Molecular Traffic Jams on DNA

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Abstract

All aspects of DNA metabolism—including transcription, replication, and repair—involves motor enzymes that move along genomic DNA. These processes must all take place on chromosomes that are occupied by a large number of other proteins. However, very little is known regarding how nucleic acid motor proteins move along the crowded DNA substrates that are likely to exist in physiological settings. This review summarizes recent progress in understanding how DNA-binding motor proteins respond to the presence of other proteins that lie in their paths. We highlight recent single-molecule biophysical experiments aimed at addressing this question, with an emphasis placed on analyzing the single-molecule, ensemble biochemical, and in vivo data from a mechanistic perspective.
INTRODUCTION

Long stretches of naked DNA are unlikely to persist in vivo; rather, genomic DNA is coated with proteins that facilitate DNA compaction, organization, and maintenance. For example, in \textit{Escherichia coli}, the nucleoid-associated proteins H-NS, HU, Fis, IHF, and StpA are each present in copy numbers of 20,000 to 60,000 proteins per cell (∼50–150 μM) and are some of the most abundant cellular proteins (4). Furthermore, up to ∼30% of the \textit{E. coli} genome is occluded by DNA-binding proteins, suggesting that regions of protein-free DNA are unlikely to be longer than ∼100 base pairs (bp) (4, 61, 105). Similarly, eukaryotic genomic DNA is organized into chromatin, a dense mesh of proteins and nucleic acids. The basic unit of chromatin is composed of nucleosomes, each of which is made up of ∼147 bp of genomic DNA wrapped around two copies of the core histones proteins H2A, H2B, H3, and H4. Most of the eukaryotic genome is coated in nucleosomes and nucleosome-free DNA segments are, on average, only ∼20–30 bp in length (14).

Many essential aspects of DNA metabolism in both prokaryotes and eukaryotes—transcription, replication, and repair—must occur within these highly crowded environments. Despite the central role of nucleosomes and bacterial nucleoid-associated proteins in all aspects of DNA metabolism, a basic understanding of how DNA-binding motor proteins function in these crowded settings is still largely lacking. Furthermore, many of these processes must occur concurrently on the same stretch of DNA (Figure 1). In rapidly dividing prokaryotes, replication, transcription, and repair must all occur simultaneously. In eukaryotes, the replication machinery must also traverse highly transcribed sites (6) and will encounter megabase-length transcripts that require more than one cell cycle for complete synthesis (40). Thus, conflicts between transcription and replication must be resolved frequently in both prokaryotes and eukaryotes while somehow maintaining genome organization and stability.

How proteins are able to move along crowded nucleic acid tracks remains poorly understood and continues to be a topic of intense research. However, establishing a rigorous understanding of the outcomes of collisions on DNA is challenging because these events are transient, unsynchronized, and difficult to observe via traditional biochemical techniques. Methods such as single-molecule fluorescence imaging, single-molecule fluorescence resonance energy transfer (smFRET), optical and magnetic tweezers, and atomic force microscopy (AFM) offer the...
DNA replication, transcription, and repair must occur simultaneously on a crowded nucleic acid substrate. Conflicts between these molecular machines must be resolved rapidly to maintain cell viability and to avoid genomic instability.

Potential for observation and manipulation of biochemical reactions with unprecedented spatial and temporal resolution, thus capturing transient intermediates that are obstructed by traditional ensemble-averaging approaches.

In this review, we summarize recent developments in understanding how DNA-binding motor proteins function in crowded environments. Where possible, we focus on single-molecule studies that have revealed new insights into protein-protein collisions on DNA, and we integrate these results with findings that have arisen from ensemble biochemical and in vivo-based methods. Thus, a combination of complementary bulk biochemistry, single-molecule biophysics, and in vivo approaches is helping unravel the puzzle of how multiple DNA-metabolism activities occur simultaneously on a crowded DNA lattice.

**HELICASES**

Helicases are enzymes that hydrolyze nucleotide triphosphates (NTPs) to translocate along a nucleic acid track, and catalyze the unwinding of duplex RNA or DNA into individual single strands (87). Helicases participate in nearly all aspects of nucleic acid metabolism, including ribosome biogenesis, RNA folding, DNA replication, repair, and transcription (87, 90). This large group of enzymes has been organized into subfamilies based on structural, biochemical, and mechanistic classifications (93).

Early biochemical studies established that some helicases are capable of unwinding duplex DNA that is decorated with other DNA-binding proteins (46, 72). Helicases have even been found to disrupt a biotin-streptavidin interaction, which is one of the strongest noncovalent linkages found in nature (15, 72). In fact, many enzymes that contain canonical helicase motifs or display helicase activity in vitro may be primarily involved in the removal of other proteins from nucleic acids in vivo. Enzymes that are able to remove other proteins from a nucleic acid track are now frequently referred to as stripases (82). Helicases may also associate with larger macromolecular machines such as the replisome, where they act as stripases to help clear protein obstacles that would otherwise interfere with DNA replication (see Replication, below) (10, 67).

In this section we highlight recent advances in understanding how DNA helicases respond to protein obstacles in their path. We highlight single-molecule studies of three enzymes that show different behaviors when they are challenged with protein obstacles: (a) RecBCD, an *E. coli* helicase/nuclease involved in DNA double-strand break (DSB) repair, pushes and evicts roadblocks as it moves along double-stranded DNA (dsDNA) (28); (b) PcrA, a monomeric helicase from *Bacillus stearothermophilus*, translocates along single-stranded DNA (ssDNA) and disrupts RecA-ssDNA nucleoprotein filaments (76); and (c) XPD, a helicase involved in nucleotide excision repair, translocates along replication protein A (RPA)-bound ssDNA without displacing the protein obstacle (43).
RecBCD Removes Protein-Bound Obstacles

RecBCD is a heterotrimeric *E. coli* helicase/nuclease required for processing DNA DSBs, degrading invading phage DNA, and rescuing stalled replication forks (18). RecBCD is a fast (∼1,000–2,000 bp sec⁻¹) and processive (≥30,000 bp) enzyme that uses the energy of ATP hydrolysis to travel along DNA (7). RecB and RecD are members of the Superfamily 1 (SF1)-type helicases with 3'→5' (SF1a) and 5'→3' (SF1b) directionality, respectively (93). A central pin in the RecC unit bisects the DNA duplex and funnels the 3'- and 5'-ssDNA strands toward the RecB and RecD motors, respectively. As the ssDNA strands exit from their respective motor cavities, the ssDNA strands are degraded by an additional nucleolytic domain on RecB (18).

