Examples of dynamic polymerase exchange have been previously characterized in model systems provided by coliphages T4 and T7. Using a dominant negative D403E polymerase (Pol) IIIα that can form initiation complexes and sequester primer termini but not elongate, we investigated the possibility of exchange at the *Escherichia coli* replication fork on a rolling circle template. Unlike other systems, addition of polymerase alone did not lead to exchange. Only when D403E Pol III was bound to a τ-containing DnaX complex did exchange occur. In contrast, addition of Pol IV led to rapid exchange in the absence of bound DnaX complex. Examination of Pol III with varying contrast, addition of Pol IV led to rapid exchange in the absence of clamp processivity factor (holoenzyme (HE) of DNA polymerase III)*, but Not Polymerase IV, Must Be Bound to a τ-Containing DnaX Complex to Enable Exchange into Replication Forks*

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Received for publication, March 2, 2016, and in revised form, April 7, 2016 Published, JBC Papers in Press, April 7, 2016, DOI 10.1074/jbc.M116.725358

DNA Polymerase III, but Not Polymerase IV, Must Be Bound to a τ-Containing DnaX Complex to Enable Exchange into Replication Forks

As with all cellular replicases, the DNA polymerase (Pol) III holoenzyme (HE) of *Escherichia coli* is tripartite with a sliding clamp processivity factor (β3), a clamp loader (DnaX complex, DnaXβδχλφ), and an associated replicative polymerase (Pol, αεθδ) (for a review, see Ref. 2). In *E. coli* and many other bacteria, DnaX encodes two products: a shorter γ polymerase that has ATPase activity and the ability to support clamp loading on single-stranded DNA and a longer τ polymerase that has additional domains that interact with the DnaB replication helicase and Pol III. Cells that cannot express τ, the interdomain loop of one subunit of the sliding clamp and gaps between fragments, possessed the ability to challenge pre-established replication forks, and displayed equivalent susceptibility to challenge by exogenous D403E Pol IIIα. These findings reveal that redundant interactions at the replication fork must stabilize complexes containing only one τ. Previously, it was thought that at least two τs in the trimeric DnaX complex were required to couple the leading and lagging strand polymerases at the replication fork. Possible mechanisms of exchange are discussed.

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An *E. coli* rolling circle replication system has been developed that exploits a 409-nt flapped circular template that has a 50:1 asymmetric GC content, allowing convenient distinction, quantification, and labeling of leading and lagging strands. The asymmetric GC distribution allows specific slowing of lagging strand synthesis by substitution of dGDPNP for dGTP. The V max for insertion of this analog is lower than the natural nucleotide, and the K m is higher, allowing “dialing in” a desired elongation rate without reducing the nucleotide to a level where its concentration is depleted during the progress of the reaction. Because the lagging strand half of the Pol III HE cycles when a new primer is made, even when the preceding Okazaki fragment is incomplete, gaps are left between fragments when the lagging strand rate is decreased to a level where cycling is induced before fragment synthesis is complete (4).

The concept of dynamic processivity and dynamic polymerase exchange was first established in the bacteriophage T4 system (5, 6). T4 replication is highly processive, but exogenous polymerase can exchange into the fork and inhibit coupled rolling circle replication more rapidly than the dissociation rate of the T4 DNA polymerase, gp43. These results suggested that gp43-D408N actively displaces gp43 rather than passively binding template after dissociation of wild-type gp43. It was proposed that the C terminus of the incoming polymerases binds the interdomain loop of one subunit of the sliding clamp and displaces the polymerase at the replication fork (6).

Similarly, with bacteriophage T7, a complex of wild-type DNA polymerase (gp5) and its processivity factor, *E. coli* thioredoxin, readily exchanges with excess mutant gp5-thioredoxin complexes (gp5-Y526F/thioredoxin) without affecting processivity. Replacing tyrosine 526 with phenylalanine in the nucleotide binding site makes the mutant gp5 resistant to inhibition of ddNTPs but does not affect its ability to bind to other protein components and elongate. Both strand displacement synthesis and leading strand synthesis in a coupled reaction
Pol III* Exchange at Replication Forks

initiated by gp5-Y526F/trx are inhibited upon addition of gp5/trx and ddNTPs, indicating polymerase exchange (7). Exchange was not observed in a single-stranded DNA replication reaction where helicase was not included. Thus, it was proposed that an exogenous polymerase binds the hexameric T7 helicase and exchanges with polymerase that transiently dissociates (7).

