Transcription elongation
Heterogeneous tracking of RNA polymerase and its biological implications

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Abbreviations: bp, base pair; ChIP-chip, chromatin immunoprecipitation followed by microarray; CPD, cyclobutane pyrimidine dimer; dsDNA, double-stranded DNA; k_B, the Boltzmann constant; NTP, ribonucleoside triphosphate; 8-oxoG, 8-oxoguanine; Pi, inorganic phosphate; PPi, pyrophosphate; RNAP, RNA polymerase; T, absolute temperature; TEC, ternary elongation complex

Regulation of transcription elongation via pausing of RNA polymerase has multiple physiological roles. The pausing mechanism depends on the sequence heterogeneity of the DNA being transcribed, as well as on certain interactions of polymerase with specific DNA sequences. In order to describe the mechanism of regulation, we introduce the concept of heterogeneity into the previously proposed alternative models of elongation, power stroke and Brownian ratchet. We also discuss molecular origins and physiological significances of the heterogeneity.

Introduction

For over 50 years, the mechanism of transcription by DNA-dependent RNA polymerase (RNAP) has been intensively investigated. Based on the accumulated knowledge, transcription is divided into three major stages, namely, initiation, chain elongation and chain termination. Initiation consists of RNAP binding to a specific promoter DNA and strand separation of the DNA that allows NTP binding to the template strand at the enzyme active site. Elongation is a repetitive but temporally discontinuous formation of phosphodiester bonds. Termination describes the complete dissociation of the RNA strand from the RNA-DNA hybrid retained in RNAP in response to specific signals.1 Different from initiation or termination, which occurs at specific DNA sites, elongation involves a gene-scale DNA tracking system, accompanied by bond formation at every base pair. Hence, specific DNA conformations/flexibilities, DNA lesions, and DNA-binding proteins are of specific concern in the regulation of elongation.

One of the major mechanisms that regulates elongation in all kingdoms of life is sequence-dependent pausing.2-3 In bacteria, pausing is important for the coupling of transcription with translation,4 and for providing opportunities for regulatory factors to bind the elongation complex.5-6 In eukaryotes, slow elongation, achieved by pausing around an exon-intron junction, allows spliceosome assembly, which increases the efficiency of alternative splicing.7 One of the main consequences of pausing is backtracking of RNAP relative to the RNA-DNA hybrid, which extrudes the 3′ end of the RNA from the active center.8-9 A deep sequencing study detected backtracking in a large fraction of paused polymerases in the yeast genome relative to nucleosome positioning.10 Backtracking plays a role to retain polymerases in promoter-proximal regions in order to maintain association with σ70 in Escherichia coli11 or NELF in eukaryotes,12,13 as a way to control specific gene transcription. Backtracking is also a mechanism to increase fidelity by providing a chance for proofreading.14 Contrary to such positive roles for backtracking, an irreversibly and stably backtracked complex forms a roadblock to replication of genomic DNA,15 and is highly toxic to the cell.16-18 Bacterial GreA/B, archael TFS, and eukaryotic TFIIIS rescue irreversibly backtracked complexes by promoting endonucleolytic cleavage and removal of the extruded 3′ end of transcripts.19-22 Bacterial NusG increases the elongation rate by inhibiting backtracking,23,24 which, for example, is critical for uninterrupted and rapid elongation of rRNA.25 Archaea and eukaryotes have Spt5, the counterpart of NusG.26-28 Therefore, polymerase pausing and protein factors that suppress pausing, especially related to backtracking, are essential for a broad variety of transcriptional regulatory steps in all kingdoms of life. So far, polymerase elongation and pausing have been discussed as parts of a homogeneous model that considers DNA as a monotonous polymer similar to a protein filament. However, pausing is a consequence of a heterogeneous effect, namely polymerase recognition of specific DNA sequences during elongation. In this review, we summarize previously proposed models of elongation and add the concept of this sequence-specific heterogeneity into the models to explain pausing. We also discuss the Brownian ratchet mechanism, which...
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does not accurately depict polymerase elongation, as was originally pointed out by Shimamoto.29

Power stroke and Brownian ratchet mechanisms of elongation including pausing

The movement of the RNA-DNA hybrid across the surface of the catalytic cleft in the enzyme is essential for elongation and for the generation of an empty site at the active center, where NTP binds. This movement is a one base pair forward translocation of RNAP along the DNA under conditions that prevent backward translocation or backtracking. Alternative mechanisms for the forward translocation have been proposed: the power-stroke model and the Brownian ratchet model. Their core concepts were taken from studies on unidirectional movement of motor proteins, such as myosin and kinesin, that are accompanied by ATP hydrolysis (reviewed in Refs30-34). In the power-stroke model, the conversion of chemical energy into mechanical work by a motor protein is coupled in the same elementary step. Chemical energy is directly generated by ATP hydrolysis in the protein or indirectly via conformational changes due to the release of inorganic phosphate (Pi). Hence, in the power-stroke model for elongation, the forward translocation is synchronized to either phosphodiester-bond formation or PPi release, with a transition state existing between the pre- and post-translocated states. Because the chemical energy is required for driving a forward translocation, the activation energy must be much higher than 0.5 \( k_B T \), the averaged value of thermal energy per degree of freedom, where \( k_B \) is the Boltzmann constant and \( T \) is the absolute temperature. This condition is sometimes mistakenly neglected, but necessary for the description with rate equations. If this mechanism is the case for elongation by \( E. coli \) RNAP, the transition state involves PPi release, because it has been shown that a nucleotide addition precedes translocation and PPi release.35 Below, we also discuss the case for pausing in the context of the power-stroke model.