RecBCD must encounter many DNA-bound proteins in vivo, yet it was unclear how these potential obstacles might affect its progression (20). We have addressed the effects of protein roadblocks on RecBCD translocation using single-molecule fluorescence microscopy to visualize individual molecular collisions in real-time on DNA curtains (Figure 2). In the DNA curtain approach, individual DNA molecules are anchored to a fluid lipid bilayer and are aligned along the leading edges of nanofabricated chrome barriers to lipid diffusion through the application of hydrodynamic force (26, 27, 32) (Figure 2a). The DNA is stained with the highly fluorescent intercalating dye YOYO1 and can be visualized by total internal reflection fluorescence microscopy, enabling direct imaging of hundreds of individual molecules within a single field of view. Upon helicase unwinding, YOYO1 is displaced from the duplex DNA, which leads to a decrease in fluorescence (7, 35). Thus, the helicase/nuclease activity of RecBCD is directly observed as a decrease in length of the dsDNA as a function time (Figure 2b).

RecBCD was challenged with four different fluorescently labeled protein roadblocks, including *E. coli* RNA polymerase (RNAP) (28). When RecBCD collided with RNAP, the polymerase was pushed thousands of base pairs before it finally dissociated from the DNA (Figure 2b). Similar outcomes were observed for core, holoenzyme, stalled elongation complexes (ECs), and active ECs in both head-to-head and head-to-tail RecBCD-RNAP collisions. These results demonstrated that RecBCD exerts sufficient force to evict RNAP regardless of collision orientation or the transcriptional status of the polymerase, ensuring that the transcription machinery cannot impede DNA processing. Furthermore, RecBCD velocity did not decrease as it was pushing an RNAP holoenzyme (Figure 2e). These findings show that RecBCD is a more powerful stripase than even *E. coli* replisomes, which are readily obstructed by stalled RNAP (see below), and this conclusion is in agreement with the known genetic requirement for RecBCD in rescuing replication forks stalled at chromosomal roadblocks (16, 77).

We further tested RecBCD stripase activity by staging collisions with a hydrolytically defective restriction enzyme EcoRI(E111Q) (49, 106), Lac repressor (LacI), and *Saccharomyces cerevisiae* nucleosomes. Strikingly, RecBCD pushed and evicted all roadblocks with only a minor change in RecBCD velocity and processivity. Furthermore, RecBCD pushed and evicted roadblocks even when the helicase velocity was reduced 20-fold (by reducing the concentration of ATP), indicating the stripase activity remained functional under suboptimal translocation conditions (28)

A model in which the roadblock proteins dissociate from the DNA as they are forced to step from one nonspecific binding site to the next best described the experimental data (28). Figure 2d depicts four steps of RecBCD as it encounters and begins to push a protein obstacle. Protein dissociation takes place while the obstacle protein is in the weakly bound intermediate state, which predicts that the probability of roadblock dissociation is related to the cumulative number of steps that the roadblock is forced to take on the DNA, rather than the total amount of time that the roadblock spends on nonspecific DNA. In agreement with this model, we observed that the roadblocks were pushed similar distances regardless of how fast RecBCD moved (28).
Figure 2

(a) An illustration of the single-molecule DNA curtain assay used to observe RecBCD-roadblock collisions. Individual DNA molecules are tethered to a fluid lipid bilayer via a biotin-streptavidin interaction and organized at nanofabricated chrome barriers. RecBCD is loaded at the free DNA ends and fluorescent RNA polymerase (RNAP) is deposited on the native promoters along the DNA. The DNA is visualized by YOYO1, an intercalating fluorescent dye that does not interfere with RecBCD activity. The reaction is initiated by supplementing the flow buffer with ATP. As the DNA is degraded by RecBCD helicase/nuclease activity, the time-dependent decrease in DNA length serves as a readout of RecBCD translocation. (b) Kymographs of RecBCD pushing and evicting RNAP holoenzyme and elongation complexes (ECs) from DNA. In all kymographs, the tethered end of the DNA is at the top, the free end is at the bottom, and buffer flow is from top to bottom. RecBCD is able to efficiently push and eventually displace RNAP holoenzyme (top panel), and transcribing ECs (bottom panel). (c) A scatter plot of pre- and postcollision RecBCD velocities. The red line is a fit to the data with a slope of 1.0. RecBCD does not change velocity upon collision with RNAP. (d) A transition-state ejection model for roadblock displacement by RecBCD. Most roadblocks are displaced by RecBCD as they are pushed from one nonspecific DNA site to the next. Translocating RecBCD initially displaces the protein roadblock from a high-affinity specific site (S). The roadblock must pass through a much more weakly bound intermediate state (T) as it is pushed from one site to the next, followed by re-equilibration as a DNA-bound nonspecific complex (N). Subsequent steps by RecBCD continue to push the protein roadblocks from one nonspecific site to the next, until the proteins are eventually evicted from the DNA.

Importantly, our data do not rule out the possibility that EcoRI(E111Q), RNAP, and nucleosomes are structurally altered as they are pushed by RecBCD. However, even if the protein roadblocks were structurally perturbed, then this alone did not result in their eventual dissociation from DNA.

LacI differed from the other roadblocks in that, within our spatial resolution, it was immediately evicted from DNA, and the RecBCD-induced dissociation rate was comparable to the rate of spontaneous dissociation from nonspecific sites. However, we do not rule out short-range sliding by LacI, as has been observed in other single-molecule studies (21). Importantly, RNAP, EcoRI(E111Q), and nucleosomes all bind tightly to nonspecific DNA, whereas LacI binds much more weakly to nonspecific sequences, suggesting that LacI is released more rapidly from nonspecific DNA after a collision with RecBCD. Thus, the affinity of the roadblock protein for nonspecific DNA was a critical factor contributing to the outcome of the collisions. A remaining question is whether the mechanism identified for roadblock removal by RecBCD reflects a common feature of motor proteins that can disrupt stationary DNA-bound obstacles, or whether this...
Optical tweezers: technique that utilizes a tightly focused infrared laser to trap a bead, which can then be used to measure force-dependent responses of biological molecules coupled to the bead.

