effect couples local electrostatic changes during ATP hydrolysis to global conformational and dynamic changes in SecA. This network forms the foundation of the allosteric mechanochemistry that efficiently harnesses the chemical energy stored in ATP to drive complex mechanical processes.

# INTRODUCTION

Following synthesis in the cytoplasm, secreted protein domains require translocation through cellular membranes by active transport. In bacteria, the Sec system provides a mechanism for post-translational translocation of proteins that function outside of the cytoplasm. Secreted "preproteins" contain a cleavable Nterminal hydrophobic extension, called a signal sequence.<sup>1</sup> This extension targets them for delivery to SecA,<sup>2–4</sup> a cytosolic ATPase that drives their extrusion through a heterotrimeric protein-conducting channel called SecYEG.<sup>5–7</sup> SecA is a 100 kDa enzyme that contains six globular domains: an N-terminal pair of 20–30 kDa domains comprising its "DEAD-box" ATPase motor (described in detail below), an 18 kDa "preprotein crosslinking domain", a 16 kDa  $\alpha$ -helical scaffold domain (HSD), an 11 kDa  $\alpha$ -helical wing domain (HWD), and a C-terminal 2.5 kDa Zn<sup>2+</sup>-binding domain. SecA may partially insert into the pore of SecYEG as its ATPase drives preprotein extrusion.<sup>8,9</sup> Each molecule of ATP hydrolyzed by SecA leads to translocation of 10–15 amino acids of preprotein.<sup>10,11</sup> Based on mechanochemical studies of SecA and evolutionarily related ATPases,<sup>12–15</sup> this translocation reaction seems likely to be driven by mechanical rearrangements of SecA's domains coupled to its ATPase cycle. Such motions could processively push the preprotein itself through the pore of SecYEG while SecA remains bound to the surface of SecYEG,<sup>16,17</sup> or alternatively, they could control insertion and retraction of SecA from SecYEG, which would drive transport if preprotein were bound to SecA during insertion but released during retraction.<sup>12,18,19</sup> However, both of these models are speculative, and the mechanism by which SecA drives preprotein transport remains unclear despite extensive mechanistic,<sup>10,11,18–21</sup> thermodynamic,<sup>13,22–27</sup> and structural studies.<sup>12,16,28–32</sup>

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A fundamental question regarding SecA function is how hydrolysis of a single phosphodiester bond at its ATPase active site can control the functional conformational transitions of this large mechanoenzyme. The chemistry of the hydrolysis reaction is straightforward:

SecA + ATP 
$$\rightarrow$$
 SecA·ATP  
 $\rightarrow$  SecA·ADP +  $P_i$   
 $\rightarrow$  SecA + ADP +  $F_i$ 

Extensive research has shown that catalysis of this reaction is critically dependent on electrostatic interactions with ATP and its Mg<sup>2+</sup> cofactor in the active site of ATPases.<sup>33,34</sup> However, it remains unclear how these interactions and their variation during the hydrolysis reaction drive the large-scale conformational transitions that perform thermodynamic work in SecA and other ATP-driven mechanoenzymes.

The ATPase active site in SecA is located at the interface of domains called nucleotide-binding folds I and II (NBF-I and NBF-II),<sup>12</sup> which together comprise the DEAD-box ATPase motor. NBF-I and NBF-II are believed to function as an ATPdependent mechanical clamp as they close around the nucleotide bound at their mutual interface.<sup>12,28</sup> These domains bear the strongest structural and sequence similarity to the ATPase domains in superfamily I and II ATP-dependent helicases,<sup>14,35</sup> especially those in the "DEAD-box" family of RNA helicases which perform a plethora of cellular functions.<sup>36-39</sup> This family takes its name from a conserved Asp-Glu-Ala-Asp (DEAD) sequence that includes several catalytically vital residues.<sup>39,40</sup> The glutamate in this sequence (E210 in Escherichia coli SecA) is the catalytic base that activates water for hydrolytic attack on ATP.41 DEAD-box motor domains have more remote but clearly significant structural homology to a much wider variety of mechanically active ATPases including F1 ATPase,<sup>15,42</sup> AAA ATPases, and ABC transporter<sup>43-46</sup> proteins. The ATPase motor domains from these diverse families share a conserved core  $\beta$ -sheet topology as well as stereotyped Walker A and B sequence motifs<sup>47</sup> that participate in nucleotide binding. The final amino acid in the Walker B motif, an aspartate that ligates the Mg<sup>2+</sup> cofactor of ATP, is the first amino acid in the DEAD sequence. The catalytic base is found at different sites in different functional families of F1-like ATPases but occurs at the same position adjacent to this aspartate in both the superfamily I and II helicases and the ABC transporters.

Studies of the structural mechanics of all of these F1-like mechanoenzymes suggest that their homologous motor domains generally function as ATP-dependent mechanical clamps.<sup>12,14,28,29,35,15,42–46</sup> ATP binding in the interfacial active site stabilizes the closed state of the clamp via the chemical contacts made by the nucleotide to both of the flanking domains. Opening of the clamp, which prevents the nucleotide from simultaneously interacting with both domains, lowers affinity for the bound nucleotide. Therefore, ATP binding generally closes the clamp, while hydrolysis weakens interfacial contacts to permit opening of the clamp. These conformational changes are controlled by the structural interactions in the ATPase active site at the clamp interface, which includes an extensive network of electrostatic interactions between the flanking protein domains that also connects these domains to the bound nucleotide. In the case of SecA, these interactions must ultimately control the functional conformational changes

throughout this 100 kDa ATPase. While interdomain interactions and protein–nucleotide interactions have been implicated in allosteric control of SecA,<sup>12,26,27</sup> substantial uncertainty remains concerning the pathway of conformational communication and the mechanism by which this communication is controlled by ATP binding and hydrolysis.

At room temperature in the absence of nucleotide, the DEAD-box motor domains of SecA adopt a closed conformation resembling that found in the nucleotide-bound structures of homologous mechanoenzymes.<sup>14</sup> However, at temperatures slightly above physiological, SecA undergoes a global conformational transition, called the endothermic conformational transition (ECT), that significantly reduces nucleotide-binding affinity.<sup>13,22,23,48,49</sup> (Note that this widely used term implies directionality from the low-temperature conformational ground state to the high-temperature excited state, even though this transition represents a reversible equilibrium.) Based on the structural paradigms observed in related mechanoenzymes, this conformational transition is believed to open the ATPase active site at the interface between the DEAD-box motor domains (as schematized in Figure 9 below), a hypothesis supported by mutagenesis studies.<sup>12</sup> These studies furthermore demonstrate that the ECT couples opening of the DEAD-box motor domains to global changes in interdomain interactions throughout SecA.<sup>12</sup> Nucleotide binding reverses the ECT, presumably via closure of the clamp formed by the DEAD-box motor domains.<sup>12,24</sup> Therefore, the ECT is a global conformational transition allosterically controlled by nucleotide binding at the ATPase active site.

Given these properties, the ECT could emulate the ATPdriven conformational changes that mediate preprotein translocation through SecYEG. This model is attractive because otherwise SecA would have to possess a second distinct ATPmodulated conformational reaction cycle. Temperatures above physiological promote tight membrane association of SecA in *vivo*, which is consistent with the ECT gating functional interaction with SecYEG during protein transport.<sup>9,18,50</sup> This observation suggests that SecA is intimately associated with SecYEG in the nucleotide-free state, which directly contradicts data showing that the nonhydrolyzable ATP analog AMP-PNP stabilizes interaction of SecA with the channel.<sup>18,51</sup> However, AMP-PNP has been shown to have poor affinity for the ATPase active site in SecA<sup>13,23</sup> and many other mechanically active ATPases.<sup>52</sup> In several cases, this analog has resulted in misleading mechanistic inferences because it lacks sufficient binding energy to induce the proper ATP-dependent conformational changes.53-55

To elucidate the stereochemical and mechanistic effects of ATP binding to SecA and to deepen understanding of the allosteric control of global conformation by its ATPase cycle, we sought to develop alternative methods to trap SecA in an ATP-bound conformation. In ABC transporters, mechanoenzymes sharing key ATP binding and hydrolysis motifs, this goal was achieved by mutagenesis of the catalytic glutamate to glutamine,<sup>43,44,50</sup> which generally blocks ATP hydrolysis but preserves ATP-binding affinity for proteins belonging to this ATPase family. In this paper, we report that the E-to-Q mutation in the active site of either *B. subtilis* SecA (E208Q-*Bs*SecA) or *E. coli* SecA (E210Q-*Ec*SecA) has dramatically different thermodynamic effects than the equivalent mutation in ABC transporter motors. The single point-charge change caused by this isosteric mutation significantly perturbs thermodynamic couplings throughout SecA, demonstrating that an exquisitely balanced network of electrostatic interactions<sup>33,34</sup> enables small changes in active site charge to alter the global conformational state of a 100 kDa mechanoenzyme.

## METHODS

**Buffers and Reagents.** Experiments were conducted in 50 mM KCl, 20  $\mu$ M EDTA, 1.0 mM MgCl<sub>2</sub>, 25 mM Tris-Cl, pH 7.6 (KEMT) or 50 mM Tris-HCl, pH 8.0, 50 mM KCl, 5 mM MgCl<sub>2</sub> (TKM)<sup>23</sup> buffers. Methylanthraniloyl (MANT)-labeled and unlabeled nucleotides were purchased from Invitrogen (Carlsbad, CA) and Sigma/ Fluka (St. Louis, MO), respectively; concentrations were determined using  $\varepsilon_{260} = 15400 \text{ M}^{-1} \cdot \text{cm}^{-1}$  for adenine and  $\varepsilon_{356} = 5800 \text{ M}^{-1} \cdot \text{cm}^{-1}$  for MANT.