In the Brownian ratchet model, there is no transition state with significant activation energy, and there are no energy barriers larger than the order of \( k_B T \). Hence, the transition state theory and rate equations cannot be applied to describe translocation (see Ref29 for more fundamentals). The forward translocation is not synchronized to the chemical steps. A net forward bias in the translocation would be generated by a cognate NTP binding to the active site of the elongation complex, which prevents backward translocation and simultaneously progresses the elongation to the next cycle by condensation of the NTP with the transcript.36,37 In an analysis for force-distance relationship of elongation by \( E. coli \) RNAP, the effective distance over which force acts is a single base pair during elongation within a pause-free sequence, which is equal to the moving distance of polymerase to complete translocation.38 This was interpreted as the absence of an alternative mechanism, such as a pause-dependent force that is required to progress elongation.

**Figure 1.** A model of sequence-specific pausing. (A) Pause-free elongation. RNA (orange), template DNA strand (gray), catalytic Mg\(^{2+}\) (magenta circle), and two RNAP domains (blue) involving 3' RNA separation from the hybrid, i.e., Switch 3 (arrow head) and lid (triangle) domains are shown. The 3' RNA-binding site (i) and the NTP binding site (i+1) are also indicated. (B) Elongation at the pausing site. The two sequence elements involved in transcription pausing are shown: (1) 3' ACGC 5' sequence in the transcribed DNA strand (gray) corresponding to the junction between the RNA-DNA hybrid and the downstream dsDNA in the elongation complex (indicated by shaded box); this sequence increases mobility/flexibility of the RNA/DNA backbones, which promotes fraying of the 3' RNA end. (2) G residue in the RNA at the upstream end of the hybrid contributes to immobilization of the hybrid in the catalytic cleft of RNAP by interacting with the Switch 3 domain in the post-translocated state, or by interacting with the lid domain in the pre-translocated state.
of the transition state between the pre- and post-translocated states, and consistent with Brownian ratchet translocation during smooth elongation.

Interestingly, the forward translocation of RNAP is smooth or restricted, depending upon the sequence of the DNA being transcribed. We are calling this heterogeneous tracking. This is distinguished from the mechanism of homogenous tracking by the motor proteins. Therefore, RNAP pauses at specific sequences during elongation, as exemplified by a biochemical study using yeast RNAP II and a transcription factor TFIIS mutant. TFIIS is known to induce endonucleolytic transcript cleavage near the 3′ RNA end by interacting with the active center of RNAP II. The study revealed that during RNAP II elongation, the cleavage-deficient TFIIS mutant carrying alanine substitutions in the catalytic loop, TFIISAA, specifically binds to the RNAP II that transiently pauses on timescales of 100 ms to 1 s and promotes its backtracking. RNAP II encountered such sequences every 10–100 bps, where the forward translocation was restricted. This was interpreted as forward translocation being the slowest process in the single nucleotide addition and limits elongation in the position sensitive to TFIISAA. It is noteworthy that TFIISAA has dominant lethal effect on yeast cells, suggesting a physiological significance for control of the sequence-dependent pausing.

Until now, pausing and pause-free elongation have been described in terms of homogeneous tracking according to the two pawl-ratchet (Brownian ratchet) model. This model proposes two rapid Brownian fluctuations during elongation: (1) translocation fluctuation of RNA-DNA hybrid and (2) conformational fluctuation of the RNAP active site including the bridge helix and the trigger loop, elements also involved in catalysis and substrate binding. The fluctuations are supposed to occur much more frequently than formation of a phosphodiester bond during pause-free elongation. If the same rapid fluctuations are independent of the transcribed sequence, they cannot be a source of sequence-specific pausing. In contrast to pause-free sequences, when RNAP encounters RNA-DNA hybrids of an unusual conformation or flexibility, the hybrid movement through the catalytic cleft may be restrained, hindering forward translocation. At these pause sites, the movement of the hybrid may become synchronized to phosphodiester bond formation or PPI release, followed by a return to the movement uncoupled to the chemical step in pause-free sites. In other words, at pre-translocation pause sites, the energy released from PPI dissociation is utilized for the forward translocation representing a transient switch to a power-stroke translocation mechanism. The DNA sites coding for pauses may have two consecutive pyrimidine/purine duplets in the non-coding DNA strand, where the downstream pyrimidine residue corresponds to the 3′ end of the RNA, and a purine residue corresponds to the RNA residue in the upstream end of the RNA-DNA hybrid in the elongation complex (Fig. 1). These sequences have been identified as pause sites in a number of bulk biochemical and single molecule transcription studies for E. coli RNAP and yeast/human RNAP II.

Interestingly, recent NMR studies revealed that pyrimidine/purine steps within dsDNA increase mobility of the sugar-moiety of the pyrimidine nucleotide, suggesting that the pyrimidine/purine neighbor spanning the junction between the RNA-DNA hybrid and the downstream DNA increases mobility of the sugar-phosphate backbones, possibly via sugar pucker rearrangements. This unique property may promote melting or fraying of the 3′ RNA end in the hybrid from the template DNA strand and prevent forward translocation (Fig. 1B). Furthermore, in other studies, a frayed 3′ RNA end was shown to interfere with phosphoryl transfer and promote backtracking. The purine residue at the 5′ end of the RNA strand in the RNA-DNA hybrid may hinder forward translocation by a steric clash with the catalytic cleft while in the single-stranded form or stabilization in the double-stranded form (Fig. 1B). An X-ray crystal structure of the bacterial elongation complex suggested that an RNAP Switch-3 domain that is involved in the RNA separation from the hybrid tightly interacts with the 5′ RNA end of the hybrid. Thus, bulky purines, contrary to pyrimidines, could interfere with the function of the Switch-3 domain. These sequence-specific effects may explain the pauses occurring every 10–100 bps, i.e., what we call heterogeneous tracking of RNAP during elongation.

