Bacterial DnaB helicase interacts with the excluded strand to regulate unwinding

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This article contains Supplemental Table S1.

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Replicative hexameric helicases are thought to unwind duplex DNA by steric exclusion (SE) where one DNA strand is encircled by the hexamer and the other is excluded from the central channel. However, interactions with the excluded strand on the exterior surface of hexameric helicases have also been shown to be important for DNA unwinding, giving rise to the steric exclusion and wrapping (SEW) model. For example, the archaeal Sulfolobus solfataricus minichromosome maintenance (SsoMCM) helicase has been shown to unwind DNA via a SEW mode to enhance unwinding efficiency. Using single-molecule FRET, we now show that the analogous Escherichia coli (Ec) DnaB helicase also interacts specifically with the excluded DNA strand during unwinding. Mutation of several conserved and positively charged residues on the exterior surface of EcDnaB resulted in increased interaction dynamics and states compared with wild type. Surprisingly, these mutations also increased the DNA unwinding rate, suggesting that electrostatic contacts with the excluded strand act as a regulator for unwinding activity. In support of this, experiments neutralizing the charge of the excluded strand with a morpholino substrate instead of DNA also dramatically increased the unwinding rate. Of note, although the stability of the excluded strand was nearly identical for EcDnaB and SsoMCM, these enzymes are from different superfamilies and unwind DNA with opposite polarities. These results support the SEW model of unwinding for EcDnaB that expands on the existing SE model of hexameric helicase unwinding to include contributions from the excluded strand to regulate the DNA unwinding rate.

Hexameric helicases are structurally conserved toroidal enzyme complexes capable of translocating and separating double-stranded DNA (dsDNA) into two single strands (ssDNA), providing templates for DNA replication. They utilize the inherent energy from ATP hydrolysis to translocate along an encircled strand, physically displacing the opposing excluded strand. The translocation polarity of hexameric helicases differs among helicase superfamilies (SFs), defined by the organization and conservation of various folds (1). SF4 helicases from bacteria and associated phages (T4 and T7) include RecA-like folds and have 5′–3′ unwinding polarity, translocating on the lagging strand, whereas SF6 helicases from archaea and eukaryotes have AAA+ folds and 3′–5′ unwinding polarity, translocating on the leading strand (2). Although these two well-studied helicase families have globally conserved structural features, their amino acid sequences, structural folds, and unwinding polarities are not. Moreover, the precise contacts with each DNA strand to facilitate duplex unwinding are not known.

The bacterial replicative helicase, DnaB, has been shown to encircle the 5′-lagging strand in its central channel. Orientation of DnaB binding on ssDNA was shown to place the RecA motor C-terminal domain (CTD) adjacent to the duplex region and the N-terminal domain (NTD) toward the 5′-end (3). DnaB unwinds dsDNA in a steric exclusion (SE) mechanism and can even translocate over two or three strands of DNA, indicating plasticity within the central channel of the hexamer (4, 5). Currently, various X-ray and EM structures of hexameric DnaB (with and without DNA or accessory proteins) show the hexamer as either a closed ring (6–8) or a split lock washer (9, 10). ssDNA bound in the central channel adopts an A-form right-handed spiral conformation making contacts with multiple interior DNA-binding loops from different subunits to pass DNA along in a hand-over-hand mechanism (9).

In addition to specific DNA contacts that exist within the central channel, further exterior contact sites are proposed to exist for hexameric helicases to aid in DNA loading and unwinding (2). Previously, we have identified an interaction path on the external surface of the archaeal (AAA+) SsoMCM helicase with the excluded strand that both protects and stabi-
lizes the complex in a forward unwinding mode (11, 12). This interaction expanded the widely accepted SE model of unwinding to include contributions of the excluded strand in the mechanism. This new unwinding model was termed steric exclusion and wrapping (SEW). Recently, interactions with the excluded strand have been uncovered from a variety of hexameric helicase complexes in addition to archaeal MCM including papillomavirus E1 (13), SV40 large T (14), T7gp4 (15, 16), \( \text{EcDnaB} \) (17, 18), TWINKLE (19), and the eukaryotic Cdc45/MCM2–7/ GINS complex (20, 21). It is hypothesized that external interactions with the excluded strand will not only protect ssDNA but also stabilize the helicase/DNA complex and modulate the unwinding rate.

In this study, we examined whether \( \text{EcDnaB} \) has a similar specificity for exterior interactions with the excluded single-strand DNA. Single-molecule FRET (smFRET) experiments were used to directly detect \( \text{EcDnaB} \) binding to the excluded strand and to compare it with \( \text{SsoMCM} \) binding. The absolute FRET states, transition probabilities, and dwell times were strikingly similar between \( \text{EcDnaB} \) and \( \text{SsoMCM} \) even though they reside in different superfamilies and have opposite unwinding polarities. Mutation of several conserved external positively charged residues on \( \text{EcDnaB} \) differentially altered the observed FRET states and binding dynamics, consistent with disruption of the excluded strand binding path. Notably, those external SEW mutations on \( \text{EcDnaB} \) dramatically increased the dsDNA unwinding rate compared with wild type (WT). Neutralizing the negative charge on the excluded strand with a morpholino substrate increased the unwinding rate for wild-type \( \text{EcDnaB} \), consistent with electrostatic interactions regulating activity. This is the first molecular explanation for regulation of the rate of DNA unwinding through specific external surface interactions on the helicase with the excluded strand.