Behavior reflects some more specialized feature of RecBCD. Future experiments with different types of motor proteins will be critical for resolving this issue.

How much force does RecBCD exert on a protein obstacle and how much force is sufficient to disrupt these obstacles? An optical tweezers–based study measured the stall forces of a translocating RecBCD (78). The RecD subunit was immobilized on the surface of a microscope slide via a biotin-streptavidin interaction. One end of a dsDNA molecule was affixed to a polystyrene bead, and the free DNA end was engaged to the surface-tethered RecBCD. In the presence of ATP, the velocity of RecBCD as a function of opposing force could be measured by observing the displacement of the bead from the center of the optical trap. At a relatively low opposing force of ∼4–8 pN, RecBCD would slip back along the DNA and reengage forward movement if the opposing force was relaxed (78).

Although our experiments do not yield a direct readout of force, we can conclude that the force exerted by RecBCD is sufficient to overcome RNAP, which can withstand substantially higher, ∼30 pN stall forces (41). The observation that RecBCD can strip RNAP (and other proteins) from DNA would seem inconsistent with the low stall force reported for RecBCD based on optical trap measurements. However, an experimentally applied stall force is not necessarily equivalent to the force that a motor protein can apply against an oncoming obstacle. One possible explanation for this discrepancy may be that stall-force measurements place tension along an arbitrary vector of this multisubunit enzyme. This may lead to a partial mechanical unfolding of the protein, or disruption of critical intersubunit interactions and concomitant loss of translocase activity. These considerations highlight the need for new single-molecule force manipulation protocols for measuring pushing forces (as opposed to stall forces) of motor proteins.

PcrA Clears RecA Recombinase from ssDNA

Helicases can translocate on ssDNA and play crucial roles in removing unwanted ssDNA-bound proteins (17, 38, 88). For example, UvrD, PcrA, and Srs2 can promote the removal of the recombinases RecA or Rad51 from transiently exposed ssDNA regions to prevent inappropriate homologous DNA recombination (52, 76).

PcrA is an SF1 helicase that translocates along ssDNA in a 3′ → 5′ direction with a step size of 1 nucleotide (nt) (76). Monomeric PcrA preferentially binds to the ssDNA–dsDNA junction and is able to repetitively reel in the 5′-ssDNA tail. However, a dimeric or higher order form of the enzyme is required for helicase activity in vitro. The oligomeric state and in vivo function of PcrA and its E. coli homolog UvrD continue to be debated (17). Both helicases have been implicated in nucleotide excision repair, plasmid replication, and roadblock clearance ahead of the replisome, and as antirecombinases that act at stalled or collapsed replication forks (17). A smFRET approach was used to characterize B. stearothermophilus PcrA and to demonstrate that the enzyme could dislodge RecA nucleoprotein filaments from ssDNA.

Park et al. (76) used smFRET to directly observe the activity of PcrA on RecA-ssDNA filaments. A DNA oligonucleotide with a long ssDNA tail was immobilized on the surface of a passivated microscope slide via a biotin-streptavidin interaction (Figure 3a). One end of the oligonucleotide was decorated with a Cy3 dye and the other end with a Cy5 dye. Upon photoexcitation of Cy3, nonradiative energy transfer may occur to Cy5 if the dyes are within the FRET transfer distance (∼30–70 Å). The smFRET signal is observed as a photon emitted in the Cy5 fluorescence region. In the absence of all proteins, the ssDNA tail behaves as a highly flexible polymer, bringing the Cy3-Cy5 ends sufficiently close to ensure a strong smFRET signal (Figure 3b). Upon addition of RecA and ATP, a rigid nucleoprotein filament is formed, causing the smFRET signal to decrease (38, 48). When added to the flowcell, PcrA binds the ssDNA–dsDNA junction and
Figure 3

(a) A smFRET-based assay for observing the RecA-clearing activity of PcrA helicase. A DNA oligonucleotide with a Cy3-Cy5 FRET dye pair is affixed to the surface of a passivated microscope slide. Entropically driven collapse of the ssDNA brings the two dyes within efficient FRET distance. Addition of RecA leads to a rigid nucleoprotein filament, which separates the two dyes and reduces the observed FRET signal. In the presence of ATP, PcrA helicase clears RecA and reels the free DNA ends closer together. (b) smFRET data. (Top) Naked ssDNA brings both dyes relatively close, leading to a FRET signal of ∼0.5. (Middle) Addition of RecA elongates the ssDNA tail and reduces the smFRET to ∼0.3. (Bottom) In the presence of ATP, PcrA clears RecA from ssDNA as it reels the two DNA ends close to each other.

displaces RecA from ssDNA in an ATP-dependent reaction. Multiple cycles of PcrA shuttling along the same RecA-coated ssDNA strand are observed experimentally (76) (Figure 3b). The strong preference of monomeric PcrA for ssDNA-dsDNA junctions suggests that in vivo PcrA may preferentially bind at sites of stalled or collapsed replication forks and translocate along the newly synthesized lagging strand to remove recombinase proteins that might be loaded along the ssDNA. Thus, a major function of PcrA may be to act as an antirecombinase by keeping ssDNA free of RecA and thereby preventing deleterious recombination events. Importantly, although single-molecule and ensemble biochemistry studies show that PcrA and UvrD can act as antirecombinases, exactly how RecA is removed from DNA has not been delineated (60, 102). Future single-molecule studies may address critical questions such as the mechanism by which RecA is cleared from DNA, and whether individual RecA subunits are removed sequentially or as larger oligomers.