In order to explain pausing caused by Brownian motion, one has to assume Brownian motion that is as slow as or slower than the process of single nucleotide addition. Such slow Brownian motion will be observed with very rare structural configurations of the elongation complex, which occur at low energy \(-kT\). An example is a combination of the limited orientation and position of DNA duplex, RNA-DNA hybrid, bridge helix and trigger loop. Pausing can be explained by the Brownian ratchet mechanism only if these structural configurations and the process of phosphodiester-bond formation have comparable frequencies. A new optical method that directly measures changes in the protein and the hybrid motions coupled to water in a broad time domain is required for verifying the Brownian ratchet mechanism.

Endogenous and exogenous mechanisms for translocation control

During elongation, RNAP frequently encounters alterations of dsDNA, nascent RNA structure, DNA lesions, and misincorporation events at the 3′ end of the RNA. Polymerase also faces histones in nucleosomes and other DNA binding proteins in front of advancing polymerase. These different types of encounters can block forward translocation. Some of these blocks have been confirmed experimentally, whereas the others require further validation (Fig. 2).

Elongating RNAP maintains a 9–10-bp RNA-DNA hybrid in its catalytic cleft (Fig. 2A), which is essential for high transcription processivity and maintenance of the correct DNA register. In bacterial RNAP, the hybrid is 9-bp and 10-bp long in the post-translocated and pre-translocated ternary elongation complexes (TECs), respectively. An expansion of the hybrid beyond this length is limited by the lid and rudder, which are small domains of RNAP located at the 5′ end of the hybrid and by the bridge helix at the 3′ end. Shortening of the hybrid below 7 base pairs leads to termination. In the yeast RNAP II, the hybrid appears to be 1 bp shorter than in the bacterial RNAP. However, this conclusion requires additional validation because all available structures of the yeast RNAP II were generated to
contain mismatched RNA/DNA of the hybrid upstream from the -8 position, which prevents backtracking and maintains structure. In TECs that have been crystallized, the hybrid always adopts a conformation that is intermediate between the A and B forms of a double helix. However, this observed uniform structure may be misleading because TEC crystallization typically involves a laborious selection to find base pair composition and length of the hybrid that is structurally stable and inhibits RNAP translocation in the crystal lattice. Thus, the X-ray results on the length and structure of the hybrid in TEC being strongly biased by the technical limits imposed.

The X-ray structures of TECs poised at different steps during translocation and catalysis revealed multiple orientations of the 3′ RNA end, the incoming cognate and non-cognate NTPs, and the template DNA residue (i+1) near the active center of the enzyme. Apart from local protein changes in the flexible regions surrounding the RNA-DNA hybrid and the downstream dsDNA, which include the switches, fork loop, trigger loop, lid and rudder elements, the structure of RNAP showed minor variations in different translocation intermediates. One wall of the catalytic cleft is part of a mobile clamp domain from the N-terminal part of the largest subunit of RNAP. Binding of the hybrid to the catalytic cleft induces closure of the clamp, leading to encirclement of the RNA-DNA hybrid and partial clamping of the downstream DNA duplex. An inner surface of the cleft facing the hybrid is composed of the switch and fork domains from the two largest RNAP subunits (Fig. 2A). Notably, the catalytic cleft contains small flexible loops forming sequence specific contacts with the major groove of the hybrid and cavities that accommodate bulges and/or flipped out residues in the hybrid. Because of the helical structure of dsDNA, translocation includes the rotation of the RNA-DNA hybrid and partial clamping of the downstream DNA duplex. Any local irregularity in the structure of the downstream DNA duplex or of the RNA-DNA hybrid (e.g., intrinsic bending or other non-canonical double-helical forms of the double helix) will interfere with the hybrid rotation and passage during translocation through the

![Figure 2. Cis- and trans-acting factors affecting translocation. (A) Structure of RNA-DNA hybrid and dsDNA in TEC by T. thermophilus (Tth) RNAP. The structural targets for translocation regulators are indicated by arrows. (B) A schematic structure of TEC: RNAP (blue oval), upstream and downstream dsDNA (gray cylinders), RNA-DNA hybrid (brown cylinder), transcription bubble (black line) and the bridge helix (green) are shown; the active center in RNAP is represented by a circle with i and i+1 subsites. The inset displays the pre-translocated configuration of the active center with DNA lesion in i site (yellow triangle) and the 3′ RNA residue in a frayed configuration in i+1 site. The left side shows cis-acting translocation inhibitors: bending, or other structural alteration of the hybrid, the front-end DNA duplex and hairpin in the nascent RNA interacting with RNAP (shown by curved arrow). The right side displays the trans-acting inhibitors: a drug molecule bound to bridge helix reducing its mobility/bending (red dot), protein factors bound to dsDNA, nascent RNA or RNAP, and the second RNAP molecule in a head-to-tail configuration (all in magenta).](image)
narrow catalytic cleft (Fig. 2B, left panel). Because the most extensive RNAP contacts with the hybrid are localized close to the active center, translocation is particularly sensitive to the chemistry and structure of the RNA-DNA base pairing at the 3′ end of the nascent RNA, as well as to DNA lesions in the template strand entering the active site. The 3′ residue RNA-binding site (i) and the NTP binding site (i+1) subdivide the active site, each having a different capacity to accommodate different base pairs as well as 3′ RNA-DNA mismatches and DNA lesions. For instance, RNAP appears to preferentially place pyrimidine and a mismatch at the 3′ RNA end (i site), and bulky DNA lesions at the i+1, which is less restrictive than the i site. This stabilizes the pre-translocated register. A lesion with photo-activated cyclobutane pyrimidine dimer (CPD) appears not to interfere with forward translocation before entering the i site. This tolerance is explained by the recent finding that translocation occurs without loading of this lesion to i+1 site, with the CPD being maintained on top of the bridge helix in a flipped-out configuration. A strong translocation block occurs after the CPD enters the i and i-1 sites. In contrast, the i site appears to be more tolerant than the i+1 site for other types of DNA lesions such as 8-oxoguanine (8-oxoG). Thus, translocation register is dictated by the ability of different parts of the catalytic cleft to accommodate DNA lesions and RNA-DNA base pairs of variable chemical structure. Finally, formation of a secondary structure in the nascent transcript immediately upstream from the RNA-DNA hybrid also interferes with translocation by anchoring the RNA in single-strand RNA binding site on RNAP located beneath the flap domain of β subunit. A direct interaction of the nascent 5′ RNA with the coil-coiled motif at the tip of the flap may also restrict the hybrid movement during translocation (Fig. 2B).
Exogenous effects