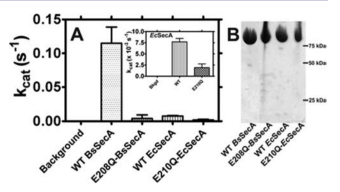
**Protein Biochemistry Methods.** Cloning and mutagenesis and protein expression, purification, and crystallization methods are described in the Supporting Information (SI). They are equivalent to those previously reported<sup>12,13</sup> except for use of an additional gel filtration column during purification. In brief, full-length proteins spanning the exact native sequences without additional residues were purified through DEAE sepharose, SP sepharose, butyl sepharose, and sephacryl S300 columns prior to concentration to 5–10 mg/mL in KEMT buffer for enzyme assays and fluorimetry, TKM buffer for calorimetry, or 300 mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 1 mM DTT, 20 mM BES, pH 7.0 for crystallization. Protein concentration was determined by absorbance using  $\varepsilon_{280} = 51\,480\,\,\mathrm{M^{-1}cm^{-1}}$  or 76 000  $\,\mathrm{M^{-1}cm^{-1}}$  for *Bs*SecA or *Ec*SecA, respectively, after dilution into 6 M guanidinium-HCl, 20 mM sodium phosphate, pH 6.5.

Assays Methods. ATPase and fluorescence anisotropy assays were conducted and analyzed as previously<sup>13</sup> described using the buffers indicated. Calorimetric methods are described in the SI. Samples for calorimetry were prepared by dialyzing 100–500  $\mu$ L of protein stock solution for >18 h against 1 L of TKM buffer in either 3.5 K or 10 K slide-a-lyzer dialysis cassettes (Pierce, Rockford, IL). Buffer was changed 3 times with each step lasting >3 h. The final dialysis buffer was used to dissolve solid Mg-ADP for use in calorimetric experiments. Amide <sup>1</sup>H/<sup>2</sup>H exchange rate measurements using mass spectrometry were conducted as described in Sharma et al.<sup>57</sup> with slight modifications described in the SI.

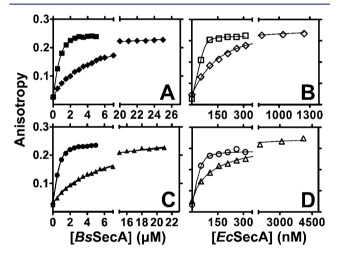
## RESULTS

E-to-Q Mutation in the Catalytic Base of SecA Impairs ATPase Activity. The basal ATPase activities of wild-type (WT) and mutant SecA enzymes from B. subtilis and E. coli were measured using the Malachite Green phosphate-release assay. Figure 1 shows that the basal turnover rate of WT-BsSecA ( $k_{cat} = 7.2 \pm 0.6 \text{ min}^{-1}$ ) is ~15-fold higher than that of WT-*EcSecA* ( $k_{cat} = 0.480 \pm 0.024 \text{ min}^{-1}$ ), consistent with previous reports.<sup>13,41</sup> In both enzymes, mutation of the catalytic base from glutamate to glutamine very strongly reduces the rate of ATP hydrolysis, to a level similar to background in comparable enzyme preparations ( $k_{cat} = 0.24 \pm 0.18 \text{ min}^{-1}$ for BsSecA and  $k_{cat} = 0.120 \pm 0.006 \text{ min}^{-1}$  for EcSecA), also consistent with previous work on the E. coli enzyme.<sup>41</sup> Note that these rates represent upper limits for the ATPase velocities of the mutant enzymes. Even though our three-column purification procedure yields highly pure enzymes (Figure 1B), the observed turnover rates of the mutants are so low that they could derive from trace contamination by an active ATPase (e.g., <0.1% by mass). It is extremely challenging to exclude this possibility with enzyme turnover rates on the order of 1 per 500 s.

**E-to-Q Mutation Reduces Nucleotide Affinity and Accelerates Binding and Release Rates.** We used fluorescence anisotropy assays to determine the effects of the E-to-Q mutation on the affinity (Figure 2) and kinetics (Figures 3 and 4) of SecA's interaction with MANT-labeled



**Figure 1.** ATP activity of SecA is reduced in the E-to-Q mutants. (A) ATP hydrolysis rates of WT and mutant SecAs were measured using the Malachite Green phosphate-release assay. Reactions at 25 °C containing  $1-2 \mu M$  SecA and 2 mM Mg-ATP in KEMT buffer were monitored at 10–15 min intervals for 30–90 min. The observed activity of E210Q-*EcS*ecA is so low that it could be attributable to contaminating enzymes despite the high protein purity (see text). (B) Coomasie-blue stained SDS-PAGE gel of purified SecA variants.



**Figure 2.** Nucleotide-binding affinity is reduced to a different extent in E208Q-*Bs*SecA vs E210Q-*Ec*SecA. SecA variants were titrated onto a fixed concentration of MANT-labeled nucleotide at 25 °C in KEMT buffer (pH 7.6), while MANT fluorescence anisotropy was monitored at 352 nm. (A) WT ( $\blacksquare$ ) and E208Q ( $\blacklozenge$ ) *Bs*SecA were titrated onto 900 nM Mg-MANT-ATP. (B) WT ( $\Box$ ) and E210Q ( $\diamondsuit$ ) *Ec*SecA were titrated onto 30 nM Mg-MANT-ATP. (C) WT ( $\blacklozenge$ ) and E208Q ( $\bigstar$ ) *Bs*SecA were titrated onto 900 nM Mg-MANT-ATP. (D) WT ( $\bigcirc$ ) and E210Q ( $\bigtriangleup$ ) *Bs*SecA were titrated onto 900 nM Mg-MANT-ATP. (D) WT ( $\bigcirc$ ) and E210Q ( $\bigtriangleup$ ) *Ec*SecA were titrated onto 30 nM Mg-MANT-ATP. (D) WT ( $\bigcirc$ ) and E210Q ( $\bigtriangleup$ ) *Ec*SecA were titrated onto 30 nM Mg-MANT-ADP. (D) WT ( $\bigcirc$ ) and E210Q ( $\bigtriangleup$ ) *Ec*SecA were titrated onto 30 nM Mg-MANT-ADP. (D) WT ( $\bigcirc$ ) and E210Q ( $\bigtriangleup$ ) *Ec*SecA were titrated onto 30 nM Mg-MANT-ADP. (D) WT ( $\bigcirc$ ) and E210Q ( $\bigtriangleup$ ) *Ec*SecA were titrated onto 30 nM Mg-MANT-ADP. (D) WT ( $\bigcirc$ ) and E210Q ( $\bigtriangleup$ ) *Ec*SecA were titrated onto 30 nM Mg-MANT-ADP. (D) WT ( $\bigcirc$ ) and E210Q ( $\bigtriangleup$ ) *Ec*SecA were titrated onto 30 nM Mg-MANT-ADP. (D) WT ( $\bigcirc$ ) and E210Q ( $\bigtriangleup$ ) *Ec*SecA were titrated onto 30 nM Mg-MANT-ADP. (D) WT ( $\bigcirc$ ) and E210Q ( $\bigstar$ ) *Ec*SecA were titrated onto 30 nM Mg-MANT-ADP.

ADP and ATP analogues at 25 °C<sup>13</sup> (Table 1). Previous research has shown that MANT-labeled analogues closely track the binding properties of unlabeled adenine nucleotides,<sup>13</sup> although they bind with ~5-fold higher affinity to *EcS*ecA. Figure 2 shows that the E-to-Q mutation reduces the affinity for Mg-MANT-ADP by ~20-fold in *Bs*SecA (increasing the  $K_d$ from 160 ± 40 nM to 3.50 ± 0.14  $\mu$ M) and by ~10-fold in *EcS*ecA (increasing the  $K_d$  from 12 ± 3 to 120 ± 11 nM). The somewhat more severe defect in *Bs*SecA compared to *EcS*ecA suggests that the influence of the E-to-Q mutation may differ in some details between the enzymes even though it produces a similar biochemical phenotype.

Figure 3 shows that the E-to-Q mutation in both enzymes produces substantially larger increases in the rate of Mg-MANT-ADP release ( $k_{off}$ ) than in its binding affinity. The mutant enzymes release this nucleotide so rapidly that that it is

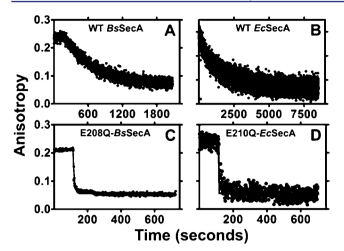
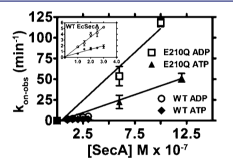


Figure 3. E-to-Q mutants of SecA do not kinetically trap ATP. Competitive displacement of MANT-labeled nucleotides by unlabeled nucleotides was monitored kinetically using fluorescence anisotropy spectroscopy at 352 nm in KEMT buffer. (A) WT-BsSecA, (B) WT-EcSecA, (C) E208Q-BsSecA, or (D) E210Q-EcSecA at 250–750 nM was equilibrated with 60 nM Mg-MANT-ATP in KEMT buffer (pH 7.6) at 5 °C for 3–5 min before adding a >100-fold excess of unlabeled Mg-ADP to initiate the exchange reaction.



**Figure 4.** Nucleotides bind to the E-to-Q mutants faster than to WT SecA. WT or E210Q *EcS*ecA were added to a cuvette containing 60 nM MANT-labeled nucleotide in KEMT buffer (pH 7.6) at 25 °C, and binding was monitored using fluorescence anisotropy spectroscopy at 352 nm. The on-rate ( $k_{on}$ ) was calculated from the observed single-exponential binding curves as explained in the SI. The labels ADP and ATP refer to Mg-MANT-ADP and Mg-MANT-ATP, respectively. The inset shows a magnified view of the data for the WT enzyme.

difficult to quantify the exact rate at 25 °C, which means that it must be at least 100-fold faster than in the WT enzymes given the configuration of our experimental apparatus. (At 5 °C, a temperature at which nucleotide-exchange kinetics are substantially slower due to factors explained below, the mutation increases  $k_{\text{off}}$  by 270-fold in BsSecA and 140-fold in EcSecA, as shown in Table 1.) Previous research<sup>13</sup> has demonstrated that the ratio of the measured off-rate for Mg-MANT-ADP release to the measured on-rate for its binding accurately recapitulates its  $K_d$  (i.e.,  $k_{off}/k_{on} = K_d$ ), indicating that binding is a well-behaved two-state process that is accurately monitored by the anisotropy of the fluorescently labeled nucleotide. Therefore, the disproportionately large increase in  $k_{\rm off}$  compared to  $K_{\rm d}$  for Mg-MANT-ADP indicates that the mutation must accelerate its rate of binding in addition to its rate of release.