### Results

**EcDnaB interacts with the excluded strand**

Previously, we have shown that interactions with the excluded strand exist for the 3′–5′ hexameric \( \text{SsoMCM} \) helicase (11, 22). Using a similar smFRET approach, we sought to examine whether analogous contacts on the exterior of \( \text{EcDnaB} \) also interact with the excluded strand despite the opposing 5′–3′ translocation polarity. Three separate model fork substrates, 30/30 (DNA43/DNA44), 40/30 (DNA111/DNA44), and 50/30 (DNA116/DNA44) composed of an 18-bp duplex with (dT)30 on the 5′-lagging strand and (dT)30, (dT)40, and (dT)50 on the 3′-leading strand, respectively, were used. DNA forks alone result in low FRET signals as a result of Cy3 and Cy5 on the termini of the fork arms not being in close proximity (Fig. 1A). Additional nucleotides on the leading strand 3′-arm of the fork further decrease the FRET efficiency as expected. Addition of \( \text{EcDnaB} \) to each of these substrates shifts the signal to higher FRET states (Fig. 1B). \( \text{EcDnaB} \) preferentially encircles the 5′-strand (3), and a titration of \( \text{EcDnaB} \) onto 30/30 showed little to no variation in the resulting histogram profiles, suggesting that only one hexamer can be accommodated by the fork substrate over a large concentration range (Fig. 1C). Should a second hexamer encircle the opposing 3′-strand, the FRET values would decrease. The occurrence of a high-FRET state is consistent with an interaction of the excluded Cy3 3′-strand on the external surface of \( \text{EcDnaB} \), analogous to \( \text{SsoMCM} \) binding to DNA and consistent with the proposed SEW model of helicase interaction and unwinding (11).
EcDnaB loaded onto the 40/30 substrate produced an almost exclusively high-FRET state (>0.9), the 50/30 fork produced a bimodal distribution of high- (>0.9) and medium (~0.5)-FRET states, and the 30/30 fork yielded a bimodal distribution of two high-FRET (0.8 and >0.9) states in the presence of EcDnaB. The interaction with the excluded strand likely differs from varying excluded strand lengths sampling slightly different external binding paths. However, in all cases, the shifts to high-FRET states correspond to a stably wrapped excluded strand that places the Cy3 dye near the Cy5 dye on the encircled strand.

Explicit probability and rate transition (ExPRT) analysis of excluded strand dynamics

To better visualize the FRET states, explicit transition probabilities, and dwell times on a single plot to compare different conditions, mutants, and helicases, we developed ExPRT plots (Fig. 2A). The positions of the circular markers correspond to transitions between specific FRET states with the initial FRET state on the x axis and the final FRET state on the y axis. The initial and final FRET states for a particular transition refer to the observed FRET states immediately preceding and immediately following the transition of interest, respectively. The size and color of each marker correspond to the probability of that transition occurring within a measured trace and the average dwell time of the state preceding the transition, respectively.

A comparison of analyses between established programs, the HaMMY and transition density plot (TDP) programs (Fig. 2B) (23) as well as the population-weighted and kinetically indexed transition (POKIT) density program (Fig. 2C) (24), and the ExPRT program is shown. Each program analyzed and visualized identical data corresponding to EcDnaB bound to the 30/30 substrate and can distinguish states (Fig. 2D). Each plot illustrates that the transitions between FRET states of ~0.8 and ~0.95 are the most frequent. The TDP program analysis works on a trace-by-trace basis and is able to reveal heterogeneities in the transition data that can be missed by programs that work on stitched data sets such as the POKIT and ExPRT programs. However, despite the TDP program’s ability to gather probability and rate values, these values are not directly visualized by the resulting plot. The POKIT program bins the probabilities and rates of each transition into user-defined ranges and produces plots that allow for some level of quantitative comparison between experimental conditions. However, these plots fail to display explicit probabilities and rate values. Determining explicit transitions, probabilities, and rates without binning is an inherent advantage of single-molecule methods that allows for extensive insights into the dynamics and kinetics of molecular interactions and enzymatic activities. The ExPRT analysis program extracts these explicit values from the smFRET data and visualizes them directly in a single plot. This allows users and readers to easily make comparisons between data sets on the most detailed level. Therefore, the ExPRT plots provide a useful advance in the investigation and comparison of the probability and kinetics of smFRET dynamics.

EcDnaB and SsoMCM interact with the excluded strand similarly

The bimodal distribution observed for EcDnaB on 30/30 fork substrate was very similar to the distribution produced by the archaeal MCM helicase on the same substrate (Fig. 3A) (11). In fact, the single-molecule traces for EcDnaB and SsoMCM exhibited strikingly similar dynamics between two high-FRET states (~0.95 and ~0.8) (Fig. 3, B and C). This is highlighted by the ExPRT plots for EcDnaB compared with SsoMCM on the 30/30 fork substrate (Fig. 3, D–F). In both cases, the excluded strand has a reversible transition between two high-FRET states (~0.8 and ~0.95) that is exhibited by ~70% of the molecules analyzed. For both EcDnaB and SsoMCM, there is a preference
for the ~0.95 FRET state, indicated by the longer dwell times measured for that state (shades of orange versus green/yellow). In addition, both data sets exhibit a reversible lower probability transition between each of the high-FRET states and a medium-FRET state of ~0.55. Altogether, it is noteworthy that similarities in the FRET states, transitions, and dwell times exist for EcDnaB and SsoMCM on the 30/30 fork even though these two helicases belong to different superfamilies with low sequence homology, exist in different domains of life (Bacteria versus Archaea), and have opposite unwinding polarities. The similarities are suggestive of a common SEW unwinding mechanism across diverse replicative helicases.