In Figure 2B, we classify several exogenous inhibitors to translocation: (1) protein roadblocks imposed downstream of RNAP, (2) protein tethers that bind to both the nucleic acids and RNAP, restraining translocation, (3) a leading RNAP that stalls during elongation to impose a roadblock to the trailing RNAP, and (4) small drug molecules that bind to the oscillating elements of RNAP, such as trigger loop, bridge helix, and forks.20,53,57,58 The first class includes nucleosomes, DNA-bound proteins such as transcription repressors (LacI, GalR, etc.), and limited movement imposed by adjacent RNAP molecules located in tandem transcribing the same DNA.66,67 A translocation block has been confirmed experimentally for RNAP II encountering nucleosomes53 and for RNAPs transcribing in tandem.67

Another class includes transcription elongation factor Nun encoded by H022 bacteriophage, which simultaneously binds RNAP, DNA and RNA-DNA hybrid at a significant distance from the RNAP active site to physically interfere with forward translocation (Vitiello et al., submitted). Nun has been shown to strongly arrest RNAP elongation in vivo and in vitro by stabilizing the enzyme in a pre-translocation register.58 Another similar example includes elongation factor NusG as found in Bacillus subtilis, which induces pausing of RNAP by interacting with the non-transcribed DNA strand in the transcription bubble as well as the upstream DNA duplex.59,70 Strepmodulin and tagetixin drugs, which act on bacterial RNAP, and α-amanitin, which acts on eukaryotic RNAP II, exemplify the third class of translocation inhibitors.57,71,72 These drugs bind to the bridge helix or the trigger loop (Fig. 2B) and reduce their mobility, which is required for translocation.36

The RNA-DNA hybrid in TECs is a target for control of translocation

Structural studies of Thermus thermophilus TECs suggested that the hybrid length increases to 10-bp immediately after bond formation and is followed by restoration of the original 9-bp hybrid after translocation.9,54 In this view, the 10-bp and 9-bp hybrids are the signatures of the pre- and post-translocated state of TEC, respectively. This assumption is consistent with the reported biochemical properties of T. thermophilus TECs assembled on a synthetic RNA/DNA scaffold containing 8–11-bp hybrids.73 TECs made with a 10-bp hybrid exhibited a high rate of pyrophosphorolysis relative to those made with a 9-bp hybrid,73 indicating that removal of 1 base pair at the -10 position of the hybrid causes forward translocation. Transcription factors and cis-acting elements in the DNA and nascent RNA may also promote or inhibit translocation by targeting the upstream end of the hybrid (Fig. 3). For instance, Nun protein of bacteriophage H022, which causes pre-translocation arrest in vivo and in vitro, likely inhibits translocation by stabilizing the -10 base pair position of the hybrid (Vitiello et al., submitted).68 In contrast, N protein of bacteriophage λ, which binds to the RNAP catalytic cleft near the RNA exit channel,74,75 may inhibit base pairing at the -10 position of the hybrid favoring forward translocation. NusG protein from E. coli interacts with the DNA duplex at the upstream end of the transcription bubble.76 A similar interaction has been proposed by Murakami and colleagues for the yeast Spt5 protein based on their structural analysis of Spt4/5 dimer bound to the clamp domain of archaeal RNAP.77 This upstream interaction may aid translocation by promoting re-annealing of DNA strands immediately upstream from the 9-bp hybrid position. NusG and Spt5 have been shown, as is the case for N protein,78 to stimulate RNAP elongation in vitro.42,75 The cis-acting hairpin formed in the RNA upstream from the -9 position of the RNA-DNA hybrid likely sequesters the -10 nt residue into the stem removing it from the hybrid (Fig. 3) during transcription termination.1 Interestingly, structural data also indicate that, in the post-translocated TEC, the -10 RNA base is sequestered in a protein pocket made by the Switch-3 domain of the bacterial β subunit (Fig. 3),49 and mutations in this domain exhibit severe translocation defects in vitro.79 Proteins such as N, Nun and NusG, bound close to the hybrid end, may alter the Switch 3 domain to decrease or increase sequestration of the -10 residue of RNA. It is not clear how the B. subtilis NusG and H022 Nun proteins have evolved to recognize the pre-translocated state of RNAP, as opposed to their homologs in E. coli, NusG and λ N, which target the post-translocated state.68,80 However, there are several lines of evidence that indicate that NusG can switch between transcription stimulation and inhibition modes depending on sequence context and/or bacterial species.69,70,81 Nun can also switch to N-like anti-termination activity in vivo52 and in vitro, as it has been shown by engineering of “hybrid” proteins between N and Nun in domain swapping experiment.82