This inference is confirmed by direct measurements of the rate of Mg-MANT-ADP binding to the WT and mutant enzymes. While the rate was difficult to quantify accurately for E208Q-*Bs*SecA due to its high velocity, it is clearly accelerated compared to the WT enzyme (data not shown). The rate of Mg-MANT-ADP binding to both WT and E208Q-*Ec*SecA could be quantified (Figure 4 and Table 1), showing that the mutation increases  $k_{on}$  by 6.6-fold (from  $18.0 \pm 0.9 \times 10^6$  to  $120.0 \pm 0.9 \times 10^6$  M<sup>-1</sup> min<sup>-1</sup>). The measured values of the on-rates match those expected from the measured values of  $K_d$  and  $k_{off}$  for both WT and mutant enzymes (i.e.,  $k_{off}/k_{on} = -K_d$  for the measured values), supporting the accuracy of our kinetic measurements. The observed increases in both  $k_{on}$  and  $k_{off}$  in both enzymes suggest that the E-to-Q mutation substantially accelerates opening of the interface between NBF-I and NBF-II, which forms the ATPase active site.<sup>12,13</sup>

Equivalent affinity (Figure 2) and kinetic (Figures 3 and 4) measurements can be done using Mg-MANT-ATP. Previously reported measurements of pre-steady-state kinetics have demonstrated that ATP is hydrolyzed very rapidly after binding to WT SecA, while release of the ADP product occurs much more slowly<sup>41</sup> (and no rapidly released population is observed in kinetic experiments, ruling out nonproductive binding of ATP). Therefore, binding experiments conducted on WT enzymes with hydrolyzable ATP species report a combination of ATP binding and ADP release rates. However, equivalent measurements on the ATPase-deficient E-to-Q mutants yield accurate estimates of binding affinity because these enzymes release ATP substantially more rapidly than they hydrolyze it (Table 1). Both affinity and kinetic measurements with Mg-MANT-ATP show equivalent trends to those observed for Mg-MANT-ADP (Table 1), including a strong acceleration in binding rate and a net reduction in apparent binding affinity. The ATP analogue binds to both WT and E-to-Q variants of EcSecA and BsSecA ~3-fold slower than the ADP analogue. While there is a strong reduction in ATP-binding affinity in both mutant enzymes, there is again a larger reduction in BsSecA compared to EcSecA (~20- vs ~5-fold), reinforcing the conclusion that the exact influence of the mutation may differ between the enzymes.

These results contrast sharply with those observed for the stereochemically equivalent mutation in the active sites of a wide variety of ABC transporter motors, which consistently kinetically trap ATP in the presence of the E-to-Q mutation (i.e., release it exceedingly slowly).43,45 The equivalent mutation in SecA clearly does not kinetically trap ATP and instead substantially accelerates its release rate, resulting in a large reduction in net binding affinity. Furthermore, introducing the equivalent mutation into MJ0669, a model DEAD-box helicase from Methanococcus jannaschii, similarly fails to kinetically trap ATP (Figure S1). Therefore, even though the motor domains of ABC transporters and ATP-dependent helicases share homology and even though both function as ATP-dependent mechanical clamps, there appear to be major differences in the role of electrostatic interactions in controlling protein conformation or dynamics.<sup>33,34</sup> Therefore, we determined the crystal structure of E208Q-BsSecA to examine whether the mutation perturbs active-site stereochemistry.

**Perturbations in Active-Site Stereochemistry Are Unlikely to Account for Reduced Nucleotide-Binding Affinity.** The nucleotide-free *B. subtilis* mutant protein crystallized isomorphously with the previously reported structure of the WT enzyme, indicating no significant change in interdomain interactions when the protein is packed in the crystal lattice. The structure of the mutant crystal was solved using molecular replacement and refined to a free *R* factor of

Table 1. Kinetic and Therm	odynamic Parameters of Nucleotide	Interactions with WT and Mutant SecA
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parameter	ligand	method	temp.	WT BsSecA	BsSecA E208Q	WT EcSecA	EcSecAE210Q
n	Mg-ADP	ITC	25 °C	0.89	0.92	0.93	_
$\Delta H^\circ$ (kcal mol <sup>-1</sup> )	Mg-ADP	ITC	25 °C	$-13.00 \pm 0.07$	$-14.0 \pm 0.1$	$-20.0 \pm 0.1$	_
$\Delta S^{\circ} \; (\text{kcal mol}^{-1})^a$	Mg-ADP	ITC	25 °C	$-13.0 \pm 0.3$	$-24.0 \pm 0.3$	$-34.0 \pm 0.4$	_
$\Delta G^\circ$ (kcal mol $^{-1}$ )	Mg-ADP	ITC	25 °C	$-9.20 \pm 0.04$	$-6.90 \pm 0.02$	$-9.80 \pm 0.04$	_
$K_{\rm d}$ (nM)	Mg-ADP	ITC	25 °C	$180 \pm 10$	$7700 \pm 300$	62 ± 4	_
$k_{\rm cat} \ ({\rm min}^{-1})^b$	Mg-ATP	$MG^{c}$	25 °C	$7.2 \pm 0.6$	$0.24 \pm 0.18^{d}$	$0.480 \pm 0.024$	$0.120 \pm 0.006^d$
$K_{\rm d}$ (nM)	Mg-MANT-ADP	anisotropy	25 °C	$160 \pm 40^{e}$	$3500 \pm 140$	$12 \pm 3$	$120 \pm 11$
$K_{\rm d}$ (nM)	Mg-MANT-ATP	anisotropy	25 °C	$190 \pm 50^{f}$	$3800 \pm 120$	$22 \pm 7^{f}$	$120 \pm 9$
$K_{\rm d}$ (nM)	Mg-MANT-ADP	anisotropy	5 °C	$250 \pm 30$	$460 \pm 10$	$17 \pm 7$	16 ± 9
$k_{\rm off}  ({\rm min}^{-1})$	Mg-MANT-ATP	anisotropy	5 °C	$0.120 \pm 0.003^{f}$	$32 \pm 3$	$0.0400 \pm 0.0004^{f}$	$5.6 \pm 0.6$
$k_{\rm off}  ({\rm min}^{-1})$	Mg-MANT-ADP	anisotropy	25 °C	$1.1 \pm 0.1$	_	$0.070 \pm 0.003$	_
$k_{\rm off}  ({\rm min}^{-1})$	Mg-MANT-ATP	anisotropy	25 °C	$0.80 \pm 0.03^{f}$	_	$0.0500 \pm 0.0007^{f}$	_
$k_{\rm on} \ ({\rm M}^{-1} \ {\rm min}^{-1})$	Mg-MANT-ADP	anisotropy	25 °C	$11.0 \pm 0.9 \times 10^{6}$	-	$18.0 \pm 0.9 \times 10^{6}$	$120.0 \pm 0.9 \times 10^{6}$
$k_{on} (M^{-1} min^{-1})$	Mg-MANT-ATP	anisotropy	25 °C	$5.0 \pm 0.6 \times 10^{6}$	-	$6.0 \pm 0.6 \times 10^{6}$	$40.0 \pm 0.4 \times 10^{6}$
$K_{\rm d} = k_{\rm off} / k_{\rm on} \ ({\rm nM})$	Mg-MANT-ADP	anisotropy	25 °C	$100 \pm 20$	-	$4.0 \pm 0.4$	-
$K_{\rm d} = k_{\rm off}/k_{\rm on} \ ({\rm nM})$	Mg-MANT-ATP	anisotropy	25 °C	$150 \pm 20^{f}$	_	$8 \pm l^{f}$	_
$T_{\rm m}$ (°C)	none	DSC	N/A	$40.0 \pm 0.1, 49.0 \pm 0.1$	$27.0 \pm 0.1, 41.0 \pm 0.0$	$\begin{array}{c} 41.00 \pm 0.03, \\ 44.00 \pm 0.05 \end{array}$	$39.0 \pm 0.1, \\ 44.0 \pm 0.1$
$T_{\rm m}$ (°C)	Mg-ADP	DSC	N/A	$47.0 \pm 0.4, 51.0 \pm 0.4$	$32.00 \pm 0.03,$ $41.00 \pm 0.03$	$47.00 \pm 0.03$	$41.0 \pm 0.1,$ $44.0 \pm 0.1$

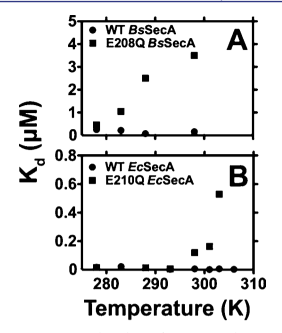
<sup>*a*</sup>Calculated by the ORIGIN ITC module from  $\Delta G^{\circ} = \Delta H^0 - T\Delta S^{\circ}$ , with  $\Delta G^{\circ}$  being calculated from the optimized value of  $K_d$  from curve-fitting the ITC data. <sup>*b*</sup> Previous research has shown that  $k_{cat}$  is equal to  $k_{off}$  for Mg-ADP release, <sup>13</sup> meaning that Mg-ADP release is rate limiting in the basal ATP hydrolysis cycle of SecA. The  $k_{cat}$  values reported here for Mg-ADP do not match the  $k_{off}$  values reported below for Mg-MANT-ADP because the fluorescently derivatized nucleotide binds to SecA tighter and is released more slowly than Mg-ADP itself. <sup>*c*</sup>Measured via Malachite Green phosphate-release assays calibrated with inorganic phosphate. <sup>*d*</sup>This level of ATP turnover may reflect the activity of contaminating enzymes; see text. <sup>*c*</sup>This value was obtained using 900 nM [Mg-MANT-ADP]. WT *Bs*SecA showed evidence of a modest reduction in  $K_d$  value in experiments conducted at substantially lower nucleotide and therefore protein concentrations, suggesting that dissociation of the *Bs*SecA dimer may increase nucleotide-binding affinity slightly. <sup>*f*</sup>Mg-MANT-ATP affinities and off-rates for the WT enzymes reflect the release rates of Mg-MANT-ADP.