YFP-tagged E. coli Pol III α proteins have been detected by single-molecule microscopy in live cells bound near the replication fork (8). Repetitive bursts of fluorescent Pol III α were observed in living cells that were interpreted as new polymerases continuously exchanging with the DnaX complex organizer at replication forks with the synthesis of each Okazaki fragment. This appeared to be inconsistent with the 4.9-h mean lifetime (k_off = 5.7 × 10^{-5}) observed for α-τ DNA dissociation (10) and is the primary subject of this investigation. In addition to a putative exchange of Pol III into the replication fork, exchange of other polymerases, particularly Pol IV, has been well documented (11–16). Two different mechanisms have been proposed for Pol IV exchange. One, called the toolbelt modification in light of a recent elegant cryo-EM structure that shows that both of the canonical binding sites within the one of the two canonical clamp binding sites, whereas the other predicts recruitment mediated by a direct contact of Pol IV with both Pol III* and an accessory site on the rim of the β2 sliding clamp, subsequently displacing Pol III to allow binding by Pol IV (12–15, 17–19). If operative, the toolbelt model will require modification in light of a recent elegant cryo-EM structure that shows that both of the canonical binding sites within β2 are occupied by interacting Pol III α and ε subunits of the Pol III HE (20). We pursued this issue in this work and compared the requirements for exchange with those of Pol III to determine whether similar mechanisms were used.

Recently, we have established that the major cellular form of Pol III HE is Pol III_{2γδδ′}χψ, disproving proposals that the cellular replicase contains three τs and three polymerases (3). One of the observations that was initially used to support the three-polymerase model was that three Pol III assemblies synthesized DNA in a coupled fork system with smaller gaps between Okazaki fragments (21). We reinvestigated this observation using a minicircle replication system we developed (4). Our findings are presented here.

Experimental Procedures

Nucleotide Analogs and Radioactive Nucleotides—dGDPNP was custom-synthesized for this project by TriLink Biotechnologies. The radioactive nucleotides α-[³²P]dATP (3000 Ci/mmol), α-[³²P]dCTP (3000 Ci/mmol), and α-[³²P]dGTP (3000 Ci/mmol) were from PerkinElmer Life Sciences.

Proteins—The following proteins were purified as described: D403E Pol III α (22), ε (23), θ (24); Pol III (25); wild-type and mutant DnaX complexes, including τ_{γ}, τ_{γ2}, and τ_{γχ}, Pol IV (DinB) (27); β2 (28); SSB4 (29); DnaB6 (30); DnaC (4); DnaG (29); DnaT3 (4); PriA (4); and PriB2 (4). The purifications of the various forms of the DnaX complexes were performed by high-resolution FPLC chromatography on Mono S columns, providing baseline resolution between the forms with no cross-contamination, as revealed by rechromatography (26). D403E Pol III was assembled by incubating D403E Pol III α, ε, and θ at a molar ratio of 1:1:1 on ice for 15 min. Three forms of Pol III* containing D403E Pol III α were assembled: D403E Pol III_{2γδδ′}χψ, D403E Pol III_{2γδδ′}χψ, and D403E Pol III_{2γδδ′}χψ. In each case, they were assembled by incubating D403E Pol III and the corresponding DnaX complex (τ_{γ}, τ_{γ2}, or τ_{γχ}) on ice for 15 min at a ratio of 6:1 for D403E Pol III_{2γδδ′}χψ and DnaX complex, at a ratio of 4:1 for D403E Pol III_{2γδδ′}χψ, and at a ratio of 2:1 for D403E Pol III_{2γδδ′}χψ. Likewise, wild-type Pol III* (Pol III_{2γδδ′}χψ) and mutant Pol III* bearing the ATPase K515E mutation in the γ subunit (Pol III_{2γδδ′}χψ) were assembled by incubating Pol III and the corresponding DnaX complex (τ_{γγδδ′}χψ, or τ_{γγγ′δδ′}χψ) on ice for 15 min at a ratio of 4:1.