Eukaryotic UvrD/PcrA homologs have also been implicated in roadblock disruption and antirecombinase activity (64, 71, 103). The S. cerevisiae Srs2 helicase is an SF1 DNA-dependent ATPase that translocates along ssDNA to remove illegitimate Rad51-ssDNA filaments (64, 103). Intriguingly, ATP hydrolysis by Rad51 is required for efficient removal by Srs2, suggesting that Srs2 provokes displacement of Rad51 from ssDNA by taking advantage of the lower affinity of Rad51 for DNA while in the ADP-bound state (2). Furthermore, Srs2 physically interacts with the replisome, suggesting that it functions to prevent Rad51 accumulation on ssDNA patches that are transiently generated during DNA replication (3, 79). Does monomeric Srs2 prefer ssDNA-dsDNA junctions and does it repetitively shuttle along ssDNA to displace Rad51? Is Rad51 displaced as monomeric units or as oligomers, and how does its ATP hydrolysis affect Srs2 activity? Single-molecule techniques are poised to add crucial new mechanistic details to this intriguing family of proteins.
Figure 4

A single-molecule dye-quenching assay for XPD translocation along replication protein A (RPA)-coated DNA. (a) A DNA oligonucleotide is immobilized on the surface of a passivated flowcell and decorated with Cy5-labeled RPA. The FeS cluster of XPD helicase quenches Cy5 fluorescence in a distance-dependent manner. (b) When ATP is added, XPD translocation toward RPA initially quenches the Cy5 fluorescence (arrow 1), and eventually XPD bypasses the RPA (arrow 2), which is revealed as an eventual recovery of the Cy5 intensity. (c) Examples of intensity trajectories showing XPD approaching (arrow 1) and moving past (arrow 2) RPA.

XPD Can Bypass RPA on the Same ssDNA

Not all encounters between a helicase and a DNA-bound protein must end in roadblock eviction. For example, Spies and colleagues (43) used a novel single-molecule fluorescence quenching approach to demonstrate that Ferroplasma acidarmanus XPD helicase is able to translocate on protein-coated ssDNA without disrupting some of the bound proteins. XPD is an SF2 helicase involved in nucleotide excision repair and translocates along ssDNA with a 5′ → 3′ polarity (43, 84). XPD encodes an FeS cluster domain that is important for coupling ATP hydrolysis to translocation and also forms a secondary ssDNA-binding site (84, 86). To monitor XPD movement along DNA, the authors cleverly exploited the observation that the FeS cluster of XPD quenches nearby Cy3 and Cy5 dyes in a distance-dependent fashion. A schematic of the experimental setup is illustrated in Figure 4a (43). A Cy3-labeled ssDNA oligonucleotide was immobilized on the surface of a passivated microscope slide via biotin-streptavidin interactions. In the absence of XPD, the Cy3-fluorescence intensity remained constant. When wild-type XPD and ATP were added to the flowcell, XPD began translocating along the ssDNA, and the Cy3 signal was quenched by the approaching FeS cluster (Figure 4b). Thus, XPD velocity could be characterized on naked DNA by using Cy3 quenching as a proxy for translocation (43).

The authors went on to characterize the translocation of XPD on protein-coated ssDNA. *F. acidarmanus* encodes two distinct proteins that bind ssDNA: RPA1 occludes 20 nt of ssDNA and contains two OB folds and a Zn-finger domain; RPA2 contains a single OB fold and occludes 5 nt of ssDNA per monomer of RPA2. Both proteins bind ssDNA with nanomolar affinity. XPD
translocated on RPA1-coated ssDNA at the same velocities as seen for naked ssDNA, indicating that RPA1 did not interfere with XPD translocation. Furthermore, RPA1-ssDNA binding lifetimes were significantly reduced when XPD was introduced into the flowcell. This result suggested that XPD competed with RPA1 for ssDNA and cleared RPA1 from the ssDNA during translocation.

The outcome was remarkably different in reactions with RPA2. In the presence of RPA2, XPD translocation velocities displayed a bimodal distribution with two distinct rates: a rate of 13 nt s\(^{-1}\), as was also observed on naked ssDNA, and a second rate of just 6 nt s\(^{-1}\) (43). Strikingly, RPA2 lifetimes on ssDNA were not affected by XPD translocation, suggesting that RPA2 was not displaced even as XPD was translocating along the same ssDNA. The authors then used a smFRET assay with Cy3-ssDNA and Cy5-RPA2 to verify that XPD could motor along an RPA2-coated ssDNA lattice without displacing RPA2 from the DNA. This is the first direct observation of a protein translocating along crowded DNA without displacing other bound proteins. This result is in agreement with other studies suggesting that XPD helicase activity is stimulated by RPA2 but not by RPA1 (85). The authors suggest that some helicases may specifically interact with specialized ssDNA-binding proteins to promote their biochemical activity at the proper location. Further structural information on XPD will be required to understand how the helicase is able to achieve this feat, and additional single-molecule and ensemble biochemistry work will clarify whether other types of motor proteins can also function on nucleic acid substrates without displacing proteins that might lie in their paths.

**REPLICATION**

The replication machinery must duplicate genomic DNA that is decorated with nucleosomes, transcription factors, RNAPs, and other nucleoprotein complexes (70). Replication fork collapse at a nucleoprotein barrier may lead to compromised replication and genomic instability (19, 40). Conflicts between replication and transcription are particularly deleterious, as they must occur frequently (67, 70). Below, we highlight recent ensemble biochemical studies that have helped define the strategies that organisms use to avoid replisome collapse at protein roadblocks.

**Nucleosomes**

In vitro biochemical studies established that replisomes are strongly inhibited by protein roadblocks, and these studies were among the first to highlight the conceptual importance of protein-protein collisions on DNA (12, 81). In the first example of this work, Alberts and colleagues (12) demonstrated that a bacteriophage T4 replisome requires the addition of Dda, a helicase that is not required for replication of naked DNA, to replicate past a nucleosome. The authors concluded that Dda acts as a roadblock-clearing enzyme in front of the T4 replisome (12). Although the role of Dda in T4 phage replication has been reexamined (53, 62), this original work spurred numerous other studies that have now highlighted the roadblock clearance role of accessory nonreplicative helicases in both prokaryotes and eukaryotes (see Accessory Helicases, below) (65).