26.7% at 3.4 Å resolution (Table S1 and Figure S2A). Electron density in the active site, which is the most stably ordered region of the structure, was clearly interpretable despite the limited resolution of the diffraction data (data not shown). While local stereochemistry is mostly preserved, the side chains of Arg-489, Arg-517, and Arg-525 show shifts in conformation resulting in altered hydrogen-bonding (H-bonding) interactions in the active site in the presence of the E-to-Q mutation (Figure S2A). Disruption or rearrangement of cooperative H-bonding interactions involving these Arg residues, which are all directly or indirectly involved in nucleotide binding, could potentially interfere with nucleotide binding.

To evaluate this possibility, we mutated Arg-489 to lysine<sup>51,58</sup> in WT-BsSecA and E208Q-BsSecA to change the stereochemistry of this residue's H-bonding interactions while preserving net charge in the active site. The crystal structure of E208Q-R489K-BsSecA was solved and refined to a free *R* factor of 26.9% at 3.3 Å resolution (Figure S2B and Table S1). It shows no electron density for the side chain of residue 489, indicating conformational disorder and confirming that the R489K mutation changes cooperative H-bonding interactions in the active site. However, this mutation does not significantly perturb Mg-MANT-ADP binding affinity in either the WT or the E208Q background (Figure S3A), showing that the detailed H-bonding interactions of Arg-489 are not an important determinant of nucleotide binding energy.

Active-Site Stereochemistry Controls Nucleotide-Release Rate. We evaluated next the influence of net electrostatic charge in the active site on nucleotide-binding affinity and kinetics, focusing on reversal of the charge difference caused by the E-to-Q mutation. Ser-211 is spatially adjacent to the catalytic glutamate E208 in the active site of BsSecA. Therefore, Ser-211 was mutated to aspartate or glutamate in both WT and E208Q-BsSecA. These mutations, which modestly reduce Mg-MANT-ADP binding affinity in the WT background and modestly increase it in the presence of the E208Q mutation (data not shown), greatly accelerate Mg-MANT-ADP release rate in the WT background (Figure S3B,D) and fail to reverse the accelerated release rate in the presence of the E208Q mutation (Figure S3C,E). These experiments varying active-site electrostatic charge demonstrate that active-site stereochemistry is a much more important determinant of nucleotide release rate than net electrostatic interaction with the nucleotide. This rate was previously inferred to depend on global conformational fluctuations in SecA based on the parallel effects of temperature changes on release rate and protein conformation.<sup>13</sup> Taken together, these observations suggest that perturbations in active-site stereochemistry, like those produced by the E-to-Q mutation, might change global conformational couplings within SecA.

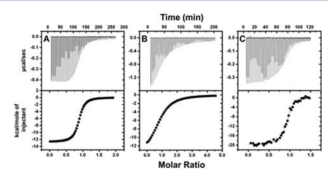
**Mg-MANT-ADP Binding To the E-to-Q Mutants Is Temperature Dependent.** SecA is known to undergo an ECT at physiological temperatures. This conformational change reduces nucleotide-binding affinity and accelerates nucleotide-release rate.<sup>13,59</sup> Facilitation of this transition would be expected to reduce nucleotide-binding affinity, so such an effect could contribute to the defects in nucleotide binding observed in the E-to-Q mutants. Therefore, we measured the Mg-MANT-ADP binding affinities at temperatures from 5 to 35 °C for WT and E-to-Q mutants of *B. subtilis* and *E. coli* SecA (Figure 5). The E-to-Q mutants both show dramatic changes in nucleotide-binding affinity in this temperature range, while the WT proteins show no significant variation except for a very slight increase in  $K_d > 33$  °C due to



**Figure 5.** Temperature dependence of  $K_d$  in WT and E-to-Q SecA. The Mg-MANT-ADP-binding affinity of WT ( $\bullet$ ) or mutant ( $\blacksquare$ ) variants of BsSecA (panel A) or EcSecA (panel B) were measured at temperatures from 5 to 30 °C in KEMT buffer (pH 7.6) using the fluorescence anisotropy assay shown in Figure 2 with 900 nM Mg-MANT-ADP.

an increasing population of protein in the domain-dissociated, low-affinity conformation produced by the ECT. Most strikingly, at 5 °C, E208Q-BsSecA binds Mg-MANT-ADP with comparable low-nanomolar affinity to the WT enzyme, so reducing temperature completely suppresses the defect in Mg-MANT-ADP-binding affinity. However, the  $K_d$  of the mutant begins to increase around 15 °C and is in the micromolar range at 25 °C. E210Q-EcSecA shows a qualitatively similar dependence of  $K_d$  on temperature, although the  $K_d$  only begins to increase above 20 °C. These data are consistent with the affinity measurements presented above showing a greater reduction in binding affinity for E208Q-BsSecA compared to E210Q-EcSecA at 25 °C. Moreover, the observed temperature dependence of Mg-MANT-ADP-binding affinity supports the hypothesis that the E-to-Q mutation changes global conformational couplings within SecA and facilitates the ECT. Specifically, this dependence suggests that the mutation lowers the temperature at which the ECT occurs and that it thereby produces a reduction in Mg-MANT-ADP-binding affinity at temperatures at which the mutant enzyme has undergone the ECT while the WT enzyme remains in the conformational ground state and retains maximal binding affinity (Figure 5). To evaluate this hypothesis more directly, we undertook calorimetric studies of nucleotide binding and enzyme conformation.

**ITC Shows Mg-ADP Binding Is Coupled to a Larger Entropy Loss in E208Q-BsSecA.** We used isothermal titration calorimetry (ITC) to measure the enthalpy and entropy of Mg-ADP binding to WT-BsSecA, E208Q-BsSecA, and WT-EcSecA at 25 °C (Figure 6 and Table 1). We were unable to obtain interpretable data for E210Q-EcSecA because this protein displayed irreproducible behavior in ITC experiments (as documented in Figure S4). However, high-quality ITC data were obtained for WT-EcSecA as well as both WT and E208Q-BsSecA. WT-EcSecA was reported to have ~5-fold



**Figure 6.** ITC reveals that large entropy losses accompany nucleotide binding to E208Q-*Bs*SecA and WT-*Ec*SecA. ITC experiments were performed at 25 °C in TKM buffer (pH 8.0). Mg-ADP was titrated onto 20  $\mu$ M WT-*Bs*SecA (panel A), 30  $\mu$ M E208Q-*Bs*SecA (panel B), or 10  $\mu$ M WT-*Ec*SecA (panel C). Table 1 gives the thermodynamic parameters inferred from curve-fitting these data.

lower affinity for Mg-ADP compared to the MANT-labeled analog based on competitive fluorescence binding experiments.<sup>13</sup> Our ITC data confirm this inference for *EcS*ecA and demonstrate that Mg-ADP binds to WT *BsSecA* with roughly comparable affinity either with or without MANT derivatization. Underivatized Mg-ADP binds to BsSecA ~3-fold more weakly than to EcSecA (180 ± 10 vs 62 ± 4 nM), but this difference is physiologically insignificant given the fact that cytosolic Mg-ADP concentration is at least 1000-fold greater than the higher of these two  $K_d$  values and cytosolic Mg-ATP concentration is even higher in growing bacterial cells.<sup>60</sup>

The detailed thermodynamic data generated by the ITC experiments (Figure 6 and Table 1) demonstrate that there is a much higher entropy loss during binding of Mg-ADP to E208Q-BsSecA compared to the WT enzyme at 25 °C ( $\Delta S^{\circ}$  of  $-24 \text{ vs} - 13 \text{ cal mol}^{-1} \text{ K}^{-1}$ ), while there is a comparatively small change in binding enthalpy  $(-14 \text{ vs} - 13 \text{ kcal mol}^{-1})$ . Thus, the E208Q mutant modestly increases the enthalpy gain upon binding, perhaps due to reduced electrostatic repulsion of the phosphates of ADP, while it doubles the unfavorable entropy of binding. The most straightforward explanation for this entropic effect, which accounts for the strong reduction in binding affinity produced by the E208Q mutation in BsSecA, is that the mutant enzyme is more disordered than the WT enzyme in the apo conformation while it is not yet underway in the Mg-ADPbound conformation. This explanation is consistent with the inference made above based on the temperature dependence of Mg-MANT-ADP-binding affinity that the E208Q mutation facilitates the ECT, causing it to be underway at 25 °C in the mutant enzyme while it is not yet underway in the WT enzyme. The conformation produced by this transition, which greatly reduces Mg-ADP-binding affinity, is believed to be more dynamic than the low-temperature ground-state conformation.<sup>12,26</sup> Therefore, because reversal of the ECT must precede Mg-ADP binding, the entropy loss involved in reversing the transition is coupled to Mg-ADP binding to E208Q-BsSecA. This entropy loss accounts for the lower net affinity of binding to E208Q-BsSecA compared to WT BsSecA, because the WT enzyme is in the conformational ground state at 25 °C and the ECT does not have to be reversed before it can bind Mg-ADP at this temperature.