Rolling Circle Reactions—Rolling circle reactions were performed as described previously (4). 20 nm minicircle DNA template, 2 μM SSB, 100 nM βγ, 72 nM DnaB6, 100 nM DnaG, 2.5 nM Pol III*, 160 nM PriA, 50 nM PriB2, 333 nM DnaTγ, and 108 nM DnaC were incubated with 5 μM ATPγS, 200 μM CTP, 200 μM UTP, and 200 μM GTP for 5 min at 30 °C to form an initiation complex. The reaction buffer was 10 mM magnesium acetate, 70 mM KCl, 50 mM Heps (pH 7.5), 100 mM potassium glutamate, 20% glycerol, 200 μg/ml bovine serum albumin, 0.02% Nonidet P-40, and 10 mM dithiothreitol. The reaction was started by addition of 1 mM ATP and 100 μM dNTPs. After 4 min, α-[³²P]dCTP or α-[³²P]dGTP were added to quantify leading and lagging strand synthesis, respectively. After an additional 4 min, the reaction was quenched with EDTA (140 mM final concentration), and samples were analyzed by liquid scintillation counting after trichloroacetic acid precipitation as described previously (31).

Competition Experiments with D403E Pol III—For control experiments that analyzed competition of D403E Pol III, wild-type Pol III and varying D403E Pol III were combined first and then mixed with the other components of the rolling circle reaction to form an initiation complex as described in the preceding paragraph. Wild-type Pol III was added at a 2-fold molar excess of the DNAx complex (2.5 nM final concentration) and depended on the stoichiometry of the τ subunit. For τ_{γ} complex, Pol III was added at 15 mM; for τ_{γ2} complex, Pol III was added at 10 mM; and for τ_{γχ} complex, Pol III was added at 5 mM. After 5 min, 1 mM ATP, 100 μM dNTPs, and α-[³²P]dCTP or α-[³²P]dGTP were added to start the reaction. After 8 min, the reaction was quenched with 140 mM EDTA final concentration. For experiments that analyzed competition of D403E Pol III added to an ongoing reaction, reactions were started with non-radioactive nucleotides. After 4 min of elongation to establish coupled replication, competitor D403E Pol III was added along with α-[³²P]dCTP or α-[³²P]dGTP, and the reaction was allowed to continue for an additional 4 min. The data shown in this figure are the results of experiments conducted in a similar format multiple times with the same conclusions.
used to label the leading and lagging strands in reactions containing Pol III* reconstituted with four different DnaX complexes: $\tau_2 \delta \delta \gamma' \psi$, $\tau_2 \gamma \delta \delta \chi \psi$, $\tau_2 \gamma \delta \delta \chi' \psi$, and $\gamma_3 \delta \delta \chi' \psi$. The reaction products were analyzed for strand-specific incorporation by liquid scintillation counting of trichloroacetic acid-precipitable products. Rolling circle reactions were also conducted in the presence of either 100 $\mu$M dGTP or dGDPNP (30, 60, 120, and 240 $\mu$M) using Pol III* reconstituted with four different DnaX complexes ($\tau_2 \delta \delta \chi' \psi$, $\tau_2 \gamma \delta \delta \chi' \psi$, $\gamma_3 \delta \delta \chi' \psi$, and $\gamma_3 \delta \delta \chi' \psi$) using $\alpha$-[$^32$P]dATP to label both strands. The reaction products were analyzed by alkaline-agarose gel electrophoresis.

Rolling Circle Replication with Pol III* Containing Wild-type and Mutant $\gamma$ DnaX Subunits—Rolling circle reactions were set up using two different forms of Pol III*: one with a wild-type form of the $\gamma$ subunit ($\tau_2 \gamma$) and one with K51E$\gamma$ ($\tau_2 \gamma^{M}$, ATPase-defective). Both forms of the polymerase were functional in holoenzyme reconstitution experiments (26). For incorporation experiments, the amount of dGTP was varied (0, 1.6, 3.1, 6.3, 12.5, 25, 50, and 100 $\mu$M). For analysis of rolling circle reaction products on alkaline-agarose gels, reactions were set up in the presence of $\alpha$-[32P]dCTP, $\alpha$-[32P]dGTP, or $\alpha$-[32P]dATP. The nucleotide $\alpha$-[32P]dCTP specifically labels the leading strand, $\alpha$-[32P]dGTP specifically labels the lagging strand, and $\alpha$-[32P]dATP labels both the leading and lagging strands. A total of 50,000 cpm was loaded to each lane.