Physiological significances of pausing in bacteria and in eukaryotes

The physiological role of transcriptional regulation via RNAP II pausing has been extensively investigated in eukaryotic cells. Promoter-proximal pausing of RNAP II controls expression of heat shock genes and proto oncosgenes in Drosophila and mammalian cells, respectively.2 ChIP-chip analysis indicated that these pauses are also detected at a large number of untranscribed genes in Drosophila and mammalian cells under variable experimental conditions.12,84-86 There are factors that induce promoter-proximal pausing of RNAP II, namely, DSIF (DRB sensitivity-inducing factor) and NELF (negative elongation factor).2,13,87 The involvement of these factors in the control of pausing implies that transcription pausing induced by translocation blocks followed by backtracking of RNAP may not be sufficient to explain all promoter-proximal pausing in vivo, although it has been proposed that pausing in Drosophila does involve backtracking of RNAP II.88 Below, we summarize recent findings on promoter-proximal pausing and discuss the pausing mechanism in the context of heterogeneous tracking of RNAP over specific sequences that can modulate the tracking.

A prominent role of promoter-proximal pausing was initially proposed to explain the rapid transcription response to environmental stimuli, such as heat-shock or cell differentiation, where the transcription initiation step is bypassed.84 However, recent studies indicate more divergent regulatory roles of promoter-proximal pausing; in Drosophila, pausing is employed to maintain a basal level of expression of genes coding for membrane receptors, transcriptional regulators involved in immune response, and factors affecting signal transduction pathways.89 Promoter-proximal
pausing is a key step in the concerted activation of genes involved in embryogenesis,\textsuperscript{30} and for shutting off the heat shock genes by NELF-mediated pathway during recovery from heat shock.\textsuperscript{31} Promoter-proximal paused RNAP II may additionally contribute to gene repression by blocking promoter access by other RNAP II molecules.\textsuperscript{32}

The recent development of high-throughput sequencing technology\textsuperscript{33} has allowed short RNA species retaining a 7-methylguanosine cap at their 5′ ends to be deep-sequenced.\textsuperscript{13} A large fraction of the 3′ ends of these short RNA species was mapped to Drosophila genomic positions that coincided with sites susceptible to permanganate, revealing that RNAP II often paused at positions located between +25 and +60 bps relative to transcriptional start sites throughout the genome.\textsuperscript{13} This result was consistent with the permanganate-ChIP-seq data, which indicated that promoter-proximal pausing occurs between +20 and +60 bps from transcriptional start sites at thousands of Drosophila promoters.\textsuperscript{93}

The positions of the sequenced 3′ ends were shifted downstream in cells defective for TFIIIS, indicating that the paused RNAP II may be backtracked.\textsuperscript{13} Interestingly, the estimated DNA melting temperature of the regions surrounding the 3′ end of these short RNA species was higher than those of most transcribed regions of the genome.\textsuperscript{13} The authors claimed that RNAP II pauses transiently within the downstream regions, where the RNA-DNA hybrid is less stable, and backtracks upstream from these regions to generate a more stable hybrid;\textsuperscript{13} however, the mechanical link to a translocation block of these sequences remains unclear.

In E. coli, promoter-proximal pausing has also been reported for rplN and ompX genes, as well as for the genes for λ and 82 phages.\textsuperscript{94,95} Gre factors are required for escape from these pause sites. A permanganate footprint analysis in E. coli revealed that RNAP associated with σ′\textsuperscript{79} frequently pauses at promoter-proximal regions carrying a -10 like sequence.\textsuperscript{96,97} This pausing was detected at 10-20% of the E. coli promoters, and it was enhanced by deletion of the greA gene.\textsuperscript{95} This suggested that promoter-proximal pausing caused by RNAP backtracking in E. coli is a general phenomenon. ChIP-chip analysis also indicated that RNAPs undergoing pausing near the transcription start sites in more than 20% of transcribed genes in E. coli.\textsuperscript{98} A fraction of paused RNAPs detected by ChIP-chip analysis likely includes “moribund complexes” that retain σ′\textsuperscript{98} and short abortive transcripts.\textsuperscript{99,100}

Formsations of the moribund complexes have been shown to depend on promoter sequence.\textsuperscript{101,102} The abortive type of RNA synthesis by the moribund complex is much slower than RNA synthesis by a productive elongation complex.\textsuperscript{93} The release of abortive RNA transcripts may require backtracking.\textsuperscript{29} Because the RNA retained in the moribund complex is shorter than 20-nt and has a 5′ triphosphate,\textsuperscript{103} deep sequencing of nascent transcripts that are cleaved into short fragments and have either 5′ triphosphate or 5′ monophosphate ends could allow discrimination of the moribund complexes from any paused elongation complexes.

ChIP-chip analysis revealed that promoter-proximal pausing in S. cerevisiae and B. subtilis is not as robust as in Drosophila and E. coli.\textsuperscript{104,105} In yeast, these pauses are typically sensitive to transcription cleavage factor TFIIIS: RNAP II pause sites identified by deep sequencing of nascent 3′ transcripts are shifted several bps downstream from their original location in cells lacking the dst1 gene coding for TFIIIS.\textsuperscript{10} This result indicates that pausing in yeast involves backtracking of RNAP II followed by TFIIIS-stimulated cleavage of the 3′ proximal transcript, which is required to resume elongation. ChIP-chip analysis also detected increase in promoter-proximal pausing caused by deletion of the greA gene in B. subtilis.\textsuperscript{105} These observations argue that promoter-proximal pausing due to RNAP backtracking appears to be a common theme in prokaryotes and eukaryotes and that the activity of TFIIIS or Gre factors that rescue backtracked RNAP plays a key role in gene regulation at transcription pause sites. Future characterization of the variable cis-acting DNA signals and trans-acting protein factors that regulate elongation by RNAP in vivo will help to establish a firm link between transcription pausing and heterogeneous tracking of polymerase.

