Single-molecule observation of nucleotide induced conformational changes in basal SecA-ATP hydrolysis

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SecA is the critical adenosine triphosphatase that drives preprotein transport through the translocon, SecYEG, in Escherichia coli. This process is thought to be regulated by conformational changes of specific domains of SecA, but real-time, real-space measurement of these changes is lacking. We use single-molecule atomic force microscopy (AFM) to visualize nucleotide-dependent conformations and conformational dynamics of SecA. Distinct topographical populations were observed in the presence of specific nucleotides. AFM investigations during basal adenosine triphosphate (ATP) hydrolysis revealed rapid, reversible transitions between a compact and an extended state at the ~100-ms time scale. A SecA mutant lacking the precursor-binding domain (PBD) aided interpretation. Further, the biochemical activity of SecA prepared for AFM was confirmed by tracking inorganic phosphate release. We conclude that ATP-driven dynamics are largely due to PBD motion but that other segments of SecA contribute to this motion during the transition state of the ATP hydrolysis cycle.

INTRODUCTION

More than 30% of newly synthesized proteins in Escherichia coli are exported across the cytoplasmic membrane to their final destinations, where they acquire their natively folded and functional state (1, 2). Several distinct export mechanisms have evolved to accomplish this protein maneuvering, but only the general secretory (Sec) system is found in all domains of life. Proteins to be exported are synthesized as precursors with an N-terminal leader sequence and are translocated through the Sec pathway in E. coli. SecA, the adenosine triphosphatase (ATPase) of the Sec system, binds to the translocon SecYEG to form a translocase. In a process that is currently not well understood, this complex mediates chemomechanical conversion that results in translocation of precursor proteins across the cytoplasmic membrane (3, 4).

Peripheral membrane protein SecA is a large multidomain protein (901 aminoacyl residues; Fig. 1A) with numerous binding partners including the translocon SecYEG, chaperone SecB, precursor proteins, and lipids (5–8). The multiple and complex functions that this macromolecule performs in a cell have likely led to the remarkable dynamics and flexibility that the domains of SecA exhibit (1, 9). For example, the precursor-binding domain (PBD) is thought to move approximately 80° relative to other domains (Fig. 1B). Several models have been proposed to explain protein translocation with different roles for SecA, and interdomain allosteric regulation of ATPase activity is a common theme among them (1). SecA acts as a processive motor in a “piston” model where extended precursor protein segments move along its clamp (10, 11). The tip of the two-helix finger [alternatively referred to as intramolecular regulator of ATPase (IRA1); Fig. 1, A and C, orange] has been proposed to latch onto the preprotein chain and push it through the SecYEG pore. The precursor is then translocated through the channel via adenosine triphosphate (ATP) hydrolysis–driven power strokes in a stepwise fashion (4, 12). In another model, SecA controls SecYEG pore opening and closing through ATP hydrolysis, hence acting as an allosteric channel regulator. This facilitates a Brownian ratchet mechanism where the precursor protein undergoes proton motive force driven diffusion through the SecYEG channel, and SecA acts as a “brake” to prevent backsliding (3, 13).

How SecA harnesses the energy released during ATP hydrolysis and translates it into functional conformational transitions to drive precursor protein chains through the translocon remains unclear. Major insight has come from x-ray crystallographic studies (14, 15). The N-terminal portion of SecA consists of three large domains, namely two nucleotide-binding folds [Fig. 1, A (red, NBD1; blue, NBD2) and B (gray)] and the PBD (Fig. 1, A to C, pink). ATP binds at the interface of NBD1 and NBD2, forming a superfamily 2 helicase motor (1). The energy released during hydrolysis has been proposed to allosterically mobilize the PBD and the C-terminal segment [which includes most of the helical scaffold domain (HSD) and all of the helical wing domain to the C terminus; Fig. 1, B and C (crimson) (residues 620 to 901)] (2, 9). SecA has been crystallized in the presence of different nucleotides and has exhibited three major conformational states: wide-open, open, and closed (Fig. 1B). It has thus been hypothesized that the PBD of SecA can swivel from wide-open to a closed state in a ligand-dependent manner (1, 10, 14, 16, 17).