**Exterior surface mutations of EcDnaB alter excluded strand wrapping**

As positively charged residues on the surface of SsoMCM had previously been shown to support an external interaction with the excluded strand (11), similar surface-exposed and conserved residues were identified based on a homology model for EcDnaB (Fig. 4). Four EcDnaB surface positions (Arg-74, Arg-164, Lys-180, and Arg-328/Arg-329) that exist in positively charged electrostatic patches were mutated to alanine, overexpressed, and purified. All mutant proteins were consistent with a hexamer as the major peak after gel filtration (data not shown). smFRET DNA fork binding assays were performed for the wild type and each mutant on each of the three fork templates (30/30, 40/30, and 50/30). The results were analyzed and compared using traditional histograms and ExPRT plots. The mutations gave rise to several important differences in the hexamer-excluded strand interactions and dynamics. Across all the excluded strand lengths tested, EcDnaB (R74A) does not sample the highest observed FRET state ($E_{\text{app}} = 0.95$) observed for the WT and the other mutants. This can be seen clearly in the histograms (Fig. 5, B versus A) and the ExPRT plots (Fig. 5, G, L, and Q versus F, K, and P). R74A also produces fewer transitions between FRET states compared with the WT for the 30/30 and 40/30 forks because the most frequent transitions observed for the WT data are those between the highest FRET state ($E_{\text{app}} = 0.95$) and lower FRET states. The absence of the 0.95 FRET state for R74A across all substrates tested suggests that Arg-74 is necessary to close the connection of the excluded strand to the NTD traversing the entire longitudinal length of EcDnaB (9).

![Figure 3. Comparison of the excluded strand interactions of SsoMCM and EcDnaB by smFRET.](image)
The EcDnaB (R328A/R329A) mutant shows similar FRET state transitions, probabilities of those transitions, and dwell times on the 30/30 fork when compared with the WT (Fig. 5, I versus F). However, there are extreme differences in the binding states and dynamics on the longer DNA strands compared with WT (Fig. 5, T versus P). WT EcDnaB bound to the 50/30 fork shows a small fraction of traces that transition between high- and medium-FRET states. In comparison, EcDnaB (R328A/R329A) produces almost entirely medium-FRET states that are very dynamic with many transitions and relatively short dwell times, indicative of severe destabilization of binding. In contrast to WT, very little high-FRET signal from the R328A/R329A mutant on the 30/50 substrate is observed. These results indicate that Arg-328 and Arg-329 may be required to stabilize longer excluded strands (40- and 50-mers) along the waist of the hexamer and mediate interactions between the excluded strand and other regions such as NTD where Arg-74 is located and responsible for the highest FRET state ($E_{\text{app}} \approx -0.95$).

Although we primarily tested the effect of eliminating positive charge on the exterior surface of EcDnaB and its effect on excluded strand binding, there may also be additional noncovalent binding interactions defining a path. To directly test whether electrostatic interactions exclusively define the excluded strand binding path, we titrated NaCl into WT EcDnaB prebound to a 30/30 fork in our smFRET experiments. Increased salt concentrations resulted in increased dynamics (shorter dwell times), but the FRET states were not significantly affected as visualized using ExPRT plots (Fig. 7). Even though the FRET states and transition probabilities remained constant, the decreased dwell times suggest that EcDnaB utilizes electrostatic interactions to mediate wrapping but that other noncovalent interactions are also important.

**SEW mutants of EcDnaB have enhanced DNA unwinding activity**

Mutating positively charged residues involved in the excluded strand interaction inhibited SsoMCM's unwinding activity (11, 22). Gel-based fluorescent DNA unwinding assays were performed to determine whether these EcDnaB SEW mutants have any effect on activity. Fig. 8A shows a representative 6-min time point; however, quantification of the steady-state unwinding rates occurred over multiple time points for each mutant (Fig. 8B). All mutants had increased unwinding rates compared with the WT EcDnaB. Specifically, R74A (26 ± 1 nm s$^{-1}$) and R164A (53 ± 13 nm s$^{-1}$) have 3- and 6-fold increases, respectively, over WT (9 ± 1 nm s$^{-1}$); whereas K180A (199 ± 16 nm s$^{-1}$) and R328A/R329A (191 ± 13 nm s$^{-1}$) have more than 20-fold increases in unwinding activity.

ATPase assays were performed for WT and mutant EcDnaB proteins, and the rates were quantified in the absence and presence of DNA (Fig. 8C). R74A, K180A, and R328A/R329A have similar basal rates to WT; however, R164A had a 2.5-fold enhancement over WT. Similar to WT EcDnaB, both R74A and R164A were stimulated 1.5–2.0-fold in the presence of DNA, consistent with previous results (25). Interestingly, the fastest unwinding mutants, K180A and R328A/R329A, were only weakly stimulated further in the presence of DNA. Previously, the R328A/R329A mutant was investigated for its potential role...
DnaB-excluded strand interaction

in a leucine zipper motif (26), although it was later discounted (27) and also found not to have DNA-stimulated ATPase activity. No stimulation in ATPase rate with DNA is sometimes indicative of a perturbation in DNA binding; however, these mutants show stimulated unwinding abilities, and fluorescence anisotropy experiments showed no significant differences in $K_d$ values measured for mutants binding to fork DNA compared with WT (data not shown).

To validate whether alteration of electrostatic interactions are responsible for the increased unwinding rates in the mutants compared with WT EcDnaB, we performed DNA unwinding reactions with a 3′-morpholino (Morph) strand. Morpholino nucleic acids have standard base pairing properties but have morpholine rings linked through phosphorodiamidate groups that lack negative charge and are as stable or more stable than an equivalent DNA duplex (28, 29). Previously, the homologous hexameric T7 gp4 DNA helicase was shown to unwind excluded strand morpholino substrates with a greater rate and efficiency than for DNA (16). They attributed this enhanced unwinding activity to the disruption of the helicase’s interaction with the displaced strand that limits its activity.