Progress on understanding how the eukaryotic replication machinery proceeds through a nucleosome has been hampered by a paucity of reconstituted eukaryotic replisomes. SV40 replication continues to be the only fully reconstituted eukaryotic replisome that can be carried out in both cell-free extracts and in a highly purified in vitro system. SV40 replisomes efficiently replicated chromatinized templates (45), and upon passage of a replication fork, nucleosomes were repositioned behind the replisome with an equal probability of being placed on either the leading or lagging strand (97). In addition, the large T antigen, which is the SV40 replicative helicase, is capable
of unwinding nucleosome-containing DNA, suggesting the replicative helicase may contribute to nucleosome displacement from DNA during replication (89). However, whether the nucleosome core particle is actually disrupted during replication remains controversial. Several studies concluded that H2A/H2B is lost upon histone transfer behind the replisome (30, 33), although chemically cross-linked nucleosomes are also efficiently transferred behind the SV40 replisome (104).

The polycomb group of epigenetic DNA-binding proteins was shown to transfer efficiently behind an SV40 replisome during replication in vitro, suggesting that this group of proteins, which makes critical contributions to epigenetic inheritance, has evolved to survive collisions with replisomes without being displaced from DNA (29). More work needs to be carried out to understand whether and how nucleosomes and other epigenetic factors remain bound to chromosomes during eukaryotic replication, and single-molecule observations may play a key role in understanding their fates when faced with an oncoming replisome.

Accessory Helicases

Nonreplicative accessory helicases have been identified as critical roadblock-clearing motors that participate in DNA replication in T4 phage, E. coli, Bacillus subtilis, and S. cerevisiae (12, 36, 67). In an elegant biochemical and cell biology study, McGlynn and colleagues (36) identified how two nonreplicative helicases help E. coli replisomes get past protein roadblocks. For this work, EcoRI(E111G), a hydrolytically defective restriction enzyme that binds its cognate site with high affinity, was employed as a DNA-binding protein roadblock (49). Eight EcoRI(E111G) repeats were found to sufficiently block progression of an in vitro–reconstituted E. coli replisome (66). The authors then tested eight nonreplicative helicases for their ability to stimulate replication through the EcoRI(E111G) nucleoprotein barrier. Two E. coli–encoded helicases, Rep and UvrD, which are 3′→5′ SF1 helicases that, in the absence of the replisome, cannot dissociate EcoRI(E111G) from DNA, greatly stimulated replication past the EcoRI(E111G) barrier in vitro. In addition, PcrA, a homologous SF1 helicase from B. subtilis, promoted replication past the barrier. Importantly, although Dda is capable of clearing multiple proteins from DNA and can even disrupt the very strong biotin-streptavidin interaction, it did not stimulate replication past EcoRI(E111G) (15).

In vivo analysis of rep uvrD cells confirmed the in vitro results. For example, both Rep and UvrD were required for growth on rich media (i.e., conditions under which conflicts between replication and transcription are expected to occur most frequently), and although rep uvrD colonies could be recovered at very low plating dilutions on rich media, these colonies were found to harbor suppressor mutations that reduce RNAP backtracking (75). Backtracked RNAP is likely the most dangerous natural replication barrier and a source of genomic instability (see below); thus, survival of the rep uvrD mutants was contingent upon elimination of a naturally occurring protein roadblock to DNA replication through the acquisition of suppressor mutations (19, 75).

Accessory helicases such as Rep, UvrR, and PcrA are capable of disrupting different types of protein roadblocks ahead of the replisome. For example, rep uvrD cells could not tolerate 34 chromosomal tandem LacI-operator repeats, although this same nucleoprotein barrier was well tolerated in wild-type cells. Expression of PcrA, but not Dda, in a rep uvrD background rescued the double-knockout phenotype.

Either Rep or UvrD may dislodge protein roadblocks to release stalled replication forks, and a model that describes the role of both Rep and UvrD at the replication fork is summarized in Figure 5. The C terminus of Rep interacts directly with DnaB, the replicative helicase, suggesting Rep is an integral component of the E. coli replisome. If the replisome stalls behind a nucleoprotein obstacle, Rep is proposed to load onto the strand opposite of DnaB where it can then displace the roadblock. UvrD does not physically interact with the replisome, but it is present at a very
Figure 5

(a) Accessory helicases interact with the replisome to remove protein roadblocks. In *Escherichia coli*, the C terminus of the accessory, nonreplicative Rep helicase physically interacts with DnaB. Accessory helicases can aid replication past protein roadblocks such as LacI repeats, hydrolytically inactive EcoRI, and head-on RNAP collisions. (b) Upon encountering a backtracked RNA polymerase (RNAP) in a codirectional orientation, the replisome can reinitiate from the 3′ end of the mRNA R-loop. A subsequent round of replication converts the resulting nick into a double-strand break. Abbreviation: EC, elongation complex.

There is growing support for the hypothesis that eukaryotes also employ nonreplicative helicases for roadblock clearance (10). The Rrm3/Pif1 family of helicases is involved in roadblock clearance at the *S. cerevisiae* replisome (10). Analogous to Rep, Rrm3 physically interacts with...
DNA polymerase epsilon (ε) and PCNA (92), making it an integral component of the yeast replisome. Rrm3 translocates in a 5′→3′ direction, which is opposite the 3′→5′ polarity of the MCM replicative helicase. Thus, a combination of genetic and biochemical data strongly underscores the need for accessory helicases in replisome progression past protein roadblocks.

How accessory helicases are regulated at the fork and how they physically remove proteins from DNA remain uncertain. Single-molecule biophysical studies are ideally suited for parsing out the mechanistic requirements of accessory motors in replisome progression. For example, single-molecule fluorescence methods can observe whether accessory helicases are a stable part of the replisome or are transiently recruited to stalled forks. Furthermore, it will be important to address the actual mechanism of roadblock clearance, such as determining whether multiple helicases are required to clear tough roadblocks, as has been observed for Dda on ssDNA (15).