The ITC experiments demonstrate that Mg-ADP binding to WT-*EcS*ecA at 25 °C involves a substantially larger entropy loss  $(-34 \text{ cal mol}^{-1} \text{ K}^{-1})$  and a substantially larger enthalpy gain  $(-20 \text{ kcal mol}^{-1})$  than binding to WT or even E208Q *Bs*SecA

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(Figure 6 and Table 1). These large entropy and enthalpy changes almost entirely compensate for one another, because the net binding affinities of the two enzymes differ by only  $\sim 10$ fold (Table 1). The ITC experiments were conducted at a temperature well below the onset of the ECT in the apo form of either WT enzyme (see below), so the high entropic cost of Mg-ADP binding to *EcSecA* cannot come from coupling to the reversal of the ECT. It must reflect some other molecular process involving a reduction in dynamics, either a reduction in protein dynamics in the low-temperature ground-state conformation of the enzyme and/or a change in solvent dynamics upon nucleotide binding. Both processes typically involve strong enthalpy-entropy compensation of the kind observed in comparing Mg-ADP-binding to EcSecA vs BsSecA. The process responsible for the much larger entropy loss upon Mg-ADP binding to EcSecA must be either absent or greatly reduced in magnitude in BsSecA, making it unlikely to be an essential feature of the mechanochemistry of the enzyme. NMR studies have suggested that helicase motifs V and VI in the ATPase active site have elevated dynamics in apo EcSecA,<sup>26,61</sup> and immobilization of these segments during Mg-ADP binding to EcSecA could potentially account for the larger entropy loss observed during binding to this enzyme compared to BsSecA (Table 1). However, data presented below demonstrate that both of these segments adopt predominantly folded conformations in apo EcSecA at 25 °C on the time scale of amide hydrogen-deuterium exchange (HDX), making it unlikely that a simple unfolding transition in motifs V or VI is responsible for the much larger entropy loss upon Mg-ADP binding to EcSecA vs BsSecA.

E208Q Mutation in BsSecA Destabilizes the ECT by 13 °C. We used thermal titrations monitored by intrinsic tryptophan fluorescence spectroscopy<sup>12</sup> (Figure S5) and differential scanning calorimetry (DSC; Figures 7 and S6) to verify the hypothesis that the E-to-Q mutation facilitates the ECT in BsSecA. Previous research<sup>23,24</sup> demonstrated that WT-BsSecA undergoes two endothermic transitions between 20 and 60 °C. The first, with a midpoint of 40 °C, represents the ECT and is fully reversible, while the second, with a midpoint at 49 °C, involves the irreversible unfolding of some protein domain, probably NBF-I.<sup>23,24</sup> Addition of Mg-ADP leads to a progressive increase in the temperature of the first transition due to inhibition of the ECT by nucleotide binding<sup>23</sup> (i.e., to 47 °C in the presence of sufficient Mg-ADP to saturate the active site). Our DSC data on WT-BsSecA recapitulate these earlier observations (Figure 7A,B). Strikingly, as predicted based on the temperature-dependence of Mg-MANT-ADP-binding affinity, DSC analysis shows that the ECT is shifted to lower temperature in E208Q-BsSecA (Figure 7C). The observed 13° downshift to 27 °C reflects a dramatic alteration in the global conformational properties of the apo enzyme in this temperature range based on a single isosteric point mutation in the active site. The second endothermic transition in BsSecA is also shifted to lower temperature by  $\sim 8^{\circ}$ , consistent with the previous interpretation that this transition involves unfolding of NBF-I (i.e., the domain containing the E208Q mutation).

Addition of Mg-ADP to E208Q-*Bs*SecA results in an increase of 5 °C in the  $T_{\rm m}$  of the first transition (i.e., shifting the ECT to 32 °C) as well as an increase in its enthalpy (Figure 7D). However, both of these changes are substantially attenuated compared to those observed upon Mg-ADP binding to the WT enzyme, consistent with the data presented above showing that the E-to-Q mutation strongly perturbs nucleotide interactions

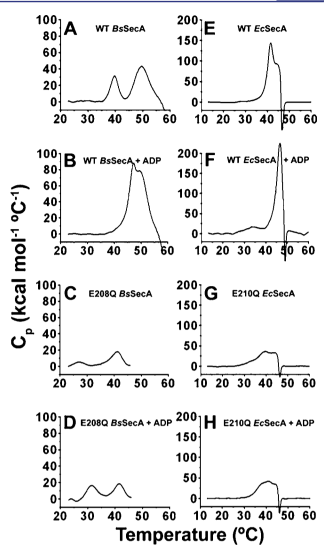


Figure 7. DSC of WT and E-to-Q BsSecA and EcSecA. All DSC scans were conducted on the same instrument at 1 °C per min in TKM buffer (pH 8.0). The same vertical scale is used here for all experiments conducted on enzymes from the same organism, while Figure S6 shows the same data replotted with the vertical scale adjusted to enable clear visualization of the details of the transitions in each experiment. (A,B) 10  $\mu$ M WT-BsSecA without (panel A) or with (panel B) 1 mM Mg-ADP. (C,D) 20 µM E208Q-BsSecA without (panel C) or with (panel D) 1 mM Mg-ADP. (E,F) 10 µM WT-EcSecA without (panel E) or with (panel F) 1 mM Mg-ADP. (G,H) 20 µM E210Q-EcSecA without (panel G) or with (panel H) 1 mM Mg-ADP. Protein aggregation following the second conformational transition, which is probably unfolding of NBF-I, likely contributes to the very large magnitude of  $C_p$  in some experiments. This phenomenon prevents reliable integration of enthalpy. However, it does not interfere with observation of the temperature and approximate magnitude of the initial transition corresponding to the ECT. The data shown here for WT BsSecA accurately recapitulate previously published results.<sup>23,24</sup>

at the high-affinity ATPase site in BsSecA. The  $T_{\rm m}$  of the second transition in E208Q-BsSecA is essentially unchanged by the addition of Mg-ADP; this observation provides further evidence that nucleotide release effectively accompanies the ECT, because residual nucleotide binding to the Walker motifs in NBF-I after the occurrence of the ECT would be expected to inhibit its unfolding during the second transition and thereby increase the  $T_{\rm m}$  of this transition.

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DSC analysis of EcSecA in the absence and presence of the E210Q mutation shows qualitatively similar trends (Figure 7E-H). The mutation reduces the  $T_{\rm m}$  of its ECT and substantially attenuates the increase in the  $T_{\rm m}$  and enthalpy of this transition upon Mg-ADP binding (Figure 7G,H vs E,F). Notably, the single endothermic transition in WT-EcSecA is split into at least two sequential transitions by the E210Q mutation. This split makes the calorimetric behavior of E210Q-EcSecA more similar to that of WT BsSecA, although a broad shoulder on the lowtemperature side of the first transition in E210Q-EcSecA indicates that several sequential endothermic subtransitions may contribute to the ECT in this mutant enzyme. Critically, the onset of the earliest thermal transition in apo E210Q-EcSecA occurs at ~30 °C (Figure 7G) compared to ~20 °C in apo E208Q-BsSecA (Figure 7C). Therefore, while the E-to-Q mutation facilitates the ECT in both enzymes, it occurs at a significantly lower temperature when the mutation is introduced into BsSecA compared to EcSecA. This quantitative difference explains the substantially more severe impairment in the affinity of E208Q-BsSecA vs E210Q-EcSecA in the initial nucleotide-binding experiments reported above, which were conducted at 25 °C. At this temperature, the ECT is in progress in E208Q-BsSecA but not in E210Q-EcSecA. Therefore, the qualitative effects of the E-to-Q mutation, which are fundamentally similar in both enzymes, were masked by the quantitative differences in their thermodynamics properties. The next section presents further evidence supporting the similarity of the functional conformational changes and the qualitative effects of the E-to-Q mutation in the catalytic base in both enzymes.

HDX Experiments to Map the Effect of the ECT on **BsSecA Dynamics.** The DSC experiments reported above provide direct evidence showing that the E-to-Q mutation in the catalytic base of SecA perturbs the thermodynamics of the ECT. The observation that a single isosteric point mutation triggers the ECT in the apo enzyme at 25 °C provided a unique opportunity to use HDX measurements at a single temperature to characterize the global conformational changes in SecA during this transition, which is likely to be the principal nucleotide-controlled process mediating polypeptide transport. HDX provides information on protein dynamics by quantitatively measuring the rate of exchange of backbone and side chain amide hydrogen atoms for deuterium atoms from water.<sup>62,63</sup> Because chemical exchange requires exposure of the amide group to solvent, the method is sensitive to conformational fluctuations breaking backbone H-bonds (e.g., in regular protein secondary structures). Backbone HDX proceeds via two mechanisms: "EX1" involving transient conformational fluctuations exposing the backbone amides to solvent and "EX2" involving an enhanced population of a locally or globally unfolded protein conformation.<sup>62,63</sup> An increase in HDX rate via either mechanism requires an increase in net backbone dynamics because even the EX2 mechanism involves substantial fractional residency in an unfolded conformation on the millisecond time scale of the HDX process. Therefore, we used mass spectrometric HDX measurements<sup>63</sup> to probe the conformational state of the BsSecA variants at 10 and 100 s at 25 °C, where the WT enzyme is in the ground-state conformation but the E208Q enzyme is undergoing the ECT. The data from these experiments are presented in Figure 8A,B and SI, while the key conclusions are summarized schematically in Figure 9.