A complementary line of inquiry to the static crystal structures would be to observe the conformational motions of SecA molecules during ATP hydrolysis in real time and in near-native conditions. Because these motions are triggered by stochastic events, they are asynchronous and challenging to resolve. Atomic force microscopy (AFM) is a single-molecule imaging technique that is well suited to these measurements. AFM uses an extremely sharp tip to probe protein topography and has proven useful in a number of biological contexts, primarily because it achieves molecular resolution imaging in solution in real time without the need for freezing or crystalization and with minimal molecular weight constraints (18, 19). Despite its strengths, AFM generally requires proteins to be adsorbed to a supporting surface; this specimen preparation step can...
raise questions about activity. In principle, the product of surface-adsorbed enzymes can be measured to provide biochemical verification of activity.

Previously, our group applied AFM to study the translocon and translocase complexes in supported lipid bilayers. We measured the dynamics of the membrane external cytoplasmic loops of SecYEG and observed the association and dissociation of SecA with the translocon (7, 20, 21). Here, we apply single-molecule AFM imaging to study the conformational dynamics associated with basal SecA ATP hydrolysis and elucidate nucleotide-coupled conformational states.

To focus on basal activity, SecA was passively adsorbed onto mica surfaces in the absence of binding partners. This also improves precision, as lipid bilayers can add noise to the measurement, for example, through vertical conformational fluctuations (21). We demonstrate angstrom-scale changes in protein conformations that correlate with particular domains and particular nucleotide species. Conformational changes were captured in real time at the single-molecule level with up to ~100-ms temporal resolution in physiological solution. Biochemical measurements tracking the release of radiolabeled γ-phosphate confirmed that the surface-adsorbed SecA prepared for AFM measurements still hydrolyzed ATP. Together, our work demonstrates through single-molecule observation that ATP hydrolysis drives large-scale conformational changes of specific domains of SecA.

RESULTS
We shed light on the conformational dynamics of SecA during ATP hydrolysis by imaging single molecules in near-native conditions. Following previous work (7, 20, 21), proteins were adsorbed to freshly cleaved mica and imaged via tapping mode AFM in aqueous solution. Salient topographic information, including the height and area of individual macromolecules, was extracted algorithmically and collected for statistical analyses (22).
We studied first wild-type SecA (SecA-WT). SecA-WT exists in equilibrium between monomer and dimer in E. coli. We focused on conformational dynamics of SecA monomer by choosing solution conditions that bias the system away from dimer (23). Figure 1D shows an AFM image of about 100 SecA-WT molecules. Detailed topographic images of individual proteins are also shown [Fig. 1, E to I; fig. S1 shows phase images (24)]. Simulated AFM images were constructed using x-ray crystal coordinates in conjunction with an assumed tip geometry (see Materials and Methods for details). Simulated images showed good overall agreement with the experimental data, and analysis implied that the orientation of the SecA on the mica was not random but had two preferred orientations (figs. S2 and S3).

We determined the relationship between specific topographical populations and the functionally important PBD of SecA. The distribution of feature height for individual SecA-WT molecules in the apo state (i.e., not exposed to any nucleotide after purification) is presented (Fig. 1, black). These data show that most of the conformations exhibit a maximum height of around 4 nm. A closer inspection reveals a minimum separating two subpopulations that are spaced ~0.5 nm apart. Although small, this separation is above the ~0.1-nm vertical resolution of the technique (25). Figure S4 demonstrates that the separation between the two peaks is robust. We observed an additional subpopulation in the height distribution as a distinct shoulder at about 5.1 nm (Fig. 1, light blue shade). This shoulder population was substantial, comprising 29% of the total. When a variant of SecA with the PBD deleted (SecAAPBD) was analyzed, the population at 5.1 nm was diminished significantly (Fig. 1K, light blue shade). At the same time, the population at 4.3 nm was enhanced by roughly the same amount (compare Fig. 1, J and K, green shades), and the other three populations were essentially unchanged compared to the WT enzyme (Fig. 1, J and K; dark blue, orange, and pink shades). We emphasize that the peak positions of the two major populations match exactly; hence, SecA-WT and the mutant lacking the PBD are likely to be in the same overall orientation on the mica. This notion is buttressed by similarities in equipotential surface maps between the two SecA species (fig. S5). Hence, the conformational state at 5.1 nm appears to be directly related to the PBD. We note that cells require the PBD of SecA; SecAAPBD was overproduced, and the purified sample contained <10% residual SecA-WT.