**Replication-Transcription Conflicts**

RNAP is one of the most pervasive and potent roadblocks to DNA replication (6, 22, 23, 40, 68, 80, 81). In eukaryotes, the transcription of many genes, including the core histones, is highly upregulated during S-phase (34). Furthermore, transcription of the longest genes present in higher eukaryotes takes more than one full cell cycle; therefore, conflicts between RNAP and the replication machinery must occur at these genes (40). In prokaryotes, transcription is 20-fold slower than replication, guaranteeing that replisomes will frequently collide with the transcribing RNAPs. Several recent reviews have summarized replication-transcription conflicts and how they may be resolved (65, 67, 69).

Replisomes can collide with RNAP in either head-on or codirectional orientations. Head-on collisions are thought to arrest replisome progression (67, 80, 81), and most bacteria minimize head-on collisions by organizing highly transcribed genes codirectionally with DNA replication (37, 94). In support of this view, Pomerantz & O’Donnell (81) demonstrated that in vitro, head-on collisions between an *E. coli* replisome and RNAP caused fork stalling and required the Mfd ATPase to assist in dislodging RNAP. In contrast, codirectional collisions were resolved when RNAP was displaced by the replisome itself, and replication could then resume by using the 3′-OH of the nascent transcript as primer for continued synthesis (80).

A recent in vivo study by Nudler and colleagues (19) demonstrated that codirectional, but not head-on, collisions lead to DNA DSBs and genome instability in *E. coli*. For these assays, a unidirectional origin of replication was used to stage head-to-head or codirectional collisions on plasmid substrates. Transcription was under the control of a temperature-sensitive lambda repressor and the resulting products were identified using primer extension PCR assays. The transcript also included a ribosome-binding site, and the translation state was modulated by the inclusion or omission of an ATG start codon. Replication could also be suppressed by supplementing the media with sublethal levels of hydroxyurea (HU) (19). Interestingly, the authors observed that backtracked RNAP resulted in replication-dependent DSBs. *E. coli* employs multiple mechanisms to prevent RNAP backtracking, including GreA/GreB-dependent transcription restart, transcription-translation coupling, and ATPase-dependent RNAP clearance (67, 75), and DSB levels were significantly increased in greA and greB cells. Conversely, when the ATG start codon was introduced to couple transcription to translation, the occurrence of DSBs was reduced to near-background levels, even in a greA/greB mutant background. Thus, the ribosome plays a dominant role in suppressing the negative impact of backtracked RNAP, thereby promoting successful DNA replication (83).

The same study found that DSBs were almost completely absent in head-on collisions, despite the fact that head-on collisions have long been considered the most deleterious to DNA
Replication forks. The authors explain this puzzling observation by suggesting that displacement of a backtracked RNAP during a codirectional collision may lead to a stable R-loop that can then be used by the replisome to reprime downstream DNA synthesis (Figure 5b). The RNA-DNA hybrid must then be cleared by RNase H and the resulting ssDNA resynthesized before the next round of replication (1). However, a DSB may form if the next round of replication reaches the R-loop before this repair is completed. In support of this model, overexpression of RNase H alleviated DSBs in codirectional collisions. Even though head-on collisions can stall the replisome, they do so without leading to DNA damage (19). Thus, codirectional conflicts may be particularly dangerous sources of genomic instability and must be minimized or resolved in vivo.

Reconstructing replication-transcription conflicts at the single-molecule level will be necessary to address many of the remaining mechanistic questions. For example, is the backtracked RNAP evicted immediately from the DNA or is it pushed ahead of the replisome, and how does the replisome resolve multiple collisions at heavily transcribed genes, such as the rDNA operon? The development of single-molecule replication assays promises to help put these experiments within our technical capabilities (31, 99, 101).

TRANSCRIPTION

Gene expression is intimately related to chromatin organization. RNAP must gain access to genomic DNA that is organized in nucleosomes, and positioned nucleosomes regulate the transcription program on a whole-genome level. Thus, it is crucial to understand the mechanisms of transcription through nucleosomes.

Pioneering studies by Studitsky et al. (95) established that phage SP6 RNAP transcribed efficiently through a nucleosome, revealing that the nucleosome could be transferred in cis onto the DNA behind the moving polymerase. In the DNA-looping model, the RNAP EC bends DNA sufficiently to facilitate formation of new nucleosome-DNA contacts behind the EC. As the EC peels DNA away from the surface of the nucleosome, the newly exposed histone contacts are transferred to the naked DNA present behind the progressing EC (95).

Reconstitution of in vitro transcription by S. cerevisiae RNA polymerase III (pol III) confirmed that the DNA-looped intermediate mechanism was also valid for eukaryotic RNAPs (96). Pol III, an enzyme that transcribes relatively short rRNA and tRNA genes, may be unlikely to transcribe past large numbers of nucleosomes in vivo. In vitro, however, pol III transcribes through a nucleosome in a manner analogous to that of phage RNAPs (96).

Does RNAP Actively Disrupt Nucleosomes?

In eukaryotes, RNA polymerase II (pol II) transcribes mRNA and may need to frequently contend with nucleosome barriers. However, in vitro, nucleosomes were a potent barrier to both yeast and human pol II transcription (11, 50, 51, 55). Furthermore, the mechanism of transcription and the fate of the nucleosomes differed markedly compared to what was seen with the simpler phage RNAPs (50, 55).

Bustamante and colleagues (8, 42) employed a high-resolution single-molecule optical tweezers assay to study the mechanism of how pol II traverses the nucleosome barrier. The study used a dual-optical trap approach as illustrated in Figure 6. One polystyrene bead was affixed to the end of a DNA molecule via a digoxigenin-antidigoxigenin interaction. A pol II EC was assembled and affixed to a second streptavidin-coated bead. A single nucleosome was assembled on a strongly positioning Widom 601 DNA sequence downstream of the RNAP promoter (Figure 6a). The
Figure 6
An optical trap assay to investigate RNA polymerase II (RNAP II) transcription past a nucleosome. (a) A schematic of the DNA construct used in this study. One end of the DNA was affixed to a polystyrene bead via a DIG–anti-DIG interaction. Biotinylated pol II was assembled onto a promoter (blue) and affixed to a second streptavidin-coated bead. A nucleosome was reconstituted on the strong nucleosome-positioning Widom 601 DNA sequence (brown). (b) An illustration of the experimental setup. RNAP transcription was monitored as a change in the DNA length between the two optical traps. (c) Individual pol II transcription traces as a function of KCl concentration. At low salts, pol II stalls at the nucleosome barrier. As KCl concentration is increased, pol II transiently pauses and eventually transcribes past the nucleosome. (d) A histogram of nucleosome-dependent arrest probability. The polymerase pauses as it encounters the nucleosome but can transcribe past transiently disrupted histone-DNA interactions at higher salt concentrations. (c,d) Transcription traces on naked DNA are shown in black; transcription traces on DNA with nucleosomes are shown in colors. The shaded regions of the graphs represent the location of the nucleosome positioning sequence. Abbreviations: EC, elongation complex; DIG, digoxigenin.
reactions were then initiated by the addition of all four rNTPs, and transcription by pol II was revealed as a change in the DNA length between the two beads.