Alkaline-Agarose Gel Electrophoresis—For the analysis of leading and lagging strand products, samples were mixed with 30 mm NaOH, 2 mm EDTA, 2% glycerol, and 0.02% bromphenol blue and fractionated on 0.6% alkaline-agarose gels (10 $\times$ 14 cm) for 16 h at 60 mA ($\sim$20 V) in a running buffer of 30 mm NaOH and 2 mm EDTA. Gels were fixed in 8% (w/v) trichloroacetic acid, dried onto DEAE paper, autoradiographed on storage phosphor screens, and scanned with a Phosphorimager. Okazaki fragment length was determined as described previously (4).

PolIII* Exchange Assay on a Single-stranded Template—This assay measures the ability of a Pol III* competitor to exchange with a Pol III* undergoing DNA synthesis on a single-stranded template. Holoenzyme reactions (22) were assembled on ice using WT Pol III* (40 nM, experimentally determined to be at saturation), $\beta_2$ (6 nM), DnaG primase (60 nM), SSBs (0.6 $\mu$m), M13Gori DNA (2.3 $\mu$m as circle), four NTPs (0.2 mm each), and 1 mm ATP. The reaction buffer was 10 mm magnesium acetate, 70 mm KCl, 50 mm Hepes (pH 7.5), 100 mm potassium glutamate, 20% glycerol, 200 $\mu$g/ml bovine serum albumin, 0.02% Nonidet P-40, and 10 mm dithiothreitol. Initiation complexes were formed at 30 $^\circ$C for 3 min. Control reactions demonstrated that initiation complexes were completely formed by 1 min. The reaction was initiated by adding dNTPs (48 $\mu$m each dATP, dCTP, dGTP; 18 $\mu$m dTTP; 100 cpm [3H]/pmol dNTPs) plus competitor Pol III* (D403E Pol III*, WT Pol III*, or buffer control) at various concentrations (0, 24, 48, 94, 188, 375, and 750 nM). Reactions were performed in triplicate. DNA synthesis was quenched after 30 s by the addition of EDTA (140 mm), and samples were analyzed by liquid scintillation counting after trichloroacetic acid precipitation. 1 unit of Pol III* activity is 1 pmol of total nucleotide incorporated into acid-insoluble DNA per minute.

Results

Exogenous D403E Pol III Does Not Exchange with Pol III at the Replication Fork—A proposal has been made that free Pol III can exchange with the replication fork by dissociation and reassociation with $\tau$ (8). This observation seemed to be in conflict with the observation that the Pol III $\alpha$-$\tau$ interaction has a lifetime of 4.9 h (10). We employed a dominant negative mutant E. coli Pol III $\alpha$ (D403E) that is able to form an initiation complex at the replication fork but cannot elongate (22, 32) to test this proposal. When D403E Pol III was premixed with wild-type E. coli Pol III to form an initiation complex, rolling circle DNA replication was inhibited in proportion to the concentration of D403E Pol III as expected (Fig. 1A). When D403E Pol III was added to an ongoing rolling circle DNA replication reaction, inhibition was not observed (Fig. 1B), indicating that exchange of Pol III does not occur over the 8-min time course of the reaction.

Exogenous D403E Pol III* Exchanges with Pol III* at the Replication Fork—We next tested whether exogenous D403E Pol III* was able to compete with wild-type Pol III* at the replication fork. In a control reaction, D403E Pol III* inhibited a rolling
circle reaction when premixed with wild-type Pol III* to form the initiation complex (Fig. 2A). Surprisingly, D403E Pol III*, when added after initiation, inhibited an ongoing rolling circle reaction. Lagging strand synthesis was marginally more susceptible to challenge, being reduced 50% with half the level of D403E Pol III* challenge compared with leading strand synthesis. (Fig. 2B).

DNA Pol IV Exchanges into the Replication Fork with Different Protein Requirements than Pol III—We repeated the experiments we performed to assess exchange of free Pol III, substituting Pol IV. Pol IV incorporates nucleotides very slowly, so inhibition of elongation can be used as an assay for exchange (13, 14, 19). We observed that Pol IV exchanges just as effectively whether added before or after initiation complex formation by Pol III HE (Fig. 3). In both cases, lagging strand synthesis is more susceptible to a challenge by Pol IV than leading strand synthesis. (Fig. 2B).