Before characterizing enzyme topography during ATP hydrolysis, we sought to verify the ATPase activity of SecA samples as prepared for AFM study. To this end, we performed thin-layer chromatography to track the dephosphorylation of [γ-32P]-labeled ATP. The approach was based on prior work (26) modified to probe surface-adsorbed enzymes. We prepared SecA-WT samples on mica in a manner identical to that used for AFM imaging, but at a higher concentration (2.4 μM) to increase the signal. At this concentration, most of the SecA will be dimer [Kd (dissociation constant) ≤ 1 μM] (2); however, basal ATPase activity of SecA is weakly coupled to oligomeric state (27). The generation of inorganic phosphate (Pi) from mica-adsorbed SecA-WT over time is shown (Fig. 2). We carried out control experiments on mica in the absence of SecA. The results show that SecA actively hydrolyzes ATP when adsorbed to mica, even over extended time periods (>2 hours) that are frequently used for AFM study.

What fraction of the SecA molecules retained ATPase activity when bound to the mica surface? To estimate this, we used AFM data and molecular dimensions to determine that approximately 3 × 10⁻¹² mol of SecA was available on the mica disk during the surface-adsorbed ATPase assay (fig. S6). Figure 2C shows a time course of mol ATP hydrolyzed divided by mol SecA available. The slope represents the normalized ATP turnover rate. For comparison, we carried out a traditional solution-based assay with SecA at a monomer concentration of 2.4 μM (Fig. 2D). The results indicate that there is only a modest decrease of 6% in activity when the molecules are adsorbed onto mica. This unexpected result suggests that approximately 9 of 10 SecA molecules remain active for ATP turnover when adsorbed on mica. We note that, in addition to the aforementioned estimate of SecA copy number, there are other uncertainties in this result. For example, it is possible that the surface itself can act as a mild destabilization agent; it has been shown that low levels of the denaturant urea stimulate the ATPase activity of SecA (28). In the face of caveats, our results suggest that a significant fraction of SecA molecules remain competent for ATP hydrolysis when adsorbed onto mica for AFM analysis.

After confirming that ATPase activity was maintained, we studied the dynamic topography of SecA during ATP hydrolysis. The SecA-WT and the SecAAPBD mutant were subjected to four different ATP assays: (i) apo (Fig. 3, black); (ii) preincubated with 100 μM ATP for 15 min before deposition on mica and then imaged in the absence of ATP (Fig. 3, red); (iii) preincubated with 100 μM ATP in solution for 15 min and then imaged with 100 μM ATP in the imaging buffer (Fig. 3, green); and last, to monitor the long time-scale recovery of the enzyme after ATP exposure; (iv) preincubated with 100 μM ATP in solution for 4 hours and then imaged in the absence of ATP in the imaging buffer (Fig. 3, blue). The concentration of ATP chosen for this study was significantly higher than the reported affinity for ATP (Kd ~ 300 nM) (29). Representative images of SecA-WT and SecAAPBD from the four assays are shown (fig. S7).

We determined the footprint area of each protein (i.e., the area the macromolecules projected onto the local background). These footprint area distributions are presented as smoothed histograms in Fig. 3 [A (SecA-WT) and C (SecAAPBD)]. When exposed to ATP, the areal footprint of SecA-WT increased in all conditions studied [compare primary peak position for the apo state of 200 nm² (black) to 270 nm² (red), 300 nm² (green), and 280 nm² (blue); Fig. 3A]. The observation of increased area suggests that SecA-WT explores a wider conformational space upon ATP exposure. Furthermore, the area increases remained 4 hours after ATP exposure; hence, the ATP-induced changes were persistent.