As expected, transcription on naked DNA is subject to pausing and backsliding of the RNAP EC (59, 75). However, the pol II pause propensity and pause durations increased markedly as the polymerase approached the nucleosome, and pol II could not transcribe past the nucleosome at low KCl conditions (42) (Figure 6c). At higher KCl concentrations, where the nucleosome-DNA electrostatic interactions are significantly weakened, transcription could proceed past the nucleosome, albeit with reduced velocities (42) (Figure 6).

The data is consistent with a model in which pol II acted as a ratchet that invades a spontaneously unwrapped nucleosome, but is inconsistent with models invoking active pol II–dependent disruption of the histone-DNA contacts (42). Polymerase pausing was significantly increased as the EC approached the dyad axis of the nucleosome; high-resolution single-molecule nucleosome unwrapping experiments have indicated that this region has particularly strong electrostatic histone-DNA interactions and that these interactions are disrupted at higher ionic strengths (39). Upon reaching the nucleosome dyad axis, pol II either backtracks (see below) or continues transcription through the rest of the nucleosome. Importantly, the data ruled out a model in which pol II used the energy of NTP hydrolysis to actively unwind the nucleosome; rather it seems as though pol II waits for the nucleosome to spontaneously unravel (42).

The nucleosome barrier to transcription was further analyzed in terms of individual contributions from the histone-DNA interactions, the underlying DNA sequence, and the histone tails (8). Bustamante and colleagues demonstrated that each of these components of the nucleosome barrier has a profound and distinct effect on the rate of transcription through a nucleosome. Removing the histone tails stimulated transcription past a nucleosome by increasing the rate of spontaneous histone-DNA wrapping-unwrapping. In contrast, the underlying DNA sequence may cause pol II pausing, and these pauses are amplified at a nucleosome barrier. The nascent RNA transcript was found to prevent pol II backtracking, and removing the transcript with RNase A dramatically reduced transcription through a nucleosome (8).

The strength of the nucleosome barrier to transcription is strongly affected by histone-DNA interactions. This viewpoint is supported by the finding that a single point mutation in the histone H3 could rescue yeast cells that were missing an otherwise essential chromatin remodeling factor (SWI/SNF) (54). This mutation, known as sin (Swi-INdependent), disrupts a strong nucleosome-DNA interaction, thus allowing more ready displacement of nucleosomes even in the absence of the SWI/SNF chromatin remodeler (54, 73). Importantly, transcription by human pol II was significantly more efficient through sin nucleosomes than wild-type nucleosomes, and was even stimulated to near naked-DNA levels by the addition of the transcription elongation factors TFIIS and TFIIF (44, 63). When examined at the single-molecule level, sin mutations in H3 and H4 stimulated transcription through a nucleosome by decreasing the spontaneous histone-DNA rewrapping rate (8). Thus, the strength of the histone-DNA interactions plays a key role in modulating transcription in vitro and in vivo.

The fate of the nucleosome has also been investigated through the use of single-molecule AFM imaging. The rate of transcription may dictate the fate of the nucleosomes (9). Rapidly transcribing RNAP facilitates nucleosome dissociation by not leaving sufficient time for the unwrapped nucleosome to transfer behind the RNAP. For slower transcribing pol II, H2A/H2B disruption may result in a hexasome that is deposited behind the EC (9). These observations may explain the differences in histone loss seen between rapidly transcribing phage RNAPs and pol II (95), and are in agreement with in vivo evidence for rapid H2A/H2B exchange at moderately transcribed genes (see below) (56, 58).
A Trailing RNAP Assists in Overcoming Nucleosome Barriers

Backtracking by the RNAP EC is a universal feature of all RNAPs and may interfere with transcription past protein roadblocks and result in genome instability (19, 75). In prokaryotes, four major mechanisms can rescue a backtracked RNAP EC: (a) Translation of the nascent transcript prevents RNAP backtracking (83); (b) factors such as greA/greB rescue stalled ECs by clipping the backtracked message to realign the 3′-OH mRNA end with the RNAP catalytic site (74); (c) ATP-ases such as Rho and Mfd can clear stalled RNAP from DNA (13, 25, 91); and (d) cotranscription by two polymerases decreases backtracking by the leading EC (24, 47).

Cotranscription by two closely associated T7 RNAPs can also increase transcription through a nucleosome (47). An optical tweezers–based DNA unzipping assay was used to investigate how two T7 RNAPs collaborate to transcribe past a nucleosome barrier (Figure 7). By unzipping duplex DNA, the authors directly mapped the position of the RNAP active site. Gel electrophoresis of the RNA message was used to determine the length of the transcript with single-nucleotide sensitivity. A comparison of the RNAP position to the length of the mRNA transcript specified how far the RNAP backtracked upon encountering a nucleosome barrier.

The positions of individual backtracked T7 RNAP after a 10-s round of transcription through a nucleosome are shown in Figure 7a. Gel electrophoresis of the mRNA indicated that transcription stalled at the −60 bp position relative to the nucleosome dyad. However, the RNAP backtracked an additional −15 bp away from 3′-mRNA end upon encountering a nucleosome. Furthermore, run-off transcription through a nucleosome was significantly compromised, even after a 30-minute transcription reaction (47).