Interestingly, WT EcDnaB also unwinds excluded strand morpholino substrates with a profoundly enhanced rate compared with a DNA/DNA duplex (Fig. 9). The rate of unwinding for the Morph/DNA is at least 1000-fold faster with a $>0.7$ amplitude after 1 min, whereas the DNA/DNA duplex is only $>0.4$ unwound after 45 min. No unwinding or strand separation is seen when ATP is excluded from the experiment. Accurate quantification of the unwinding rate would require rapid quench experiments, but it is apparent that EcDnaB unwinds excluded strand morpholinos rapidly. The unwinding rates for the Morph/DNA are also significantly faster ($>10$-fold) than even those seen above for the fastest SEW mutant (K180A) on a DNA fork (Fig. 8). This may not be surprising as the SEW mutants only affect contact at one specific mutated site on the exterior surface, whereas the excluded strand Morph eliminates electrostatic contacts throughout the longitudinal length of EcDnaB.

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**Figure 5.** Histograms and ExPRT plots of WT EcDnaB and mutants bound to DNA forks. Histograms (A–E) report the population of molecules as a function of FRET states on DNA forks with a 30-base 5′-strand and a 30-base (blue), 40-base (green), or 50-base (red) 3′-strand for WT, R74A, R164A, K180A, and R328A/R329A, respectively. Yellow, blue, and red regions highlight low-, medium-, and high-FRET populations, respectively. Corresponding ExPRT plots are shown for 30/30 (F–J), 40/30 (K–O), and 50/30 (P–T) forks for each of the respective EcDnaB helicases.
Discussion

Although hexameric DNA replication helicases have global structural conservation, their amino acid sequences are not conserved, allowing for the classification of these helicases into different superfamilies. We have shown previously that the 5’-excluded strand makes important external surface interactions that aid in the mechanism of unwinding for the SF6 archaeal SsoMCM helicase (3’–5’) to develop the SEW of unwinding (11). In this report, we can now show that the bacterial replication helicase, EcDnaB, with opposite unwinding polarity (5’–3’) and of a different family (SF4) and organismal domain, has similar conformational states and dynamics of binding the excluded strand that also regulate DNA unwinding (Fig. 10). The combined results highlight the importance and conservation of the SEW model for hexameric helicase unwinding of DNA and reveal external surface residues required for regulating the activity of the EcDnaB helicase.

It is striking that the absolute FRET states, transition probabilities, and dwell times visualized by the ExPRT plots are extremely similar between SsoMCM and EcDnaB hexameric helicases bound to fork DNA. In both cases, the large increase in FRET observed is consistent with encircling of the translocating strand and exclusion of the other along the exterior surface. For EcDnaB, there have been reports of hexamers loading on opposing strands in opposite orientations (3, 30–32). The consequence of loading two hexamers would ultimately separate the strands further, resulting in a decrease in the FRET signal, which is opposite to what we observe in the smFRET experiments even at high concentrations of EcDnaB (Fig. 1B). Both helicases are of similar size and oligomeric state and are thought to engage their respective translocating strands in a similar way, so the DNA-bound states of each helicase may be structurally equivalent even with opposing translocation polarities. Therefore, EcDnaB binding of DNA includes both the encircling of the 5’-strand and the exclusion and external interaction of the 3’-strand in a similar manner to SsoMCM and the SEW model for unwinding.

Without an appropriate DNA-bound crystal structure of EcDnaB, we had to infer binding positions for the excluded 3’-strand based on amino acid homology and electrostatics from crystal structures that represented a closed ring (6, 7) or a split lock washer structure (9). Because ssDNA was contained in the central channel of the split lock washer structure, we used this conformation as a primary model to interpret interactions with the excluded strand. This restricted the definition of any precise or specific exterior binding path, and rather we can only conclude general binding to the CTD, waist, and NTD. That along with the specific residues that were mutated and the smFRET data informed our interpretation of excluded strand binding.

Figure 6. Example smFRET kinetic traces. A–F, comparison of WT (A–C) and K180A (D), R74A (E), and R328A/R329A (F) EcDnaB FRET efficiencies as a function of time on the 30/30, 40/30, and 50/30 forks, respectively. The calculated FRET values (blue) are overlaid with the ideal state fits (red).
To test the specificity of this external interaction, mutations of conserved residues were found to both disrupt and alter the binding states. In particular, Arg-74 was found to be necessary for stable interactions of the excluded strand at the NTD, giving rise to the highest FRET state ($0.95$). R164A and K180A exhibited somewhat different dynamics than WT, generally fitting to more FRET states, which may reflect alternate binding paths on the helicase exterior. It is likely that these two residues...

Figure 2. [Diagram of NaCl titration onto EcDNA B-bound fork. A–D] ExPRT plots of [NaCl] titration onto EcDNA B (250 nM) prebound to 30/30 DNA fork.

Figure 3. [Gel Unwinding Rate] A, [ATPase] DNA unwinding (nM.min$^{-1}$) B, ATPase (nM.s$^{-1}$)

Figure 8. [Biochemical properties of SEW mutants of EcDNA B] A and B, a representative gel for 6-min time point is shown for EcDNA B (WT and mutants, 3 µM) unwinding assays performed on a fluorescein-labeled fork DNA (DNA14F/DNA15) (15 nM) (A) and quantified over multiple time points (B) is shown. Throughout, data for EcDNA B constructs are consistently colored (WT, red; R74A, ochre; R164A, orange; K180A, green; R328A/ R329A, blue). Error bars represent the standard error from at least three independent experiments.