In contrast to SecA-WT, three of the four principal populations of the SecAAPBD footprint area remained essentially unchanged after ATP exposure [compare the primary peak position of 230 nm² (black) to 230 nm² (red) and 235 nm² (blue); Fig. 3C]. This implies that the aforementioned increases in the area observed for SecA-WT are due to PBD motion. We note that SecA lacking the PBD still hydrolyzes ATP (30). Hence, an overall destabilization of the macromolecule due to the energy released upon ATP hydrolysis can be ruled out. The primary peak observed at 300 nm² in the case of the SecAAPBD mutant when imaged in the presence of ATP (Fig. 3C, green) could be attributable to motion of the NBD1 and NBD2 domains, but we think that it is more likely to be the C-terminal segment that moves. The C-terminal segment of SecA has been shown to be highly mobile and is thought to be an additional regulator of ATP hydrolysis that inserts into membrane during translocation (9, 17). Statistics of the area footprint distributions and heights for SecA-WT and SecAAPBD in different conditions are presented (see tables S1 to S4).
In addition to area analyses, the height distributions also indicated that SecA-WT was conformationally more dynamic as compared to the SecAΔPBD mutant. Figure 3 (B and D) shows height distributions of SecA-WT and SecAΔPBD, respectively. The full width at half maximum (FWHM) of SecA-WT heights was larger than that of the mutant in all the conditions studied. For example, the FWHM in the presence of ATP was 2.3 nm for SecA-WT compared to 1.5 nm for the SecAΔPBD mutant (green curves in Fig. 3, B and D). This difference can be attributed to the PBD and motions thereof. Differences in conformational dynamics (evident by FWHM of the height distributions) were pronounced even in the absence of ATP. The FWHM of apo SecA-WT was 50% larger than that of apo SecAΔPBD.

Preincubation in ATP, followed by imaging in the absence of ATP (Fig. 3, red), shifted the overall height of SecA downward by ~0.4 nm for both SecA-WT and the SecAΔPBD mutant. When imaged in ATP, this shift was eliminated for SecA-WT, but not for SecAΔPBD. A similar conformational change was reported for a different enzyme (adenylate kinase) when exposed to ATP and was attributed to a partially closed conformational state that would allow rapid product-substrate exchange (31). The overall similarity between the three SecA-WT height distributions is notable (Fig. 3B); all conditions showed two principle height peaks and a prominent shoulder at ~5 nm.

To further investigate the conformational changes, we exposed SecA-WT and the SecAΔPBD mutant to two nonhydrolyzable nucleotides: adenosine diphosphate (ADP) and ADP-AlF₃, an ATP analog that conformationally stabilizes or traps the ATPase in its transition state (32, 33). Enzymes were exposed to ADP or ADP-AlF₃ for 15 min before imaging and then imaged in the absence of nucleotides. Representative images of SecA-WT and SecAΔPBD after ADP or ADP-AlF₃ exposure are shown (fig. S8). Footprint area distributions are presented in Fig. 4 [A (SecA-WT) and B (SecAΔPBD)]. SecA-WT footprint areas increased when compared to the apo state: After ADP exposure, the primary peak position was at 220 nm² (Fig. 4A, cyan); after ADP-AlF₃ exposure, the primary peak was at 235 nm² (Fig. 4A, gray). The magnitudes of these increases were smaller when compared to preincubation in ATP (Fig. 4A, red dashed). In the case of SecAΔPBD, the area distributions remained essentially the same after exposure to ADP or the nonhydrolyzable analog ADP-AlF₃ [primary peak positions: 220 nm² (cyan) and 220 nm² (gray); Fig. 4B]. This provides further evidence that the corresponding increases in footprint area observed with SecA-WT were due to PBD motion.