A second T7 RNAP EC significantly stimulated transcription of the leading RNAP through a nucleosome. The DNA construct and the location of the leading RNAP active site for this construct are shown in Figure 7b. The leading RNAP active site shifted by +15 bp, relative to the single T7 construct shown in Figure 7a. Furthermore, a trailing RNAP could stimulate transcription through the nucleosome fivefold and reduced pausing and backtracking of the leading RNAP. On the basis of these findings, the authors proposed a model whereby the trailing RNAP provides an assisting force that rescues the backtracked leading RNAP; the combined action of two concurrently acting motors promotes transcription through the nucleosome (47).

The results with two codirectional T7 RNAPs were in strong agreement with biochemical studies of both E. coli and yeast RNAP transcription through a nucleosome (24, 57). A trailing pol II significantly stimulated nucleosome read-through by the leading EC and had a strong influence on the fate of the nucleosome (57). When the RNAPs were close to each other, the nucleosome was efficiently transferred behind the trailing EC. When the polymerases were staged farther apart, the nucleosome was lost. The authors proposed a model whereby if two RNAPs are close, the nucleosome can physically interact with the trailing pol II and transfer behind both ECs. However, if there is space on the DNA between the two ECs, then the first RNAP deposits...
a Single-promoter template

Single molecule

b Two-promoter template

Single-molecule leading RNAP

f = 10 s
a hexasome behind itself, and this hexasome is then completely displaced from the DNA upon encountering the trailing polymerase.

Taken together, these observations are yielding a more general picture of how RNAPs can transcribe through nucleosomal barriers. Rapidly transcribing phage RNAPs and yeast pol III are able to elongate past a nucleosome efficiently, and transcription is modulated largely by transient nucleosome unwrapping. In contrast, the slower *E. coli* RNAP and eukaryotic pol II pause and backtrack at a nucleosome barrier, while antibacktracking mechanisms stimulate pol II progression past the nucleosome.

The situation is more complicated in living cells, and many additional chromatin remodeling factors, such as Asf1 and FACT, participate in transcription regulation on chromatinized templates (98, 100). In vitro reconstitutions show that these enzymes associate with RNAP II and aid in transcription past nucleosomes. Future single-molecule experiments may help determine the role of these and other accessory factors in regulating transcription through nucleosomes.

**FINAL THOUGHTS**

All DNA-binding translocases must contend with chromosomes that are not just naked DNA, but rather highly complex and dynamic nucleoprotein structures. Collisions between motor proteins on DNA with other DNA-bound roadblocks can lead to deleterious events such as replication collapse, transcription deregulation, and genomic instability. Detailed mechanistic pictures of how different motor proteins function on crowded substrates will provide crucial insights necessary for understanding DNA replication, transcription, genome maintenance, and epigenetic inheritance. Why do some motor proteins, such as RecBCD, remove obstacles, whereas others seem unable to move past barriers? Do motor proteins use common mechanisms for obstacle displacement, or will each have unique and specific attributes related to their function and regulation? Is the ability to exert force a defining factor in dictating the outcomes of collisions on DNA, and to what extent do motor proteins vary in terms of the amount of force they can exert on an obstacle? What are the roles of translocases as accessory enzymes in larger macromolecular machines? How do different motors contend with nucleosomes without disrupting crucial genomic organization? Many recent breakthroughs in single-molecule imaging and manipulation techniques are allowing the field to tackle increasingly more challenging biochemical systems, which will begin to address these complex questions. Future studies will lead to a more detailed molecular understanding of how these remarkable enzymes can translocate through the dense mesh of obstacles in their path.

**SUMMARY POINTS**

1. The DNA within living cells is bound by various proteins, such as nucleoid-associated proteins in bacteria and nucleosomes in eukaryotes. DNA motor enzymes, including helicases and polymerases, must be capable of acting in nucleic acid substrates bound by these potential obstacles.

2. Traditional experimental approaches can often discern whether a particular motor protein is capable of traversing DNA bound by stationary obstacles, but it is very difficult to obtain detailed mechanistic information because these events are transient, unsynchronized, and difficult to detect. Single-molecule methods offer the opportunity to probe these collisions in real time with high spatial and temporal resolution, thus capturing transient intermediates that are otherwise undetectable.
3. Single-molecule imaging studies have revealed RecBCD as a powerful DNA translocase that disrupts a range of protein–nucleic acid complexes and acts through a mechanism in which the obstacle proteins are evicted as they are pushed from one nonspecific site to the next. The generality of this mechanism for other DNA-binding motor proteins remains unexplored.

4. Replication forks are subject to stalling upon collisions with RNAP in either head-on or codirectional collisions. Bacterial genomes have evolved to help avoid head-on collisions, which are thought to be the most deleterious, and specialized accessory helicases also assist obstacle bypass during DNA replication.

5. Nucleosomes are likely the most common obstacle encountered in eukaryotic cells.

6. An important challenge in the field is to push toward even more complex biochemical systems (such as fully recapitulated replication forks or eukaryotic RNAPs acting in the presence of a full complement of chromatin remodeling factors) using experimental conditions that mimic highly crowded in vivo environments as closely as possible.

**DISCLOSURE STATEMENT**

The authors are not aware of any affiliations, memberships, funding, or financial holdings that might be perceived as affecting the objectivity of this review.

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**LITERATURE CITED**


36. Helped reveal how the secondary helicases Rep and UvrD act at replication forks to promote roadblock bypass in living cells.

43. Provided the first suggestion that a translocase could move along a nucleic acid substrate while bypassing, but not displacing, other proteins.

52. Showed that the Srs2 helicase from *S. cerevisiae* acts to remove the recombinase protein Rad51 from ssDNA molecules.
mRNA as a primer.

synthesis by using the 3′ replisome resuming are resolved with the replisomes and RNAP between the codirectional collisions 80. Showed that ssDNA loops.

ssDNA while generating translocating along the junctions and ssDNA/dsDNA anchoring to the RecA filaments by reeling in DNA in uniform steps.

repeatedly dismantle molecules of PcrA could 76. Showed that single molecules of PcrA helicase dismantles RecA filaments by reeling in DNA in uniform steps. Cell 142:544–55


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