Figure 9. [Unwinding of morpholino forked substrates] A and B, a representative unwinding time course for 250 nM EcDNA B on either an 18-bp DNA/DNA (D:D) (A) or Morph/DNA (M:D) (B) substrate (15 nM) with a 7-base 3'-excluded strand flap is shown. Single-turnover experiments were initiated with ATP and a single-strand trap identical to the radiolabeled strand as described under “Experimental procedures.” C, averaged unwinding data for DNA/DNA (black C) were plotted and fit with a linear regression to give $k_1 = 0.009$ min$^{-1}$, and those for Morph/DNA (gray C) were plotted and fit using Equation 3 to give $k_1 = 11$ min$^{-1}$. The inset plot highlights data within the 1st min. Error bars are the standard error from three independent experiments.
unwinding of morpholino strands was also seen with the homologous T7 gp4 hexameric helicase (16). In both cases, interactions on the outer surface of the helicase with the excluded strand will act to regulate the unwinding rate. However, this external interaction is not entirely electrostatic as increasing ionic strength in the smFRET experiments resulted in decreased dwell times but did not significantly affect the FRET states.

Previous single-molecule work has detailed the single-molecule force contributions of each DNA strand to unwinding by EcDnaB using either a hairpin or fork substrate (33). In that study, it was concluded that the unwinding rate was controlled by both force-induced destabilization of the duplex and interactions of the excluded strand with the exterior surface. The main apparent discrepancy between our work and theirs is that when the excluded strand is sequestered because of constraints in the hairpin assay the rate is slower than when it is allowed to interact with the exterior surface in the fork assay. This would imply that contacts of the excluded strand with the external surface of EcDnaB increase unwinding, whereas we show that specific external contacts restrain unwinding. However, it is probable that the force applied to the excluded strand in the fork assay artificially alters the interaction with the exterior surface in a way analogous to the altered DNA binding paths and kinetics for the R74A and R164A mutants. Therefore, measured increases in unwinding in both studies can be explained by altered DNA binding paths on the exterior surface. It is notable that a variety of recent biophysical techniques monitoring EcDnaB activity and binding have detected an elusive external interaction with the excluded strand (18, 33). We can now conclude that this SEW interaction and the precise binding path regulate the speed of unwinding for EcDnaB.

Clearly, interactions with the excluded strand are acting as a regulator to control the speed of unwinding. It remains to be seen whether this is because of a greater increased force applied by the motor domain for EcDnaB that is modulated by the excluded strand or whether discrete external binding paths or polarity dictates the rate of unwinding. Further experiments will be needed to more specifically define the exterior binding path. For EcDnaB, the excluded strand interaction may act as a “molecular brake” to control the amount of exposed ssDNA or provide a platform for accessory helicases, e.g., Rep, to assemble and rescue stalled forks (34, 35). Coupled DNA synthesis by the leading strand polymerase (polymerase III) could sequester the excluded strand from the exterior surface of EcDnaB and explain the increased rate of unwinding by the coupled replisome (36–38). In fact, a recent report has shown that when the bacterial helicase and polymerase become decoupled, the unwinding rate is reduced by 80% as a fail-safe “dead man’s switch” (39). This can be explained at the molecular level by our data in which decoupling of the polymerase engages the excluded strand with the exterior surface of EcDnaB, slowing its unwinding progression. In addition to controlling the unwinding rate, the external DNA-binding sites on both helicases are likely to contribute during the loading mechanism for encircling of the translocating strand to maintain strand separation during the action and conformational changes induced by the initiation enzymes.
DnaB-excluded strand interaction

Experimental procedures

Materials

Oligonucleotides used (supplemental Table S1) were purchased from Integrated DNA Technologies, Inc. (Coralville, IA). Fluorescently labeled DNA was HPLC-purified (Integrated DNA Technologies, Inc.), and non-labeled oligos were gel-purified (40). Morpholinos were from GeneTools (Philomath, OR). SooMCM was purified as described previously and reported as hexamer concentrations (11). All other chemicals were analytical grade or better.

Cloning and protein purification of EcDnaB

The R74A, R164A, K180A, and R328A/R329A mutations of EcDnaB were created by overlap extension from pET11b-EcDnaB. The DNA primers are listed in supplemental Table S1. Mutations were confirmed by the DNA sequencing facility at the University of Pittsburgh. Mutants and WT EcDnaB were expressed in Rosetta 2 cells (EMD Millipore, Billerica, MA) and C41 cells (Lucigen, Middleton, WI) using autoinduction (41) or by induction with 0.1 mM isopropyl β-D-1-thiogalactopyranoside. Cells were pelleted, resuspended in EcDnaB lysis buffer (10% sucrose, 50 mM Tris-HCl (pH 7.5), 50 mM NaCl, 5 mM dithiothreitol (DTT)), and lysed using lysozyme and sonication. Ammonium sulfate was added to the resulting supernatant at 0.2 g/ml, pelleted, and then resuspended in EcDnaB buffer A (10% glycerol, 0.1 mM EDTA, 50 mM Tris-HCl (pH 7.5), 50 mM NaCl, 5 mM DTT). The supernatant was purified using an AKTA Prime FPLC equipped with a HiTrap Mono Q column (GE Healthcare) and eluted with a stepwise gradient of buffer A with 500 mM NaCl followed by a similar procedure using the University of Pittsburgh. Mutants and WT EcDnaB were analytical grade or better.

Single-molecule fluorescence resonance energy transfer

DNA substrates labeled with Cy3 and Cy5 fluorophores were immobilized on a PEGylated quartz slide utilizing biotin-streptavidin interactions (43). A prism-based total internal reflection microscope was used to collect all smFRET data (44, 45). A 532 nm diode laser was used to excite Cy3 fluorophore, and subsequent Cy3 and Cy5 emission signals were separated by a 610-nm dichroic long-pass mirror, a 580/40-nm bandpass filter, and a 660-nm long-pass filter. An electron-multiplying charge-coupled device iXon camera (Andor, Belfast, UK) was used to image the signals. Data were acquired at 10 fps for 10 or more regions with each region containing 50–250 molecules in the presence of an oxygen radical-scavenging solution (1 mg/ml glucose oxidase, 0.4% (w/v) D-glucose, 0.04 mg/ml catalase) and 2 mM trolox. EcDnaB (250 nM) was added and given a 5-min equilibration period. All single-molecule experiments were performed in reaction buffer as described previously (11).