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Okazaki Fragment Size and Gaps between Fragments Are the Same, Regardless of the τ Content of Pol III*—Because DnaX complexes contain a total of three γ and/or τ subunits, four different stoichiometries are possible: τγ5δ5χψ, τγ5δ5χψ, τγ5δ5χψ, and γ5δ5χψ. We tested the function of all four DnaX complexes in a minicircle replication system. We found that all τ-containing DnaX complexes functioned equivalently in terms of the level of leading and lagging strand synthesis (Fig. 4A). All τ-containing DnaX complexes synthesized Okazaki fragments of the same length (Fig. 4B). The extent to which Okazaki fragments were decreased when made shorter by decreasing dGDPNP concentration was identical for all τ-containing complexes (Fig. 4B). DnaX complex lacking τ (γ complex) was unable to function under the experimental conditions used (substoichiometric Pol III*, Fig. 4).

The observation that Pol III HE containing only one τ can function in coupled synthesis was surprising. We previously thought that two τs were required to couple leading and lagging strand synthesis. In light of this finding, we further explored the stability of replication forks reconstituted with single τ Pol III HE by testing whether they are more susceptible to challenge by D403E Pol III or Pol III* (supplemental Figs. 1 and 2). We observed that, like other forms of Pol III HE, the single τ form was resistant to a challenge by Pol III alone and similarly susceptible to a challenge by D403E Pol III*. However, a single τ Pol III HE is not sufficiently stable to function in a flow system, presumably because it dissociates sufficiently frequently to be
carried away in the flow solution, removing it from an immobilized template (33). In a DnaB-dependent leading strand-tethered bead single-molecule experiment, it was observed that DnaX complex containing only one \( \tau \) was functional (34).

We also observe that forks reconstituted with single-\( \tau \) Pol III* are just as resistant to a challenge by D403E Pol III* as \( \gamma \) forms (supplemental Fig. 1). Similarly, D403E Pol III*s containing one or three \( \tau \)s are similarly effective as a challenge (supplemental Fig. 2), but “Pol III*” reconstituted with \( \gamma \) complex (no \( \tau \)) is unable to exchange into replication forks (supplemental Fig. 3).

**DnaX Complex with One Inactive ATPase Is Not Competent for Rolling Circle DNA Replication**—To determine whether one inactive DnaX ATPase among the three present in DnaX complex affects coupled leading and lagging strand synthesis, a rolling circle reaction was conducted with reconstituted Pol III* containing either one wild-type \( \gamma \) DnaX subunit or one mutant K51E \( \gamma \) DnaX subunit. The mutation occurs within the Walker A motif and abolishes ATP binding (26). In the presence of Pol III* containing mutant \( \gamma \), the total level of rolling circle DNA synthesis is decreased 7-fold (Fig. 5A). Specific examination of leading and lagging synthesis using alkaline gel electrophoresis shows that some leading strand synthesis occurs but no detectable lagging strand synthesis (Fig. 5B).

**D403E Pol III* Can Also Exchange into Elongating Pol III HE on Long Single-stranded Templates**—To determine whether DnaB6 or other components unique to replication forks were required for exchange, we challenged Pol III HE elongating a long single-stranded template. We preformed initiation complexes using wild-type Pol III* and \( \beta \), on DnaG primase-synthesized primers on SSB-encoated M13 circles containing a G4 origin and then added challenge D403E Pol III* together with dNTPs to initiate the reaction. We observe the same level of inhibition on these simpler single-stranded templates as we do in the coupled rolling circle assays (Fig. 6). Thus, the DnaB6 helicase is not a required component to enable exchange.

**Discussion**

Unlike the model T4 and T7 systems, core Pol III from *E. coli* cannot exchange into replication forks. It requires association with a \( \tau \)-containing DnaX complex. Thus, the *E. coli* system is fundamentally different from these two systems. Further contrasting the model systems, T7 dynamic exchange is thought to be mediated by a polymerase-helicase interaction (7). In our *E. coli* system, exchange occurs on single-stranded templates in the absence of helicase (Fig. 6). Exchange of Pol III with Pol IV or Pol II also occurs on single-stranded templates independently of helicase (13–15, 35, 36).