Figure 4 (C and E) shows height distributions of SecA-WT after ADP and ADP-AlF₃ exposure, respectively. The height shifted lower than that of the apo state when the WT enzyme was exposed to either of these two nucleotides. The FWHM of these data and the associated statistics are presented in table S4. The height distributions after exposure to the two nonhydrolyzable nucleotides were sharply peaked as compared to the apo state (compare cyan and gray to black dashed; Fig. 4, C and E). Thus, it appears that most of the populations observed upon ADP and ADP-AlF₃ exposure represent nucleotide-stabilized conformational states, similar to those reported previously in bulk studies (29). Height histograms acquired with either ADP or ADP-AlF₃ in the imaging buffer (fig. S9) were similar to the nucleotide-exposed conditions. Hence, conformational changes were persistent.

It is informative to compare height distribution shifts for SecA-WT and SecAΔPBD when exposed to the two nonhydrolyzable nucleotides. In the case of the SecAΔPBD mutant, the height distributions for the apo state and the ADP-exposed state were similar (Fig. 4D). In contrast, for the WT species, the ADP-exposed state was altered significantly from the apo state, exhibiting a single sharp peak compared to a bimodal apo peak (Fig. 4C). These results imply that ADP induces a conformational state change in the PBD.
The results with the transition state analog provide additional insight into the conformational dynamics. For the SecAΔPBD mutant, the ADP-AlF₃– and ATP-exposed states exhibited similar height distributions (Fig. 4F, gray and red dashed, respectively), and both nucleotides shifted the overall heights lower compared to the apo state. Analogous behavior was observed for the SecA-WT species (Fig. 4E). The shifts from the apo state to the ADP-AlF₃ state were similar in magnitude for SecA-WT and SecAΔPBD and suggestive of a common underlying origin. It thus appears that ADP-AlF₃ binding induces a conformational change, likely in the C-terminal segment of SecA, that manifests as a height change in the AFM measurements. Area analysis showed ADP-AlF₃–induced shifts for SecA-WT, but not for SecAΔPBD. We conclude that the area shift results from PBD motion, as this is only seen in the WT enzyme, whereas the height shift is common to both species and is likely due to the C-terminal segment. Together, these data indicate a concomitant conformational change: both the PBD and likely the C-terminal segment move upon ADP-AlF₃ exposure.

Imaging SecA after exposure to nonhydrolyzable nucleotides indicated that most of the molecules become stalled in “locked” states, but switching between states is of fundamental importance for hydrolysis. Hence, we followed individual SecA-WT molecules in real time and directly visualized conformational dynamics during ATP hydrolysis. To better observe conformational dynamics, we enhanced the temporal resolution to ~100 ms by repeatedly scanning in one dimension to generate kymographs. First, a traditional two-dimensional image was taken to identify a protein of interest (Fig. 5A). Then, the slow scan axis was disabled to repeatedly scan over the same protein to monitor conformational dynamics under physiological conditions in the absence of ATP (Fig. 5B). We then exploited the large difference evident in the protein area footprint distributions (shown in Fig. 3A) to monitor real-time dynamics as the substrate ATP was consumed.

Kymographs during ATP hydrolysis revealed at least two distinct conformational states exhibiting different width profiles that interconverted back and forth over time (Fig. 5C; see also line scans in Fig. 5D, which are separated by ~100 ms). This apparent conformational
“flickering” was captured over extended periods, illustrating the reversible nature of the dynamics.

Figure 5E illustrates a kymograph of SecA-WT in the presence of ATP over more than 75 s. Here, the protein interconverted between a “compact” and an “extended” state. The green trace in Fig. 5F shows the full width distribution profile of this molecule for the same period of time. The reversible switching of the molecule between compact and extended states is reflected in the histogram as two populations, which are separated by a minimum, positioned at ~29 nm. Additional kymograph data are presented (fig. S10). The data exhibit at least two conformational states, despite differing scan trajectories with respect to the adsorbed protein. Note that all width measurements are overestimates of true molecular dimensions due to tip convolution (nominal tip radius, ~8 nm). During ATP hydrolysis (i.e., with ATP present), the protein favored the extended state over the compact state. Specifically, it remained in the extended state for 84% of the time and 14% in the compact state. We note that the rate-limiting step of the SecA ATP hydrolysis cycle is ADP release. Hence, the prominent extended state is likely to represent the ADP-bound posthydrolysis state of SecA.