**Disclosure of Potential Conflicts of Interest**

No potential conflicts of interest were disclosed.

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**References**

1. Komissarova N, Becker J, Solter S, Kireeva M, Kashlev M. Shortening of RNA-DNA hybrid in the elongation complex of RNA polymerase is a pre-requisite for transcription termination. Mol Cell 2002; 10:1151-62; PMID:12453422; http://dx.doi.org/10.1016/S1097-2522(02)00738-4

2. Aidelberg K, Lis JT. Promoter-proximal pausing of RNA polymerase II: emerging roles in metazoans. Nat Rev Genet 2012; 13:720-31; PMID:22986266; http://dx.doi.org/10.1038/nrg3193

3. Landick R. The transcriptional regulatory Rtf1p stimulates RNA chain synthesis after recruitment to elongation complexes by the exposed non-template DNA strand. Cell 2002; 109:193-203; PMID:12007486; http://dx.doi.org/10.1016/S0092-8674(02)00724-9

4. Roberts JW, Yarnell W, Bartlett E, Gao J, Marr M, Ko DC, Sun H, Roberts CW. Antitermination by bacte-riophage lambda Q protein. Cold Spring Harb Symp Quant Biol 1998; 63:319-25; PMID:10384296; http://dx.doi.org/10.1101/sqb.1998.63.319

5. Shukla S, Kavak E, Gregory M, Imashimizu M, Shrutinskiy B, Kashlev M, Oberdoerffer P, Sandberg R, Oberdoerffer S. CTCF-promoted RNA polymerase II pause sites identified by ChIP-chip analysis also indicated that RNAPs undergoing pausing near the transcription start sites in more than 20% of transcribed genes in E. coli.\textsuperscript{98} A fraction of paused RNAPs detected by ChIP-chip analysis likely includes “moribund complexes” that retain σ′\textsuperscript{98} and short abortive transcripts.\textsuperscript{99,100}

6. Artsimovich I, Landick R. The transcriptional regulatory Rtf1p stimulates RNA chain synthesis after recruitment to elongation complexes by the exposed non-template DNA strand. Cell 2002; 109:193-203; PMID:12007486; http://dx.doi.org/10.1016/S0092-8674(02)00724-9

7. Roberts JW, Yarnell W, Bartlett E, Gao J, Marr M, Ko DC, Sun H, Roberts CW. Antitermination by bacte-riophage lambda Q protein. Cold Spring Harb Symp Quant Biol 1998; 63:319-25; PMID:10384296; http://dx.doi.org/10.1101/sqb.1998.63.319

8. Nudler E, Mustaev A, Lukhanov E, Goldfarb A. The RNA-DNA hybrid maintains the register of transcription by preventing backtracking of RNA polymerase. Cell 1997; 89:33-41; PMID:9094712; http://dx.doi.org/10.1016/S0092-8674(00)80180-4

9. Komissarova N, Kashlev M. RNA polymerase switches between inactivated and activated states By translocating back and forth along the DNA and the RNA. J Biol Chem 1997; 272:15329-38; PMID:9182561; http://dx.doi.org/10.1074/jbc.272.24.15329

10. Churchman LS, Weissman JS. Nascent transcript sequencing visualizes transcription at nucleotide reso-lution. Nature 2011; 469:368-73; PMID:21248844; http://dx.doi.org/10.1038/nature09652

11. Perdue SA, Roberts JW. A backtrack-inducing sequence is an essential component of Escherichia coli (70)-dependent promoter-proximal pausing. Mol Microbiol 2010; 78:636-50; PMID:20382017; http://dx.doi.org/10.1111/j.1365-2958.2010.07347.x
25. Dutta D, Shatalin K, Epstein V, Gottesman ME, Nudler E. Linking RNA polymerase backtracking to genome instability in E. coli. Cell 2011; 146:533-43; PMID:21854980; http://dx.doi.org/10.1016/j.cell.2011.03.034

26. Nudler E. RNA polymerase backtracking in gene regulation and genome instability. Cell 2012; 149:1348-45; PMID:22726433; http://dx.doi.org/10.1016/j.cell.2012.06.003

27. Aguilera A, Garcia-Muse T, R loos: from transcriptional pauses to threats to genome stability. Mol Cell 2012; 46:115-24; PMID:22541554; http://dx.doi.org/10.1016/j.molcel.2012.04.009

28. Helmrich A, Ballarino M, Nudler E, Tora L. Transcription-replication encounters, consequences and genome instability. Nat Struct Mol Biol 2013; 20:412-8; PMID:23535296; http://dx.doi.org/10.1038/nsmb.2543

29. Borukhov S, Sagitov V, Goldfarb A. Transcript cleavage factors from E. coli. Cell 1993; 72:459-66; PMID:8431948; http://dx.doi.org/10.1016/0092-8674(93)90121-6

30. Izbann MG, Luse DS. The RNA polymerase II C-terminus of Spt5. PLoS One 2009; 4:e6918; PMID:19742326; http://dx.doi.org/10.1371/journal.pone.0006918

31. Hirreiter A, Damsma GE, Cheung AC, Klose D, Grohmann D, Vojnic E, Marin AC, Cramer P, Werner F. Spri-5 stimulates transcription elongation through the RNA polymerase clamp coiled-coil motif. Nucleic Acids Res 2010; 38:40-40; PMID:20079319; http://dx.doi.org/10.1093/nar/gkq335