We examined the ability of Pol III HE to function in coupled rolling circle replication with variable \( \tau \) content. The \( \tau_3 \), \( \tau_2 \), and \( \tau_1 \) forms functioned equivalently in terms of levels of synthesis but also Okazaki fragment size distribution, even in the presence of decreasing dGDPNP concentrations (Fig. 4). At low dGDPNP concentration, lagging strand synthesis is slowed specifically, whereas leading strand synthesis progresses at a normal rate along with the synthesis of new primers by DnaG primase. This induces the lagging strand polymerase to cycle before synthesis is complete, leaving large gaps (4). That the size of fragments is identical regardless of \( \tau \) content suggests that all three forms of Pol III HE experience the same dynamics at the replication fork, initiating and cycling between Okazaki fragment synthesis at nearly equivalent rates, resulting in the same length of gaps between fragments.

It has been proposed that \( \tau_3 \) Pol III HE leaves shorter gaps between Okazaki fragments than Pol III HE containing two \( \tau \)s. These experiments were performed by a low-resolution single-molecule technique where a single pixel covered 600 nucleotides, approaching the size of the 1- to 2-kb Okazaki fragments found under physiological conditions. To surmount this obstacle, conditions were altered to increase Okazaki fragment size to 12 kb (21). Our results do not contradict these results.
obtained under non-physiological conditions but do show that gap size is the same under physiological conditions. Furthermore, a trimeric polymerase is not the primary form of Pol III HE in cells (3), making the single-molecule results less relevant.

Pol III HE-like activity, reconstituted in the presence of γ complex (no τ), was inactive in rolling circle synthesis under the experimental conditions employed: substoichiometric Pol III*.

This is consistent with the requirement for a τ-DnaB helicase association to stabilize replication forks (37). We were initially surprised by the ability of Pol III HE that contained only one copy of τ to function efficiently (Fig. 4). When free in solution, Pol III HE must contain at least two τs to interact with DnaB6 under physiologically relevant concentrations (38). However, we have observed3 that the presence of ATPγS and an associated replication fork significantly strengthens the interaction of Pol III HE with τ. Perhaps the synergy of a multimeric complex stabilizes the interaction sufficiently to permit single τ HE interaction. We do not believe that this form of HE exists in cells to a significant extent, but its function may be a reflection of a redundancy of stabilizing interactions at the replication fork.

Our finding that Pol III* can exchange into replication forks may reconcile the conflict between our finding that free polymerase cannot exchange with Pol III at the replication fork and the single molecule experiments that were interpreted to indicate that free Pol III could dynamically exchange at replication fork (8). The bursts of fluorescence they observed could have resulted from Pol III* exchange at the replication fork because it is not possible to distinguish between Pol III and Pol III* from the fluorescence assay they conducted. Alternatively, they could have observed exogenous Pol III required for other processes associated with the nascent replication product, such as mismatch repair. Involvement of the DnaX complex in Pol III* exchange indicates that interactions mediated by the DnaX complex are important for polymerase exchange.

DnaX complexes containing a mixture of inactive and wild-type DnaX subunits can function on long single-stranded templates in 5-min reactions (26). Examination of the kinetics of initiation complex formation reveals a 3500-fold lower rate for DnaX complexes containing a single inactive ATPase subunit (39). We exploited the rolling circle system we recently developed that permits use of substoichiometric Pol III* (4) to provide a sensitive test for function under physiological conditions. We observe a modest level of leading strand synthesis, consistent with the ability of mixed mutant complexes to function on long single-stranded templates, but could not detect any lag-

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ging strand synthesis, suggesting that Pol III HE must have three active ATPases to support the rapid initiation complex formation and cycling required for lagging strand synthesis (Fig. 5). Because of the frequent initiation complex formation on the lagging strand in a rolling circle reaction, slowing the rate of initiation complex formation by the inactive ATPase exerts a much more severe effect than in a single-stranded DNA replication reaction.