**DISCUSSION**

Single-molecule AFM studies of enzymes undergoing catalysis have been challenging, as many molecules can remain inactive during studies (34). In this work, we measured the biochemical activity of surface-adsorbed SecA prepared for AFM study. Tracking the release of Pi verified ATP hydrolysis over timescales commensurate with AFM imaging and suggested that a significant fraction of SecA was undergoing ATP hydrolysis at a similar rate to that measured in solution. With the assurance that SecA retained ATPase activity when adsorbed to mica, we applied AFM to capture conformations and conformational dynamics associated with ATP hydrolysis.

The height distribution for individual SecA-WT molecules in the apo state exhibited a maximum at around 4 nm, in overall agreement with previously reported distributions for SecA when bound to SecYEG in membrane using the same technique (7, 21). To aid interpretation, we generated simulated AFM images of SecA monomers using crystal structure coordinates and an assumed tip geometry based on the manufacturer’s nominal dimensions. Simulated images (fig. S2) were in overall agreement with the experimental data.

Exposing SecA-WT to ATP increased the areal footprint of the enzyme significantly and did so over long timescales (~4 hours). What is the origin of these long-lived ATP-dependent conformational changes? Structures have suggested that the PBD undergoes a large-scale swivel motion (approximately 80°; Fig. 1B) (1). But, it is important to note that, in the x-ray structures of SecA from *E. coli* (PDB code: 2FSF), the PBD region is not resolved. This is likely an indication that the PBD populates diverse conformations without having one state significantly more stable than the others. However,
these dynamics have proven challenging to observe. Nuclear magnetic resonance (NMR) studies on full-length SecA have provided limited information (9, 35). Molecular dynamics simulation results have suggested that the transition from open to closed state occurs in two distinct phases. Initially, the PBD and the C-terminal region move as a single unit, and then the PBD moves alone in the second phase, toward NBD2 to the fully closed state (17). AFM imaging indicated that the PBD explored a much wider conformational space when SecA was exposed to ATP compared to the apo state and provided evidence of PBD motion as well as PBD motion coupled with motion of other protein segments occurring in a nucleotide-dependent manner.

ADP binding shifted both the area and height distributions for SecA-WT but did not alter the corresponding distributions for SecAΔPBD. Hence, ADP binding directly affected the PBD conformation. Furthermore, binding of ADP-AlF$_3$, the nonhydrolyzable ATP analog that traps SecA in its transition state, shifted the height peak for both SecA-WT and the SecAΔPBD mutant to similar values. This suggested a change in conformation of the C-terminal segment, a common element that is thought to dock onto SecY at the membrane interface and regulate ATPase activity (1, 9, 17). We note that we cannot rule out the possibility that the NBD1 and NBD2 domains, which are also common to SecA-WT and SecAΔPBD, could also be responsible for observed motion. The ADP-AlF$_3$–induced height shift was accompanied by an area shift for the WT enzyme, but not for the mutant. Together, the data imply that the transition state of the system involves conformational displacements of both