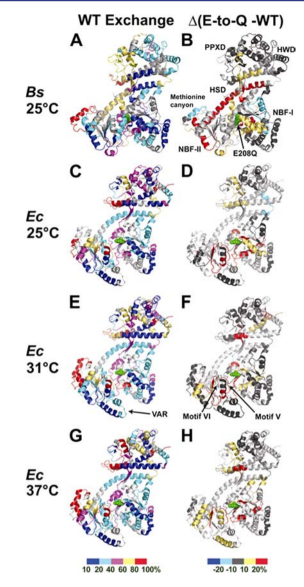
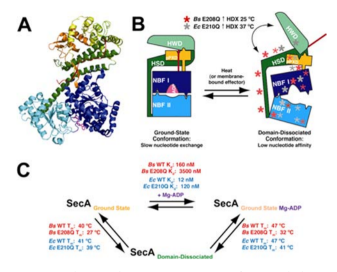


Figure 8. Summary of results from HDX-MS experiments. Average peptide <sup>1</sup>H/<sup>2</sup>H exchange levels at the 10 s time point in TKM buffer (pH 8.0) are plotted onto ribbon diagrams of the corresponding WT enzymes (PDB id 1M74 for BsSecA and PDB id 2FSG for EcSecA). The catalytic glutamates are shown in green sphere representation. (A,C,E,G) Absolute percent exchange of observable protons in peptides in the WT enzyme. (B,D,F,H) Difference in this value in the E-to-Q mutant minus that in the corresponding WT enzyme at the same time point; these latter images highlight the regions in which backbone dynamics are altered by the mutation. The exchange levels are color coded as indicated in the bar at the bottom of each column (10-20% exchange, blue; 21-40%, cyan; 41-60%, magenta; 61-80%, yellow; and 81-100%, red for the absolute exchange levels shown in the left column, and  $\geq 21\%$  reduction in exchange, blue; 20-11% reduction, cyan; 10% reduction to 10% increase; dark gray; 11-20% increase, yellow; and  $\geq 21\%$  increase in red for the differences in exchange levels shown in the right column). Experiments were conducted on BsSecA at 25 °C (A,B), where the DSC data (Figure 7A,C) indicate that the WT enzyme is in the conformational ground state but the E208Q mutant is undergoing the ECT. Experiments were conducted on EcSecA at 25 °C (C,D), 31 °C (E,F), and 37 °C (G,H). The DSC data on WT-EcSecA (Figure 7E) indicate that this enzyme is in the conformational ground state at all three temperatures, while the fine structure in the broad DSC peak for the E210Q mutant (Figure 7G) suggests that it sequentially occupies different conformational substates as it undergoes the ECT from 30 to 40 °C.



**Figure 9.** Schematic diagram summarizing conformational changes during the ECT and related features of the HDX data. (A) A ribbon diagram of the *Bs*SecA crystal structure (1M6N) bound to Mg-ADP (magenta) shows each domain colored separately to correspond to the labeled domains in panel B. The conformational transition (ECT) between the ground and the domain-dissociated states of SecA is schematized in panel B. The domain-dissociated state is marked with asterisks representing regions of increased mobility as measured by HDX in SecA mutants compared to WT. Increases in *Bs*SecA E208Q HDX are depicted in red, and increases in *EcS*ecA E210Q HDX are shown in gray. Bold asterisks represent regions with large changes in HDX rates. (C) Schematic diagram showing equilibria between the major conformational states of SecA, labeled with associated thermodynamic parameters.

In the absence of nucleotide, the WT protein shows modestly elevated amide proton exchange rates in the HWD and in local segments of the NBFs, more elevated dynamics in the PPXD and in the long central  $\alpha$ -helix in the HSD, and highly elevated dynamics in the methionine canyon in NBF-II (Figure 8A). The E208Q mutation strongly modulates amide proton exchange rates at sites throughout the protein, especially at interdomain interfaces (Figures 8A,B and 9A,B). The most dramatic change involves strong acceleration of HDX rate throughout the long central  $\alpha$ -helix in the HSD, indicating substantially greater backbone solvent exposure in this segment that serves as a physical scaffold organizing the interdomain interactions in the conformational ground state of SecA. As explained above, this observation indicates that the E208Q mutation increases backbone dynamics in this protein segment, either via enhanced conformational fluctuation or enhanced population of an unfolded conformation. The mutation produces similarly strong enhancements in HDX rates in some segments near the mutation site in NBF-I, i.e., in one  $\beta$ strand connecting this domain to the PPXD and in the first  $\alpha$ helix in NBF-II, which lies on the opposite surface of this domain from the ATP-binding site. The mutation also modestly enhances HDX rates elsewhere in the HSD and in the interface between NBF-I and NBF-II. Therefore, the HDX data verify inferences from earlier mutational<sup>12,49,64</sup> and NMR<sup>26</sup> analyses that the ECT involves a global conformational change increasing the solvent exposure and net backbone dynamics of interdomain interfaces throughout SecA, as summarized in Figure 9. The observation that this transition is induced by an isosteric glutamine-to-glutamate mutation demonstrates an

exquisitely sensitive coupling of active-site electrostatics<sup>34</sup> to global protein conformation.

While the E208Q mutation accelerates HDX rate in most regions of SecA, it modestly reduces the rate in the methionine canyon and on the distal surface of NBF-I relative to the ATPase active site. The observation of increased order at these sites suggests that they may interact with protein binding partners or other ligands specifically in the conformational state produced by the ECT. The coupling of interactions stabilizing local structure at these sites to a global conformational change generally increasing disorder elsewhere would enable such interactions to trigger the ECT and thereby coordinate the efficient progression of the conformational reaction cycle during the polypeptide transport reaction.

Equivalent Dynamic Changes in E210Q-EcSecA at Slightly Higher Temperature. Qualitatively similar results to those obtained with BsSecA are observed in HDX experiments performed on EcSecA (Figure 8C–H), suggesting that the fundamental conformational phenomena underlying SecA function are conserved between species even if the energetic consequences of the E-to-Q mutation in the catalytic base are slightly different. At 25, 31, and 37 °C, WT-EcSecA shows a similar overall HDX pattern as WT-BsSecA at 25 °C, although segments in the PPXD and HWD show some acceleration in HDX rate as temperature is raised (Figure 8C vs 8E vs 8G). These observations are consistent with the protein adopting an equivalent conformation until the temperature nears the onset of the ECT, which has a midpoint of ~41 °C in EcSecA (Figure 7E).

Although a prior NMR study showed enhanced mobility of the residues in helicase motifs V and VI in EcSecA in the absence of nucleotide,  $^{26,61}$  the HDX data support both of these motifs adopting stably folded conformations at 25 °C on the time scale of the amide-exchange process (Figure 8C and SI). Two peptides are observed that include motif V (residues 501-512),<sup>65</sup> one covering residues 500–512 and the other covering residues 502-512. These peptides respectively show ~60-80% exchange at 10 s and ~80-100% exchange at 100 s, which is similar to the levels observed for the equivalent peptide in BsSecA at 25 °C (50-60% at 10 s; 60-80% at 100 s). However, motif V adopts an extended conformation in which most residues do not participate in backbone H-bonds, so only a modest degree of protection from HDX is expected even in the folded structure. Two peptides are also observed that include motif VI (residues 563-579),<sup>65</sup> one covering residues 550-586 and the other covering residues 572-586. The longer peptide, which includes the entire motif, shows  $\sim 25\%$  exchange at 10 s and  $\sim$ 30% exchange at 100 s, while the shorter peptide shows  $\sim$ 35% exchange at 10 s and  $\sim$ 45% exchange at 100 s. These observed levels of exchange correspond roughly to the fraction of residues in these segments that do not participate in regular secondary structures and therefore would be expected to exchange even in the fully folded conformation. These HDX observations suggest that neither helicase motif V nor VI adopts a predominantly disordered conformation at 25 °C in WT EcSecA, although they do not exclude the possibility of transient unfolding in a minor fraction of the population.

HDX rates in E210Q-*EcS*ecA show progressively greater changes compared to the WT protein as temperature is raised from 25 °C (Figure 8D) to 37 °C (Figure 8H), consistent with the strong reduction in its nucleotide-binding affinity (Figure 5B) and the onset of the ECT (Figure 7G) in this temperature range in the mutant protein. While HDX data were not

obtained from E210Q-*EcS*ecA for the long central  $\alpha$ -helix in the HSD, enhanced exchange rates are observed in the regions of the structure with which it interacts, with stronger effects being observed at 31 °C (Figure 8F) and 37 °C (Figure 8H) where the protein is undergoing the ECT. These data demonstrate that net backbone dynamics are enhanced at the higher temperatures at widely distributed sites in E210Q-*EcS*ecA, including some residues in the cores of the HSD, NBF-I, and NBF-II. The overall pattern of the observed changes is similar to that observed in the HDX experiments on E208Q-*BsS*ecA (Figure 8B), as schematized in Figure 9. Therefore, the E-to-Q mutation in the catalytic base facilitates the ECT in both enzymes, but it does so at a higher temperature in *EcS*ecA.

In comparing apo E210Q-EcSecA to the WT enzyme, evidence is observed of local increases in HDX in some regions of the protein, while other segments involved in the ECT remain relatively unperturbed. These observations suggest that the broad shoulder at 25-35 °C in DSC experiments on the mutant enzyme (Figure 7G) may reflect a cascade of localized changes in protein dynamics sequentially affecting different regions of the mutant enzyme, rather than a simple two-state conformational transition. For example, segments of NBF-I proximal to the mutation site as well as motif V and an adjacent segment of NBF-II all show significant increases in HDX rates in the mutant enzyme compared to the WT enzyme even at 25 °C (Figure 8D), well prior to the midpoint of the ECT in the mutant enzyme at ~39  $^{\circ}C$  (as monitored by DSC in Figure 7G). In contrast, some regions in the PPXD and NBF-II that show enhanced exchange at the outset of the ECT at 37 °C in the mutant enzyme (Figure 8H) are unperturbed at the intermediate temperature of 31 °C (Figure 8F). These include one of the  $\alpha$ -helices in the VAR segment<sup>66</sup> (shown at the very bottom of the structure in Figure 8C-H). This segment contains a pair of  $\alpha$ -helices that packs on the surface of NBF-II in SecA orthologs from Gram-negative eubacteria like E. coli, but it is absent in orthologs from Gram-positive eubacteria like B. subtilis. Notably, this segment exchanges more slowly than any other observed region of NBF-II in EcSecA (Figure 8C,E,G), consistent with the proposal that it allosterically stabilizes the interactions between NBF-I and NBF-II that control nucleotide release.<sup>66</sup> Its presence in *Ec*SecA but absence from BsSecA therefore could contribute to the lesser effect of the E-to-Q mutation in the catalytic base of EcSecA vs BsSecA at 25 °C.