For *E. coli* Pol III to exchange into a replication fork requires sequestered DnaX complex. γ Complex, which can load β₂ clamps onto DNA but cannot bind Pol III α, does not substitute. Thus, some component of τ-containing DnaX complexes must be responsible for polymerase exchange. Because τ complexes are functional and γ complexes are not, the simplest explanation would be that domain IV and/or V, present only in τ, interacts with elongating Pol III HE and triggers exchange. However, we have discovered situations where τ is required only as a bridge to hold χτ together in the same complex with Pol III; for example, enabling replication through SSB on a single-stranded template in the absence of β₂ and for stabilizing leading strand polymerase on a forked template so that it can strand-displace, in the absence of helicase, by interaction with SSB bound to the displaced strand (31, 40). These findings have been confirmed and extended (41, 42). It has also been observed that only τ-containing DnaX complexes will support initiation complex formation using slowly hydrolyzable ATPγS (32, 43), likely by stabilizing intermediates during slow initiation complex formation. The δ subunit is involved in either opening β₂ complexes or trapping open complexes from solution as part of the initiation complex formation reaction. The δ subunit could be involved in modulating β₂ structures to enable access by other proteins. Thus, components of the DnaX complex other than τ could facilitate exchange.

We have recently demonstrated that the signaling model is exclusively operational as a mechanism to drive Pol III* release and recycling during Okazaki fragment synthesis (4). The availability of a new primer is required as the signal to initiate cycling. DnaG primase is not required when primers are provided exogenously. In a simpler model system, we observed that τ-containing DnaX complex, exogenous primer template, and ATP, but not ATPγS, could modestly accelerate Pol III* release from chip-bound oligonucleotide templates (44). τ-Containing DnaX complexes can rapidly chaperone Pol III onto nascently loaded β₂ and accelerate the overall initiation complex formation reaction 100-fold, a requisite for the physiological rate of Okazaki fragment formation (39). Combining this information, we have proposed a working model that τ-containing DnaX complexes may also chaperone Pol III off of nascent Okazaki fragments when a primer becomes available for the next fragment (4). It is possible that the DnaX complex in exogenous Pol III* complexes competes in the process. This could explain the increased sensitivity of lagging strands to exchange by a challenging polymerase (Fig. 2).

Other mechanisms might be responsible for permitting access to primer termini on the leading strand. Because the leading strand polymerase is not free to rotate around the template strand because it is linked to the lagging strand polymerase, DNA behind the polymerase will be underwound. The polymerase may need to transiently release the primer terminus to relieve this negative superhelical tension, allowing the DNA to rotate within the channel of the bound, processively elongating Pol III HE (9, 45–47). This might grant access to the primer terminus, enabling an exchange reaction.

Alternatively, direct protein contacts between the invading Pol III* with the displaced Pol III HE might be required. Clearly, interactions of Pol IV with β₂ are important to enable exchange (13–15, 19). Mutant Pol IV that lacks its β₂ interacting motif can still exchange, albeit at higher concentrations, suggesting an interaction between Pol IV and a component of Pol III* (19). It is not clear whether these interactions act simultaneously or sequentially to mediate exchange. Interaction of Pol IV with Pol III* may trigger a conformational change so that at least one of the two mutual binding sites within β₂, sequestered by Pol III α and ε during elongation (20), become accessible. Consistent with this possibility, residues within or near the start of α helix 5 in Pol IV, which are well removed from the two distinct β₂ clamp binding motifs, are required to displace Pol III* from β₂ assembled on DNA (17). Pol III* exchange could proceed by a similar mechanism, except the primary interacting motif would have to reside within τ-containing DnaX complexes. Alternatively, the mechanism of Pol III* exchange could be different from that used by Pol IV, consistent with different protein requirements for exchange.

These issues surrounding polymerase exchange at replication forks are general and important. Unpublished results show that *Bacillus subtilis* Pol C actively evicts an otherwise highly processive DNA starting ~35 nucleotides after initiating synthesis from a nascent primer during Okazaki fragment synthesis. Similarly, Pol δ must displace Pol α during eukaryotic Okazaki fragment synthesis. Very importantly, a host of error-prone or bypass polymerases must exchange with their normal replicative counterparts to maintain genome stability in all cell types. Establishing general mechanisms for exchange should be a challenging and fruitful area of investigation for future investigators.

**Author Contributions**—C. M. designed the study. Q. Y. and P. R. D. performed the experiments. C. M. and M. D. S. wrote the paper. All authors reviewed the results and approved the final version of the manuscript.

**Acknowledgments**—We thank Diane Hager for preparation of the figures and references.

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