32. Wada T, Takagi T, Yamaguchi Y, Ferdous A, Imai T, Hirose S, Sugimoto S, Yano K, Hartzog GA, Winston F et al. DSF, a novel transcription elongation factor that regulates RNA polymerase II processivity, is composed of human Spt4 and Spt5 homologs. Gen Dev 1998; 12:343-56; PMID:9450929; http://dx.doi.org/10.1101/gad.12.3.343

33. Shimamoto N. Nanobiology of RNA polymerase: biological consequence of inhomogeneity in reactant. Chem Rev 2013; 113:8400-22; PMID:24074222; http://dx.doi.org/10.1021/cr200066b

34. Oosawa F, Hayashi S. The loose coupling mechanism in molecular machines of living cells. Adv Biophys 1986; 22:151-83; PMID:2882695; http://dx.doi.org/10.1016/0065-227X(86)90085-5

35. Tyska MJ, Warshaw DM. The myosin power stroke. Cell Motil Cytoskeleton 2002; 51:1-15; PMID:11868092; http://dx.doi.org/10.1002/cm.10014

36. Arkin RD. Thermodynamics and kinetics of a Brownian motor. Science 1997; 276:917-22; PMID:9139648; http://dx.doi.org/10.1126/science.276.5314.917

37. Block SM. Kinesin motor mechanics: bending, stepping, tracking, gating, and limping. Biophys J 2007; 92:2986-95; PMID:17352031; http://dx.doi.org/10.1529/biophysj.106.067077

38. Ishii Y, Nishiyama M, Yanagida T. Mechano-chemical coupling of molecular motors revealed by single molecule measurements. Curr Protein Pept Sci 2004; 5:81-7; PMID:15078219; http://dx.doi.org/10.2174/1389203043486888

39. Malinen AM, Turtola M, Parthiban M, Vainonen L, Johnson MS, Belogurov GA. Active site opening and closure control translation of multisubunit RNA polymerase. Nucleic Acids Res 2012; 40:7442-51; PMID:22570421; http://dx.doi.org/10.1093/nar/gks383

40. Bar-Nahum G, Epstein V, Ruckenstein AE, Rafikov R, Mustaev A, Nudler E. A ratchet mechanism of transcription elongation and its control. Cell 2005; 120:183-93; PMID:15972495; http://dx.doi.org/10.1016/j.cell.2004.11.045

41. Guajardo R, Sousa R. A model for the mechanism of polymerase translocation. J Mol Biol 1997; 265:8-19; PMID:8995520; http://dx.doi.org/10.1006/jmbi.1997.0616

42. Abbondannieri EA, Greenleaf WJ, Shavetz JW, Landick R. Block direct observation of base-pair stepping by RNA polymerase. Nature 2005; 438:460-61; PMID:16284617; http://dx.doi.org/10.1038/nature04628

43. Izmashimizu M, Kireeva ML, Lukbowska L, Gote C, Datta AR, Strathern JN, Kashlev M. Intrinsically translocating RNA polymerase II with frayed RNA. Mol Cell 2009; 34:710-21; PMID:19560423; http://dx.doi.org/10.1016/j.molcel.2009.06.002

44. Vassylev DG, Vassyleva MN, Peredenera A, Tahirov TH, Artsimovitch I. Structural basis for transcription elongation by bacterial RNA polymerase. Nature 2007; 448:157-62; PMID:17581590; http://dx.doi.org/10.1038/nature05932

45. Kireeva M, Kashlev M, Burton ZF. Translocation by multi-subunit RNA polymerase II. Biochem Biophys Acta 2010; 1799:389-401; PMID:20079318; http://dx.doi.org/10.1016/j.bbagrm.2010.10.007

46. Walmacq C, Cheung AC, Kireeva ML, Lukbowska L, Ye C, Gote D, Strathern JN, Carell T, Godel P, Kashlev M. Mechanism of transcription elongation by RNA polymerase II. Role in cellular resistance to DNA damage. Mol Cell 2012; 46:18-29; PMID:22405652; http://dx.doi.org/10.1016/j.molcel.2012.02.006

47. Epstein V, Toulme F, Rahmouni AR, Borukhov S, Nudler E. Transcription through the roadblocks: the role of RNA polymerase cooperation. EMBO J 2003; 22:4719-27; PMID:12970184; http://dx.doi.org/10.1093/emboj/cdg452

48. Bintu L, Ishihashi T, Dangkulwanich M, Wu YY, Lukbowska L, Kashlev M, Bustamante C. Nucleosomal elements that control the topography of the barrier to transcription. Cell 2012; 151:738-49; PMID:23414536; http://dx.doi.org/10.1016/j.cell.2012.10.009

49. Vassylev DG, Vassyleva MN, Zhang J, Palangat M, Artsimovitch I, Landick R. Structural basis for substrate loading in bacterial RNA polymerase. Nature 2007; 448:163-8; PMID:17581590; http://dx.doi.org/10.1038/nature05931

50. Gnat AL, Cramer P, Ju P, Bushnell DA, Kornberg RD. Structural basis of transcription: an RNA polymerase II elongation complex at 3.3 A resolution. Science 2001; 292:1876-82; PMID:11313499; http://dx.doi.org/10.1126/science.1059495
64. Vakhnin AV, Vakhnin H, Babitzke P. Function of the Bacillus subtilis elongation factor TnasG in hairpin-dependent RNA polymerase pausing in the trp leader. Proc Natl Acad Sci U S A 2008; 105:163-6; PMID:18852477; http://dx.doi.org/10.1073/pnas.0708306105.