While the HDX data on NBF-II show that portions of its core as well as the VAR segment remain predominantly folded in E210Q-*Ec*SecA at 37 °C, the coverage of NBF-II is incomplete, preventing assessment of the dynamics of many segments including helicase motif VI (which was not observed in any of the HDX experiments conducted on E210Q-*Ec*SecA). Therefore, future experiments will be required to elucidate the conformational details accounting for the broad shoulder or pretransition at 25–35 °C in DSC experiments on apo E210Q-*Ec*SecA (Figure 7G) and to determine whether it reflects sequential conformational transitions increasing the flexibility of local protein segments. Even if it does, the HDX data in Figure 8 show that the overall conformational state produced by these sequential transitions is very similar to that produced by the ostensibly discrete ECT in *Bs*SecA.

# DISCUSSION

The thermodynamic studies reported in this paper substantially enhance understanding of the physicochemical factors controlling the function of a large mechanoenzyme mediating an exceedingly complex molecular process. These studies demonstrate that altering a single electrostatic charge in the ATPase active site of SecA can trigger a global conformational change in this 100 kDa mechanoenzyme. With >250 charged amino acid residues in SecA, the network of electrostatic interactions in the enzyme must be exquisitely balanced,<sup>26,27</sup> and this balance is likely to play a central role in coupling transient electrostatic changes during ATP hydrolysis in the active site to the global conformational changes that drive processive polypeptide transport.<sup>33,34</sup> These conformational changes alter cooperative atomic interactions between different domains<sup>12,13,26</sup> and possibly also between structural modules within individual domains (e.g., helicase motifs V/VI and the methionine canyon in NBF-II). The thermodynamic data presented in this paper show a substantial increase in entropy during the principal functional conformational transition in SecA (i.e., the ECT), while the HDX data support the interpretation that this increase derives from an enhancement in net backbone dynamics in many of its domains and structural modules. Thus, the conformational reaction cycle of SecA harnesses both the electrostatic and statistical properties of the protein polymer to orchestrate the efficient progression of an extremely complex physical process that is tightly coupled to the binding and hydrolysis of ATP.

The glutamate-to-glutamine mutation in the catalytic base emulates its change in electrostatic charge state during the initial step in the ATPase reaction in which this glutamate abstracts a proton from the hydrolytic water molecule to initiate a nucleophilic attack on the  $\gamma$ -phosphate of ATP.<sup>33,34</sup> This proton-transfer event induces cleavage of the phosphoanhydride bond, which is rapidly followed by conformational rearrangement of the released inorganic phosphate and ADP in the active site. Subsequent events, including proton release from the side chain of the catalytic glutamate and release of the cleaved inorganic phosphate from the active site, are likely to involve significant changes in electrostatic and other energetic interactions in the active site. The conformational effects of the E-to-Q mutation demonstrate that there is exquisitely sensitive coupling of the global conformational state of SecA to local electrostatic interactions at the catalytic center,  $^{\rm 33,34}$  and they are likely to reflect at least in part the influence on global protein conformation of the initial proton-abstraction event during ATP hydrolysis. Because electrostatic interactions can propagate through a protein structure very rapidly, the conformational couplings of the catalytic glutamate elucidated in our experiments can help drive SecA through the proper global conformational trajectory contemporaneously during the ATP hydrolysis reaction.

The HDX data in Figure 8 demonstrate that the ECT involves a substantial increase in net backbone dynamics in widely distributed regions of SecA, and this increase is likely to be an important driving force for this critical functional conformational transition. For a reaction to be endothermic, it must involve an increase in enthalpy (i.e., input of heat) as well as entropy, because changing  $\Delta G = \Delta H - T \cdot \Delta S$  from positive to negative values by increasing temperature mathematically requires both  $\Delta H > 0$  and  $\Delta S > 0$ . The widespread increase in conformational entropy, inferred from coordinated thermody-

namic and HDX experiments on both *Bs*SecA and *Ec*SecA (Figure 8), is likely to make a critical contribution to the entropy gain driving the ECT.

The delocalized nature of these changes is noteworthy because order-to-disorder transitions in helicase motifs V and VI have previously been hypothesized to be the entropic driver for the ECT.<sup>26</sup> Our HDX data (Figure 8) suggest that the entropic driver instead involves broadly distributed increases in protein dynamics throughout the multidomain protein structure, including most interdomain interfaces in SecA. This inference is supported by the observation that the E210Q mutation in *EcSecA* produces a significant increase in HDX rate and thus net backbone dynamics near both the mutation site in NBF-I and near helicase motif V in NBF-II even at 25 °C (Figure 8C vs 8D), which is well below the temperature of the ECT in E210Q-EcSecA. In the absence of Mg-ADP, the ECT in this protein variant has an apparent midpoint at ~39 °C following a broad pretransition from 25 to 35 °C as monitored by DSC (Figure 7G). Dynamic transitions already in progress at 25 °C in this protein variant are unlikely to represent the major entropic driver for a conformational transition centered at 39 °C. Therefore, the globally distributed increases in net backbone dynamics observed in the HDX studies of both BsSecA and EcSecA represent a more likely candidate to be the entropic driver controlling the progression of the ECT.

Nonetheless, the totality of our experimental data suggests that a number of distinct conformational transitions involving alterations in protein dynamics in different regions of SecA contribute to controlling the efficient progression of its conformational reaction cycle. It seems likely that some of these transitions can proceed independently, while others have varying degrees of thermodynamic interdependency. For example, the broad shoulder preceding the ECT in DSC experiments on E210Q-EcSecA (Figure 7G) suggests that this mutant enzyme undergoes one or more endothermic conformational subtransitions preceding the global change in interdomain interactions that occurs during the ECT (Figure 8); presumably residue E210 in EcSecA strengthens the thermodynamic linkage between these sequential conformational transitions in the WT enzyme. Another example is provided by our earlier study characterizing the nucleotideinteraction properties of EcSecA,<sup>13</sup> which demonstrated that different mutations have uncorrelated effects on Mg-ADPrelease rate and the ECT. Therefore, the ECT must differ in at least some details from the conformational transition controlling Mg-ADP release. This latter transition could conceivably involve an order-to-disorder transition in helicase motifs V and VI uncoupled from global conformational changes;<sup>61</sup> this possibility is supported by the observation that the E210Q mutation in EcSecA greatly accelerates Mg-MANT-ADP-release rate (Figure 3) while also increasing the dynamics of helicase motif V (Figure 8C vs D) at 25 °C, well below the onset of the ECT in this enzyme (Figure 7G). Notably, motifs V and VI both exhibit very high B factors in crystal structures of some homologous helicases.<sup>67,68</sup> However, the large entropy-enthalpy compensation that accompanies Mg-ADP binding to WT-EcSecA compared to WT-BsSecA at 25 °C (Table 1) is likely to derive from a dynamic transition involving different protein segments or alternatively dynamics on a faster time scale than that accessible to HDX measurements, because our data support both of these segments adopting a folded conformation in apo EcSecA at this temperature on the time scale of the HDX process. While

future studies will be required to evaluate the specific structural hypotheses advanced here, the available thermodynamic and dynamic data provide strong evidence that SecA undergoes a variety of distinct conformational transitions that are likely to involve changes in protein dynamics.

### CONCLUSIONS

Although the ECT strongly increases net protein dynamics at many sites, the HDX data suggest that it simultaneously involves a reduction in protein dynamics at specific sites (e.g., the methionine canyon, as shown in Figure 8A–B), and these sites could participate in stabilizing structural interactions with ligands or other proteins that guide the progression of SecA's conformational reaction cycle. In this manner, interacting structural modules undergoing sequential conformational transitions could control the progression of a large enzyme like SecA through a sequence of different conformational states each optimized to make different molecular contacts. A molecular trajectory of this kind, controlled by an exquisitely balanced network of electrostatic interactions connecting to the ATPase active site, offers clear advantages for efficient coupling of ATP binding and hydrolysis to a complex molecular process like processive extrusion of a polypeptide chain through an interacting channel. Having a cascade of distinct dynamic transitions underlie such a sequence of conformational changes would enable the inherent plasticity in protein structure, and the diverse physicochemical factors underlying this plasticity, to be exploited for efficient execution of a remarkably complex molecular process. Future research will be required to critically evaluate these mechanistic hypotheses that emerge from the integrated biophysical studies of SecA presented in the current paper. Please see the SI for further discussion.

## ASSOCIATED CONTENT

## **Supporting Information**

An expanded discussion, the crystallographic data table, and Figures S1-S5 are provided as well as a file containing processed HDX data. This material is available free of charge via the Internet at http://pubs.acs.org.

#### AUTHOR INFORMATION

#### **Corresponding Author**

jfhunt@biology.columbia.edu

#### **Present Address**

<sup>1</sup>Department of Anesthesiology, Weill Cornell Medical College, 1300 York Ave, New York, NY, 10021

#### Notes

The authors declare no competing financial interest.