Single-molecule FRET signals were identified by fitting individual regions of signal intensity to a 2D Gaussian function and measuring the goodness of fit. These peaks were corrected for thermal drift and local background intensity using ImageJ (46) and the image stabilizer plugin for ImageJ. The resulting signal was used to calculate the apparent FRET efficiency, E_{app}, according to the following equation.

\[
E_{app} = \frac{I_A}{I_A + I_D}
\]

in which I_A and I_D are the intensity of the acceptor and donor signals, respectively.

Single-molecule FRET data analysis and ExPRT plots

Data analysis and visualization were performed using manually selected single-molecule traces that displayed anti-correlation between the donor and acceptor fluorophores and single-step fluorophore photobleaching. Traces collected under identical experimental conditions were stitched together and fit to ideal states via hidden Markov modeling using the vbFRET software package (48). Stitched traces were fit to a given number of states based on those states being more than E_{app} = 0.1 tk;4apart from one another and the variation of one state not overlapping with another. Traces were then unstitched and fed into the ExPRT analysis program. This MATLAB executable program produces transition plots where the markers are sized based on the probability of transition occurring within an observed single-molecule trace and colored based on the dwell time(s) of the state preceding the transition. The average of all dwell times for a given transition was used to determine the color of the marker. Only dwell times that were both preceded and followed by transitions were included. Stitched data as fit by vbFRET were also analyzed by the POKIT analysis program, and the resulting plot contains a legend for the ranges of rates and probabilities (24). The data were also fit using the HaMMY program, allowing the program to fit the data to up to five states. The output of the HaMMY program was subsequently analyzed by the transition density plot program (23).

EcDnaB structural homology model

Global sequence alignments were performed using ClustalW2 analysis (http://www.ncbi.nlm.nih.gov/blast/bl2seq/wblast2.cgi). The homology model of EcDnaB was created by threading the alignment onto the structure of Geobacillus stearothermophilus DnaB (Protein Data Bank code 4ESV) (9) using SWISS-MODEL (49).

ATPase assay

EcDnaB variants (350 nM) were incubated in the absence and presence of 4 μM forked DNA (DNA14/DNA15) as described previously (47). Briefly, 25-μl reactions were incubated at 37 °C for 5 min in unwinding buffer (50 mM HEPES (pH 8.0), 10 mM Mg(OAc)₂, 5 mM DTT, 0.2 mg/ml BSA), and 1 mM ATP with trace amounts of [γ-32P]ATP was added to initiate the reaction. Samples were quenched at 2, 5, 10, and 15 min after initiation in equal volumes of 0.7 M formic acid. A total of 1 μl of quenched reaction was spotted on Millipore TLC PEI Cellulose F, allowed to dry, resolved in 0.6 M potassium phosphate.
substrates at 37 °C. The 5'-DNA163/DNA161 or 3'-mM EDTA (pH 8.0), 2 mg/ml Proteinase K) followed by incubation at various time points from 1 to 15 min. Reactions were kept on ice until loading and were resolved on 20% (29:1 acrylamide:bisacrylamide) tris borate-EDTA gels electrophoresed in tris borate-EDTA buffer. The gels were imaged on a Typhoon 9400 phosphorimaging system (GE Healthcare), and the fraction unwound was calculated after background subtraction using ImageQuant software according to the following equation.

\[
F = \left( \frac{l_{s(t)} - l_{s(0)}}{l_{s(t)} + l_{s(0)}} \right) \left( \frac{l_{d(t)} - l_{d(0)}}{l_{d(t)} + l_{d(0)}} \right)
\]

where \(I_{s(t)}\) and \(I_{d(t)}\) are the intensities of the single- and double-strand bands, respectively, at time \(t\); subscript \(0\) and \(b\) indicate equivalent counts at \(t = 0\) and the boiled sample, respectively. The fraction unwound or unwinding rate (nmol min\(^{-1}\)) was calculated from a linear regression fit of the fraction unwound for each time point.

**Single-turnover DNA unwinding assays**

Single-turnover unwinding experiments were also performed by initiating the reaction simultaneously with 5 mM ATP and 150 mM ssDNA trap (unlabeled strand with the same sequence as radiolabeled strand) at various distances. The reaction was quenched using an equal volume of quench solution (20% Ficoll, 1.0% SDS, 200 mM ssDNA trap (unlabeled strand with same sequence as radiolabeled strand)) at various time points from 1 to 15 min. Reactions kept on ice until loading and were resolved on 20% (29:1 acrylamide:bisacrylamide) tris borate-EDTA gels electrophoresed in tris borate-EDTA buffer. The gels were imaged on a Typhoon 9400 phosphorimaging system (GE Healthcare), and the fraction unwound was calculated after background subtraction using ImageQuant software according to the following equation.

\[
F = C + A_1(1 - e^{-k_{ss}t}) + k_{ss}t
\]

where \(A_1\) is the amplitude associated with the initial burst rate increase \(k_{ss}\) is a steady-state rate, and \(C\) is a constant.

**Author contributions**—S. M. C., S. H. L., and M. A. T. designed the experiments. S. M. C. performed the cloning, protein purification, biochemical assays, and single-molecule experiments. S. M. C. and S. G. performed DNA unwinding assays and protein purifications. M. A. T. directed the project and together with S. M. C. wrote the manuscript. All authors edited the manuscript.