65. Yuzenkova Y, Roghanian M, Bohackeira A, Zenkin N. Tagerotin inhibits transcription by stabilizing pre-translocated state of the elongation complex. Nucleic Acids Res 2013; 41:9257-65; PMID:23935117; http://dx.doi.org/10.1093/nar/ gkt708.

66. Taike S, Sarafianos SG, Wang X, Hudson B, Sinena E, Mukhopadhyay J, Birkof Tor J, Leryn O, Ismail S, Clark AD Jr., et al. Inhibition of bacterial RNA polymerase by streptolysin: stabilization of a straight-bridge-helix-active center conformation. Cell 2005; 122:541-52; PMID:16122422; http://dx.doi.org/10.1016/j.cell.2005.07.017.

67. Kashkina E, Anikin M, Tahirov TH, Kocherlon SN, Vassylevy DG, Temiakov DK. Elongation complexes of Thermus thermophilus RNA polymerase that possess distinct transcriptional conformations. Nucleic Acids Res 2006; 34:4306-43; PMID:16914448; http://dx.doi.org/10.1093/nar/gk559.

68. Mishra S, Mohan S, Godavarthi S, Sen R. The interaction surface of the bacterial transcription elongation factor required for complex formation with an anti-terminator during transcription antitermination. J Biol Chem 2013; 288:28899-103; PMID:23916688; http://dx.doi.org/10.1074/jbc.M113.472209.

69. Gusarov I, Nueller E. Control of intrinsic transcription termination by N and NusA: the basic mechanisms. Cell 2001; 107:437-49; PMID:11719185; http://dx.doi.org/10.1016/S0092-8674(01)00582-7.

70. Tomar SK, Artsimovich I. NusG-S5 proteins—Universal tools for transcription modification and communication. Chem Rev 2013; 113:8604-19; PMID:23638618; http://dx.doi.org/10.1021/cr400064k.

71. Klein BJ, Bose D, Baker KJ, Yusoff ZM, Zhang X, Watanabe K, Bushnell DA, Kornberg RD. Structural basis of transcription termination by NusG and NusA: the basic mechanisms. J Mol Biol 1997; 273:813; PMID:9367773; http://dx.doi.org/10.1006/jmbi.1997.1327.

72. Kent T, Kashkina E, Anikin M, Temiakov D. Maintenance of RNA-DNA hybrid length in bacterial transcription. J Biol Chem 2013; 288:13497-504; PMID:19321439; http://dx.doi.org/10.1074/jbc.M113.504223.

73. Stevovyanova I, Artsimovich I. Functional analysis of the Thermus thermophilus transcription factor NusG. Nucleic Acids Res 2010; 38:7432-45; PMID:20695386; http://dx.doi.org/10.1093/nar/gkq623.

74. Robledo R, Gottesman ME, Weisberg RA. Lambda macR mutations convert HK022 Nun promoter to a terminator by transcription termination factor σ to a suppressor of termination. J Mol Biol 1990; 212:635-43; PMID:2139472; http://dx.doi.org/10.1006/jmbi.2000.3642; http://dx.doi.org/10.1006/jmbi.2000.3642.

75. Nickels BE, Mukhopadhyay J, Garrity SJ, Ebright RH, Hochschild A. The sigma 70 subunit of RNA polymerase mediates a promoter-proximal pause at the lac promoter. Nat Struct Mol Biol 2004; 11:54-50; PMID:15123345; http://dx.doi.org/10.1038/nsmb757.
98. Wade JT, Struhl K. The transition from transcriptional initiation to elongation. Curr Opin Genet Dev 2008; 18:130-6; PMID:18282700; http://dx.doi.org/10.1016/j.gde.2007.12.008

99. Kubori T, Shimamoto N. A branched pathway in the early stage of transcription by Escherichia coli RNA polymerase. J Mol Biol 1996; 256:449-57; PMID:8604130; http://dx.doi.org/10.1006/jmbi.1996.0100

100. Sen R, Nagai H, Hernandez VJ, Shimamoto N. Reduction in abortive transcription from the lambdaPR promoter by mutations in region 3 of the sigma70 subunit of Escherichia coli RNA polymerase. J Biol Chem 1998; 273:9872-7; PMID:9545328; http://dx.doi.org/10.1074/jbc.273.16.9872

101. Susa M, Kubori T, Shimamoto N. A pathway branching in transcription initiation in Escherichia coli. Mol Microbiol 2006; 59:1807-17; PMID:16553885; http://dx.doi.org/10.1111/j.1365-2958.2006.05058.x

102. Imashimizu M, Tanaka K, Shimamoto N. Comparative Study of Cyanobacterial and E. coli RNA Polymerases: Misincorporation, Abortive Transcription, and Dependence on Divalent Cations. Genet Res Int 2011; 2011:572689; PMID:22567357; http://dx.doi.org/10.4061/2011/572689

103. McClure WR. Mechanism and control of transcription initiation in prokaryotes. Annu Rev Biochem 1985; 54:171-204; PMID:3896120; http://dx.doi.org/10.1146/annurev.bi.54.070185.001131

104. Steinmetz EJ, Warren CL, Kuehner JN, Panbehi B, Ansari AZ, Brow DA. Genome-wide distribution of yeast RNA polymerase II and its control by Sen1 helicase. Mol Cell 2006; 24:735-46; PMID:17157256; http://dx.doi.org/10.1016/j.molcel.2006.10.023

105. Kusuya Y, Kurokawa K, Ishikawa S, Ogasawara N, Oshima T. Transcription factor GreA contributes to resolving promoter-proximal pausing of RNA polymerase in Bacillus subtilis cells. J Bacteriol 2011; 193:3090-9; PMID:21515770; http://dx.doi.org/10.1128/JB.00086-11