Fig. 5. Reversible conformational dynamics of individual SecA during ATP hydrolysis. (A) AFM image of SecA-WT in the absence of ATP, white dashed rectangle shows area for detailed study, and asterisk identifies molecule of interest. (B) Upon rescanning and identifying the same molecule, a kymograph is instigated: the time axis begins at the top of the area (red dashed line). (C) Detailed view of kymograph in the presence of ATP. (D) Sequential height profiles separated by 95 ms (temporal sequence: pink, black, and purple) corresponding to arrows in (C). (E) Kymograph of SecA with ATP over 75 s and (F) histogram of AFM-measured SecA full width (number of scan lines, $n = 406$) showing two subpopulations: compact and extended.
hydrolyzable ATP can drive processive preprotein translocation but smallest area we observed. Previous studies have shown that only by the observation that the apo state was compact: it exhibited the compact state is likely associated with ADP release. This is supported conformational change from the dominant extended state to the 29 2018). Hence, the 29 release, and ADP release is the rate-limiting step (29, 38). In contrast, ADP release is known to in the kymographs with ATP, we propose that this state represents the ADP-bound posthydrolysis state, whereas the compact state represents one or more of steps (i) to (iii), which occur too rapidly to be resolved in our AFM measurements. The conformational change from the compact state to extended state observed in the kymographs can be associated with ATP binding and hydrolysis. A conformational change is required for ADP release, and ADP release is the rate-limiting step (29). Hence, the conformational change from the dominant extended state to the compact state is likely associated with ADP release. This is supported by the observation that the apo state was compact: it exhibited the smallest area we observed. Previous studies have shown that only hydrolyzable ATP can drive processive preprotein translocation but not ADP or other nonhydrolyzable ATP analogs (12, 42). Further, the PBD is known to be in direct contact with precursor proteins (1, 3, 4). Therefore, the two dominant states observed in the kymographs are likely associated with translocation. We studied conformational dynamics during basal SecA-ATP hydrolysis. Our measurements were carried out in real time and real space via single-molecule AFM imaging. We showed that SecA undergoes reversible conformational changes in the presence of ATP and that the proteins can be stalled in locked states upon exposure to nonhydrolyzable ATP analogs. Evidence of nucleotide-dependent motion of the PBD alone, and the PBD in conjunction with other segments of the protein, was provided. More generally, we demonstrate here a capability of AFM to study reversible conformational dynamics at the single-molecule level for the characterization of ligand-induced changes in SecA.

**MATERIALS AND METHODS**

**Protein purification**

SecA-WT was purified following standard protocols described in reference (7). The SecAΔPBD mutant (amino acids 231 to 369 deleted) is SecAC4 [a SecA variant that lacks zinc because the coordinating cysteines have been replaced by serine (26)] engineered to remove the protein-binding domain. The construct was made commercially (Mutagenex Inc.), and the protein was purified as described (7).

**ATPase activity assay**

To mimic the sample preparation for AFM measurements, SecA was preincubated for 15 min at 2.4 µM (25°C) in adsorption solution: 10 mM Hepes (pH 7.6), 100 mM KAc, 5 mM MgAc2, and 3 mM ATP. One hundred microliters of this solution was deposited onto freshly cleaved mica (Ted Pella) and incubated for 15 min at 25°C. The sample was rinsed 10 times with 100 µl of adsorption solution to remove unbound SecA. Then, 50 µl of the adsorption solution was replaced with 50 µl of solution containing radiolabeled ATP: 10 mM Hepes (pH 7.6), 100 mM KAc, 5 mM MgAc2, 3 mM ATP, 3.3 mM [γ-32P]ATP (0.908 nCi/µl; PerkinElmer). At corresponding times, 2 µl was removed to tubes with 50 mM EDTA to stop the hydrolysis reaction and applied to a thin-layer chromatography plate (MilliPore Corporation). After application of all samples, the plate was developed in 125 mM KH2PO4. After drying, the plates were exposed and scanned using a PhosphorImager. The percentage of ATP hydrolyzed was estimated from the proportion of total radioactivity that migrated as Pi. To evaluate ATPase activity in solution, the aforementioned procedure was carried out in the absence of the mica incubation and rinsing steps.

**Sample preparation for AFM**

SecA was diluted to 20 nM and incubated for 15 min in one of two solutions: (i) no substrate (apo) conditions [10 mM Hepes (pH 7.6), 100 mM KAc, and 5 mM MgAc2] or (ii) saturating nucleotide (ATP, ADP, or ADP-αβγP) conditions [10 mM Hepes (pH 7.6), 100 mM KAc, 5 mM MgAc2, and 100 µM nucleotide]. One hundred fifty microliters of this solution was then deposited onto a freshly cleaved mica surface. After 10-min incubation on mica, the sample was rinsed 10 times with 100 µl of the appropriate solution.