**References**

1. Thomsen, N. D., and Berger, J. M. (2009) Running in reverse: the structural basis for translocation polarity in hexameric helicases. *Cell* **139**, 523–534
2. Trakselis, M. A. (2016) Structural mechanisms of hexameric helicase loading, assembly, and unwinding. *F1000Res. 5*, F1000 Faculty Rev-111
3. Zewitzka, M. J., Rajendran, S., Bujalskia, D., and Bujalski, W. (1998) Does single-stranded DNA pass through the inner channel of the protein hexamer in the complex with the *Escherichia coli* DnaB Helicase? Fluorescence energy transfer studies. *J. Biol. Chem. 273*, 10515–10529
4. Kaplan, D. L., and O’Donnell, M. (2004) Twin DNA pumps of a hexameric helicase provide power to simultaneously melt two duplexes. *Mol. Cell* **15**, 453–465
5. Kaplan, D. L., and O’Donnell, M. (2002) DnaB drives DNA branch migration and dislodges proteins while encircling two DNA strands. *Mol. Cell* **10**, 647–657
6. Bailey, S., Eliason, W. K., and Steitz, T. A. (2007) Structure of hexameric DnaB helicase and its complex with a domain of DnaG primase. *Science 318*, 459–463
7. Wang, G., Klein, M. G., Tokonzaba, E., Zhang, Y., Holden, L. G., and Chen, X. S. (2008) The architecture of a DnaB-family replicative helicase and its interactions with primase. *Nat. Struct. Mol. Biol. 15*, 94–100
8. Liu, B., Eliason, W. K., and Steitz, T. A. (2013) Structure of a helicase-helicase loader complex reveals insights into the mechanism of bacterial primosome assembly. *Nat. Commun. 4*, 2495
9. Itsathipaisarn, O., Wing, R. A., Eliason, W. K., Wang, I., and Steitz, T. A. (2012) The hexameric helicase DnaB adopts a nonplanar conformation during translocation. *Cell 151*, 267–277
10. Arias-Palomino, E., O’Shea, V. L., Hood, I. J., and Berger, J. M. (2013) The bacterial DnaC helicase loader is a DnaB ring breaker. *Cell 153*, 438–448
11. Graham, B. W., Schauer, G. D., Leuba, S. H., and Trakselis, M. A. (2011) Structural and functional analysis of the Escherichia coli DnaB helicase loader complex reveals insights into the mechanism of bacterial primosome assembly. *F1000Res. 10*, e655–e695
12. Rothenberg, E., Trakselis, M. A., Bell, S. D., and Ha, T. (2007) MCM forked substrate specificity involves dynamic interaction with the 5’-tail. *J. Biol. Chem. 282*, 34229–34234
13. Lee, S. J., Syed, S., Enemark, E. J., Schuck, S., Stenlund, A., Ha, T., and Joshua-Tor, L. (2014) Dynamic look at DNA unwinding by a replicative helicase. *Proc. Natl. Acad. Sci. U.S.A. 111*, E827–E835
14. Huang, H., Zhao, K., Arnett, D. R., and Fanning, E. (2010) A specific docking site for DNA polymerase α-prime on the SV40 helicase is required for viral primosomal activity, but helicase activity is dispensable. *J. Biol. Chem. 285*, 33475–33484
15. Satapathy, A. K., Kuczyk, A. W., Ghosh, S., van Oijen, A. M., and Richardson, C. C. (2011) Coupling dTTP hydrolysis with DNA unwinding by the DNA helicase of bacteriophage T7. *J. Biol. Chem. 286*, 34468–34478
16. Jeong, Y. J., Rajagopal, V., and Patel, S. S. (2013) Switching from single-stranded to double-stranded DNA limits the unwinding processivity of the ring-shaped T7 DNA helicase. *Nucleic Acids Res. 41*, 4219–4229
17. Malhat, C. M., and McHenry, C. S. (2015) Identification of subunit binding positions on a model fork and displacements that occur during sequential assembly of the *Escherichia coli* primosome. *J. Biol. Chem. 290*, 10828–10839
18. Galletto, R., Zewitzka, M. J., and Bujalski, W. (2004) Unzipping mechanism of the double-stranded DNA unwinding by a hexameric helicase: the effect of the 3’ arm and the stability of the dsDNA on the unwinding activity of the *Escherichia coli* DnaB helicase. *J. Mol. Biol. 343*, 101–114
19. Khan, I., Crouch, J. D., Bharti, S. K., Sommers, J. A., Carney, S. M., Brosh, R. M., Jr. (2016) Biochemical characterization of the human mitochondrial replica-
tive Twinkle helicase: substrate specificity, DNA branch migration, and ability to overcome blockades to DNA unwinding. J. Biol. Chem. 291, 14324–14339

20. Bleichert, F., Botchan, M. R., and Berger, J. M. (2015) Crystal structure of the eukaryotic origin recognition complex. Nature 519, 321–326

21. Costa, A., Ilves, I., Tambeg, N., Petjoeyvic, T., Nogales, E., Botchan, M. R., and Berger, J. M. (2011) The structural basis for MCM2–7 helicase activation by GINS and Cdc45. Nat. Struct. Mol. Biol. 18, 471–477

22. Graham, B. W., Tao, Y., Dodge, K. L., Thaxton, C. T., Olaso, D., Young, N. L., Marshall, A. G., and Trakselis, M. A. (2016) DNA interactions probed by hydrogen-deuterium exchange (HDX) Fourier transform ion cyclotron resonance mass spectrometry confirm external binding sites on the minichromosomal maintenance (MCM) helicase. J. Biol. Chem. 291, 12467–12480

23. McKinney, S. A., Joo, C., and Ha, T. (2006) Analysis of single-molecule FRET trajectories using hidden Markov modeling. Biophys. J. 91, 1941–1951

24. Abeljon, J., Blanco, M., Ditzler, M. A., Fuller, F., Aravamudhan, P., Wood, M., Villa, T., Ryan, D. E., Pleiss, J. A., Maeder, C., Guthrie, C., and Walter, N. G. (2010) Conformational dynamics of single pre-mRNA molecules during in vitro splicing. Nat. Struct. Mol. Biol. 17, 504–512