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# REFERENCES

- (1) Devillers-Thiery, A.; Kindt, T.; Scheele, G.; Blobel, G. Proc. Natl. Acad. Sci. U.S.A. 1975, 72, 5016.
- (2) Oliver, D. B.; Beckwith, J. J. Bacteriol. 1982, 150, 686.
- (3) Bieker-Brady, K.; Silhavy, T. J. EMBO J. 1992, 11, 3165.
- (4) Huie, J. L.; Silhavy, T. J. J. Bacteriol. 1995, 177, 3518.
- (5) Brundage, L.; Hendrick, J. P.; Schiebel, E.; Driessen, A. J.; Wickner, W. Cell 1990, 62, 649.
- (6) Driessen, A. J. EMBO J. 1992, 11, 847.
- (7) Hanada, M.; Nishiyama, K. I.; Mizushima, S.; Tokuda, H. J. Biol. Chem. **1994**, 269, 23625.
- (8) Eichler, J.; Brunner, J.; Wickner, W. EMBO J. 1997, 16, 2188.
- (9) Eichler, J.; Wickner, W. Proc. Natl. Acad. Sci. U.S.A. 1997, 94, 5574.
- (10) Schiebel, E.; Driessen, A. J.; Hartl, F. U.; Wickner, W. Cell 1991, 64, 927.
- (11) van der Wolk, J. P.; de Wit, J. G.; Driessen, A. J. EMBO J. 1997, 16, 7297.
- (12) Hunt, J. F.; Weinkauf, S.; Henry, L.; Fak, J. J.; McNicholas, P.; Oliver, D. B.; Deisenhofer, J. Science **2002**, 297, 2018.
- (13) Fak, J. J.; Itkin, A.; Ciobanu, D. D.; Lin, E. C.; Song, X. J.; Chou, Y. T.; Gierasch, L. M.; Hunt, J. F. *Biochemistry* **2004**, *43*, 7307.
- (14) Velankar, S. S.; Soultanas, P.; Dillingham, M. S.; Subramanya, H. S.; Wigley, D. B. *Cell* **1999**, *97*, 75.
- (15) Leslie, A. G.; Abrahams, J. P.; Braig, K.; Lutter, R.; Menz, R. I.; Orriss, G. L.; van Raaij, M. J.; Walker, J. E. *Biochem. Soc. Trans.* **1999**, 27, 37.
- (16) Sharma, V.; Arockiasamy, A.; Ronning, D. R.; Savva, C. G.; Holzenburg, A.; Braunstein, M.; Jacobs, W. R., Jr.; Sacchettini, J. C. *Proc. Natl. Acad. Sci. U.S.A.* **2003**, *100*, 2243.
- (17) Cavanaugh, L. F.; Palmer, A. G., III; Gierasch, L. M.; Hunt, J. F. Nat. Struct. Mol. Biol. 2006, 13, 566.
- (18) Economou, A.; Wickner, W. Cell 1994, 78, 835.
- (19) Rajapandi, T.; Oliver, D. Biochem. Biophys. Res. Commun. 1994, 200, 1477.
- (20) Driessen, A. J. Trends Biochem. Sci. 1992, 17, 219.
- (21) Or, E.; Boyd, D.; Gon, S.; Beckwith, J.; Rapoport, T. J. Biol. Chem. 2005, 280, 9097.
- (22) Ulbrandt, N. D.; London, E.; Oliver, D. B. J. Biol. Chem. 1992, 267, 15184.
- (23) den Blaauwen, T.; Fekkes, P.; de Wit, J. G.; Kuiper, W.; Driessen, A. J. *Biochemistry* **1996**, *35*, 11994.
- (24) den Blaauwen, T.; van der Wolk, J. P.; van der Does, C.; van Wely, K. H.; Driessen, A. J. *FEBS Lett.* **1999**, *458*, 145.
- (25) Schmidt, M.; Ding, H.; Ramamurthy, V.; Mukerji, I.; Oliver, D. J. Biol. Chem. **2000**, 275, 15440.
- (26) Keramisanou, D.; Biris, N.; Gelis, I.; Sianidis, G.; Karamanou, S.; Economou, A.; Kalodimos, C. G. Nat. Struct. Mol. Biol. 2006, 13, 594.
- (27) Karamanou, S.; Gouridis, G.; Papanikou, E.; Sianidis, G.; Gelis, I.; Keramisanou, D.; Vrontou, E.; Kalodimos, C. G.; Economou, A.
- EMBO J. 2007, 26, 2904. (28) Vrontou, E.; Karamanou, S.; Baud, C.; Sianidis, G.; Economou,
- A. J. Biol. Chem. 2004, 279, 22490.
- (29) Osborne, A. R.; Clemons, W. M., Jr.; Rapoport, T. A. Proc. Natl. Acad. Sci. U.S.A. 2004, 101, 10937.
- (30) Vassylyev, D. G.; Mori, H.; Vassylyeva, M. N.; Tsukazaki, T.; Kimura, Y.; Tahirov, T. H.; Ito, K. *J. Mol. Biol.* **2006**, *364*, 248.
- (31) Zimmer, J.; Li, W.; Rapoport, T. A. J. Mol. Biol. 2006, 364, 259.
- (32) Zimmer, J.; Nam, Y.; Rapoport, T. A. Nature 2008, 455, 936.
- (33) Jencks, W. P. Catalysis in chemistry and enzymology; McGraw-Hill: New York, 1969, pp 40, 111–115.
- (34) Jencks, W. P. Adv. Enzymol. Relat. Areas Mol. Biol. 1980, 51, 75.
  (35) Story, R. M.; Li, H.; Abelson, J. N. Proc. Natl. Acad. Sci. U.S.A.
  2001, 98, 1465.
- (36) Caruthers, J. M.; McKay, D. B. Curr. Opin. Struct. Biol. 2002, 12, 123.

(38) de la Cruz, J.; Kressler, D.; Linder, P. Trends Biochem. Sci. 1999, 24, 192.

- (39) Linder, P.; Lasko, P. F.; Ashburner, M.; Leroy, P.; Nielsen, P. J.; Nishi, K.; Schnier, J.; Slonimski, P. P. *Nature* **1989**, 337, 121.
- (40) Koonin, E. V.; Gorbalenya, A. E. FEBS Lett. 1992, 298, 6.
- (41) Zito, C. R.; Antony, E.; Hunt, J. F.; Oliver, D. B.; Hingorani, M. M. J. Biol. Chem. 2005, 280, 14611.
- (42) Abrahams, J. P.; Leslie, A. G.; Lutter, R.; Walker, J. E. Nature 1994, 370, 621.
- (43) Smith, P. C.; Karpowich, N.; Millen, L.; Moody, J. E.; Rosen, J.; Thomas, P. J.; Hunt, J. F. *Mol. Cell* **2002**, *10*, 139.
- (44) Moody, J. E.; Millen, L.; Binns, D.; Hunt, J. F.; Thomas, P. J. J. Biol. Chem. 2002, 277, 21111.
- (45) Chen, J.; Lu, G.; Lin, J.; Davidson, A. L.; Quiocho, F. A. Mol. Cell 2003, 12, 651.
- (46) Khare, D.; Oldham, M. L.; Orelle, C.; Davidson, A. L.; Chen, J. *Mol. Cell* **2009**, 33, 528.
- (47) Walker, J. E.; Saraste, M.; Runswick, M. J.; Gay, N. J. *EMBO J.* **1982**, *1*, 945.
- (48) Ramamurthy, V.; Oliver, D. J. Biol. Chem. 1997, 272, 23239.
- (49) Ding, H.; Mukerji, I.; Oliver, D. Biochemistry 2001, 40, 1835.
- (50) van der Does, C.; Manting, E. H.; Kaufmann, A.; Lutz, M.; Driessen, A. J. Biochemistry 1998, 37, 201.
- (51) Economou, A.; Pogliano, J. A.; Beckwith, J.; Oliver, D. B.; Wickner, W. Cell **1995**, 83, 1171.
- (52) Weber, J.; Senior, A. E. *Biochim. Biophys. Acta* 1997, 1319, 19.
   (53) Shimo-Kon, R.; Muneyuki, E.; Sakai, H.; Adachi, K.; Yoshida,
- M.; Kinosita, K. *Biophys. J.* **2010**, *98*, 1227. (54) Rye, H. S.; Burston, S. G.; Fenton, W. A.; Beechem, J. M.; Xu,
- Z.; Sigler, P. B.; Horwich, A. L. Nature **1997**, 388, 792.
- (55) Roseman, A. M.; Chen, S.; White, H.; Braig, K.; Saibil, H. R. Cell **1996**, *87*, 241.
- (56) Yuan, Y. R.; Blecker, S.; Martsinkevich, O.; Millen, L.; Thomas, P. J.; Hunt, J. F. *J. Biol. Chem.* **2001**, *276*, 32313.
- (57) Sharma, S.; Zheng, H.; Huang, Y. J.; Ertekin, A.; Hamuro, Y.; Rossi, P.; Tejero, R.; Acton, T. B.; Xiao, R.; Jiang, M.; Zhao, L.; Ma, L. C.; Swapna, G. V.; Aramini, J. M.; Montelione, G. T. *Proteins* **2009**, *76*, 882.
- (58) Sianidis, G.; Karamanou, S.; Vrontou, E.; Boulias, K.; Repanas, K.; Kyrpides, N.; Politou, A. S.; Economou, A. *EMBO J.* **2001**, *20*, 961.
- (59) Natale, P.; Swaving, J.; van der Does, C.; de Keyzer, J.; Driessen, A. J. J. Biol. Chem. 2004, 279, 13769.
- (60) Bennett, B. D.; Kimball, E. H.; Gao, M.; Osterhout, R.; Van Dien, S. J.; Rabinowitz, J. D. *Nat. Chem. Biol.* **2009**, *5*, 593.
- (61) Chou, Y. T.; Swain, J. F.; Gierasch, L. M. J. Biol. Chem. 2002, 277, 50985.
- (62) Bai, Y.; Englander, J. J.; Mayne, L.; Milne, J. S.; Englander, S. W. *Methods Enzymol.* **1995**, 259, 344.
- (63) Englander, J. J.; Del Mar, C.; Li, W.; Englander, S. W.; Kim, J. S.; Stranz, D. D.; Hamuro, Y.; Woods, V. L., Jr. *Proc. Natl. Acad. Sci. U.S.A.* **2003**, *100*, 7057.
- (64) Ding, H.; Mukerji, I.; Oliver, D. Biochemistry **2003**, 42, 13468. (65) Papanikou, E.; Karamanou, S.; Economou, A. Nat. Rev. Microbiol. **2007**, 5, 839.
- (66) Das, S.; Grady, L. M.; Michtavy, J.; Zhou, Y.; Cohan, F. M.; Hingorani, M. M.; Oliver, D. B. J. Bacteriol. **2012**, 194, 2205.
- (67) Caruthers, J. M.; Johnson, E. R.; McKay, D. B. Proc. Natl. Acad. Sci. U.S.A. 2000, 97, 13080.
- (68) Liu, H.; Rudolf, J.; Johnson, K. A.; McMahon, S. A.; Oke, M.; Carter, L.; McRobbie, A. M.; Brown, S. E.; Naismith, J. H.; White, M. F. *Cell* **2008**, *133*, 801.

<sup>(37)</sup> Singleton, M. R.; Wigley, D. B. J. Bacteriol. 2002, 184, 1819.