**Atomic force microscopy**

AFM images were acquired at ~32°C in tapping mode using a commercial instrument (Asylum Research Cypher) in 10 mM Hepes (pH 7.6), 100 mM KAc, and 5 mM MgAc2 or 10 mM Hepes (pH 7.6), 100 mM KAc, 5 mM MgAc2, and 100 µM nucleotide, as specified. Measurement–related bias of protein conformations and conformational dynamics is an ever present concern. Care was taken to control the magnitude of the tip sample force to ≤100 pN [estimated by comparing the free amplitude (~10 nm) to the set point amplitude]. BioLever mini tips (BL-AC40TS, Olympus) with measured spring constants of ~0.06 N/m were used. Spring constants were determined using the thermal noise method. AFM images were acquired with the following parameters: scan rate, 3.6 Hz; scan size, 512 pixels × 512 pixels, 1 µm2 × 1 µm2; scan speed, 7 nm/ms. Kymographs were acquired as follows: scan rate, 5.2 Hz; scan size, 320 pixels, 300 nm; scan speed, 3 nm/ms. We analyzed at least 17 images per condition studied, at least 3 independent experiments/mica stages/sample preparations per condition, and at least 2 different tips for each independent experiment. Note that tips were recovered after each experiment and cleaned for reuse.

**Data analysis**

Individual SecA molecules were identified and extracted from large-scale AFM images via the Hessian blob algorithm implemented in custom software (Igor Pro 7; WaveMetrics, Portland, OR) (22). A local...
Simulations of AFM images and related analysis

Simulated AFM images were produced (fig. S2) by modeling the interaction between a SecA molecule and the AFM tip during the imaging process. The action of “scanning” the AFM tip across a sample surface to produce an image was captured mathematically by the morphological dilation operation, which is responsible for the convolution effect seen in AFM imagery by tips that are not perfectly sharp. To construct a simulated AFM image of SecA, we explicitly computed the morphological dilation between three-dimensional models of SecA and an AFM tip. SecA was modeled via its x-ray crystal structure where each atom was modeled as a sphere with the van der Waals radius (43). The AFM tip was modeled as a cone with a rounded, spherical tip using nominal specifications from the manufacturer (cone angle, 17.5°; tip radius, 8 nm). A simulated AFM image was then produced using custom software (Igor Pro 7; WaveMetrics, Portland, OR). To compare simulated and experimental images (fig. S3), the residual between the images was squared and integrated, with the square root of the result then used to quantify the discrepancy between the images. The discrepancy was normalized as a percentage of the total volume of the experimental image. We then defined the percent agreement as one minus the normalized discrepancy. The agreement was maximized over all relative orientations (translations and rotations) between the experimental and simulated images. To generate equipotential surface maps (fig. S5), electrostatic potentials were calculated with the APBS software (#4) at pH 7.

SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/4/10/eaat8797/DC1.

Fig. S1. Tapping mode phase imaging.
Fig. S2. Simulated AFM images and likely orientations of SecA on surface.
Fig. S3. AFM image data corroborate with simulations.
Fig. S4. Statistically robust subpopulations.
Fig. S5. Equipotential surfaces comparing the front and back faces of SecA.
Fig. S6. Evaluation of molecular density of SecA prepared for surface-adsorbed ATPase activity assays.
Fig. S7. Representative images of SecA-WT and SecAΔPBD subject to different conditions.
Fig. S8. Representative images of SecA-WT and SecAΔPBD exposed to different nucleotides.
Fig. S9. Height histograms of SecA immersed in and exposed to nonhydrolyzable nucleotides.
Fig. S10. Additional kymograph data.
Table S1. Statistics of SecA-WT (WT) and SecAΔPBD area distributions in the following conditions.
Table S2. Mean height (±SEM) and FWHM of the SecA-WT and SecAΔPBD mutant height distributions in different conditions.
Table S3. Statistics of the SecA-WT and SecAΔPBD area distributions after exposure to ADP or ADP-AlF4.
Table S4. Mean height (±SEM) and FWHM of the SecA-WT and SecAΔPBD mutant distributions exposed to ADP and ADP-AlF4.

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