25. Biswas, E. E., and Biswas, S. B. (1999) Mechanism of DnaB helicase of Escherichia coli: structural domains involved in ATP hydrolysis, DNA binding, and oligomerization. Biochemistry 38, 10919–10928

26. Biswas, E. E., and Biswas, S. B. (1999) Mechanism of DNA binding by the DnaB helicase of Escherichia coli: analysis of the roles of domain γ in DNA binding. Biochemistry 38, 10929–10939

27. Bailey, S., Elason, W. K., and Steitz, T. A. (2007) The crystal structure of the Thermus aquaticus DnaB helicase monomer. Nucleic Acids Res. 35, 4728–4736

28. Summerton, J., and Weller, D. (1997) Morpholino antisense oligomers: design, preparation, and properties. Antisense Nucleic Acid Drug Dev. 7, 187–195

29. Tackett, A. J., Wei, L., Cameron, C. E., and Raney, K. D. (2001) Unwinding of nucleic acids by HCV NS3 helicase is sensitive to the structure of the duplex. Nucleic Acids Res. 29, 565–572

30. Jejewska, M. J., Rajendran, S., and Bujalowski, W. (1998) Functional and structural heterogeneity of the DNA binding site of the Escherichia coli primary replicative helicase DnaB protein. J. Biol. Chem. 273, 9058–9069

31. Bujalowski, W., and Jejewska, M. J. (1995) Interactions of Escherichia coli primary replicative helicase DnaB protein with single-stranded DNA. The nucleic acid does not wrap around the protein hexamer. Biochemistry 34, 8513–8519

32. Jejewska, M. J., Rajendran, S., and Bujalowski, W. (1997) Strand specificity in the interactions of Escherichia coli primary replicative helicase DnaB protein with a replication fork. Biochemistry 36, 10320–10326

33. Ribeck, N., Kaplan, D. L., Bruck, I., and Saleh, O. A. (2010) DnaB helicase activity is modulated by DNA geometry and force. Biophys. J. 99, 2170–2179

34. Atkinson, J., Gupta, M. K., and McGlynn, P. (2011) Interaction of Rep and DnaB on DNA. Nucleic Acids Res. 39, 1351–1359

35. Guy, C. P., Atkinson, J., Gupta, M. K., Mahdi, A. A., Gwynn, E. J., Rudolph, C. J., Moon, P. B., van Krippenberg, I. C., Cadman, C. J., Dillingham, M. S., Lloyd, R. G., and McGlynn, P. (2009) Rep provides a second motor at the replisome to promote duplication of protein-bound DNA. Mol. Cell 36, 654–666

36. Kim, S., Dallmann, H. G., McHenry, C. S., and Marins, K. J. (1996) Coupling of a replicative polymerase and helicase: a tau-DnaB interaction mediates rapid replication fork movement. Cell 84, 643–650

37. Manosas, M., Spiering, M. D., Ding, F., Croquette, V., and Benkovic, S. J. (2012) Collaborative coupling between polymerase and helicase for leading-strand synthesis. Nucleic Acids Res. 40, 6187–6198

38. Indyani, C., Langston, L. D., Yurieva, O., Goodman, M. F., and O’Donnell, M. (2009) Translesion DNA polymerases remodel the replisome and alter the speed of the replicative helicase. Proc. Natl. Acad. Sci. U.S.A. 106, 6031–6038

39. Graham, J. E., Marins, K. J., and Kowalczykowski, S. C. (2017) Independent and stochastic action of DNA polymerases in the replisome. Cell 169, 1201–1213.e17

40. Sambrook, J., and Russel, D. W. (2001) Molecular Cloning: A Laboratory Manual, pp. 5.51–5.54, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY

41. Studier, F. W. (2005) Protein production by auto-induction in high density shaking cultures. Protein Expr. Purif. 41, 207–234

42. Bujalowski, W., Klonowska, M. M., and Jejewska, M. J. (1994) Oligomeric structure of Escherichia coli primary replicative helicase DnaB protein. J. Biol. Chem. 269, 31350–31358

43. Roy, R., Hohng, S., and Ha, T. (2008) A practical guide to single-molecule FRET. Nat. Methods 5, 507–516

44. Fagerburg, M. V., and Leuba, S. H. (2011) Optimal practices for surface-tethered single molecule total internal reflection fluorescence resonance energy transfer analysis, in DNA Nanotechnology, Methods and Protocols (Zuccheri, G., and Samori, B., eds) 1st Ed., pp. 273–289, Humana Press, New York, NY

45. Schauer, G. D., Huber, K. D., Leuba, S. H., and Sluis-Cremer, N. (2014) Mechanism of allosteric inhibition of HIV-1 reverse transcriptase revealed by single-molecule and ensemble fluorescence. Nucleic Acids Res. 42, 11687–11696

46. Rasband, W. S. (1997–2016) ImageJ, National Institutes of Health, Bethesda, MD

47. McGeoch, A. T., Trakselis, M. A., Laskey, R. A., and Bell, S. D. (2005) Organization of the archaeal MCM complex on DNA and implications for the helicase mechanism. Nat. Struct. Mol. Biol. 12, 756–762

48. Bronson, J. L., Fei, J., Hofman, J. C., Gonzalez, R. L., Jr, and Wiggins, C. H. (2009) Learning rates and states from biophysical time series: a Bayesian approach to model selection and single-molecule FRET data. Biophys. J. 97, 3196–3205

49. Arnold, K., Bordoli, L., Kopp, J., and Schwede, T. (2006) The SWISS-MODEL workspace: a web-based environment for protein structure homology modelling. Bioinformatics 22, 195–201