A hand-off of DNA between archaeal polymerases allows high-fidelity replication to resume at a discrete intermediate three bases past 8-oxoguanine

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ABSTRACT
During DNA replication, the presence of 8-oxoguanine (8-oxoG) lesions in the template strand cause the high-fidelity (HiFi) DNA polymerase (Pol) to stall. An early response to 8-oxoG lesions involves ‘on-the-fly’ translesion synthesis (TLS), in which a specialized TLS Pol is recruited and replaces the stalled HiFi Pol for lesion bypass. The length of TLS must be long enough for effective bypass, but it must also be regulated to minimize replication errors by the TLS Pol. The exact position where the TLS Pol ends and the HiFi Pol resumes (i.e. the length of the TLS patch) has not been described. We use steady-state and pre-steady-state kinetic assays to characterize lesion bypass intermediates formed by different archaeal polymerase holoenzyme complexes that include PCNA123 and RFC. After bypass of 8-oxoG by TLS PolY, products accumulate at the template position three base pairs beyond the lesion. PolY is catalytically poor for subsequent extension from this +3 position beyond 8-oxoG, but this inefficiency is overcome by rapid extension of HiFi PolB1. The reciprocation of Pol activities at this intermediate indicates a defined position where TLS Pol extension is limited and where the DNA substrate is handed back to the HiFi Pol after bypass of 8-oxoG.

INTRODUCTION
Accurate replication of genomic DNA is an elaborate cellular process which is essential to the propagation of life. The double helix is first unwound to form a replication fork where the two strands are primed for replication. A high-fidelity (HiFi) DNA polymerase (Pol) then associates with a processivity clamp to form a replicative holoenzyme (HE) complex which utilizes the single strands of the DNA substrate as a template for accurate duplication of the genome. At times, the molecular information of the template DNA is exposed to damaging agents and becomes chemically modified to form a variety of DNA lesions. DNA repair pathways (including nucleotide-excision repair and base-excision repair) allow for correction of these genomic lesions, yet, excess exposure to DNA damaging agents allows lesions to persist and then be encountered during replication. These template lesions cause HiFi Pols to stall, leading to genomic instability.

However, a specialized set of evolutionarily conserved Y-family Pols can perform translesion synthesis (TLS) by incorporating deoxynucleotides (dNTPs) across template lesions (1). Although these TLS Pols are prone to errors during replication (2), they provide a pathway for DNA damage tolerance and allow replication to proceed in the presence of unrepaired DNA lesions. Previous studies have im-
licated two DNA damage tolerance pathways which involve TLS Pols: by ‘re-priming’, a new primer is synthesized downstream of the replication-stalling lesion (3–5). This allows the HiFi Pol to skip the lesion and continue synthesis downstream. However, this discontinuous process creates a single-stranded gap which must be resolved by either template-switching homology-directed gap repair (a TLS-independent pathway) or post-replicative gap filling by TLS Pols (6). Alternatively, TLS may occur ‘on-the-fly’, in which a TLS Pol is recruited directly to a stalled replication fork to perform lesion bypass prior to a re-priming event (7). Recent in vitro studies have suggested that this continuous ‘on-the-fly’ TLS pathway is an early response to replication stalling lesions, whereas re-priming serves as a slower mechanism which is likely utilized during conditions of excessive DNA damage (8–10).

The model for ‘on-the-fly’ TLS presents two distinct events where the DNA substrate is ‘handed-off’ between the HiFi and TLS Pols. Upon encountering a lesion, a HiFi Pol stalls one template base prior to the lesion (−1) as a result of competing polymerase and exonuclease activity (11). From this stalled position, the DNA must be handed-off from the stalled HiFi Pol to a TLS Pol, allowing for bypass of the template lesion. The mechanism of this hand-off may occur indirectly by distributive ‘exchange,’ where an active Pol fully dissociates from the DNA prior to equilibrium-driven recruitment of the next Pol from solution (12); or, the DNA may be passed directly through a ‘switch’ within a coupled suprapholoenzyme (SHE) complex consisting of multiple Pols bound to a single processivity clamp (i.e. the tool belt model) (13–16). The TLS Pol synthesizes a nascent segment of DNA which bypasses the lesion and extends to an intermediate position downstream. However, the exact length of this TLS patch and the position of the second hand-off (to resume synthesis by the HiFi Pol) has not been investigated. It is understood that the length of the TLS patch must be long enough to prevent lesion recognition and exonucleolytic degradation by the HiFi Pol, but continued extension beyond the lesion by low-fidelity TLS Pols must be limited to minimize error-prone synthesis (17). Limitations to TLS Pol extension may indeed be explained by their characteristically low processivity (18), but the complex multieukaryotic cell provides multiple factors which may regulate the TLS Pol. A previous study demonstrated that a TLS patch length of five nucleotides beyond an acetylaminofluorine adduct was sufficient for HiFi Escherichia coli Pol III to resume replication; however this is not implicated as the position of the second hand-off, as TLS Pol V is capable of much longer processive stretches of synthesis (19). In eukaryotes, synthesis of this TLS patch is more intricate, involving posttranslational modifications of PCNA (20) and additional hand-offs among the various TLS inserter and extender Pols which possess unique activities for inserting across and extending beyond different types of DNA lesions, respectively (21). This is further complicated by a recent study revealing competition between Polδ and Polη in a stalled replisome that is decoupled from the CMG, even on a leading-strand lesion (10). Nevertheless, the extent of combined TLS Pol activity beyond the lesion is not fully understood. This current knowledge gap encourages additional investigation into the molecular mechanisms which limit error-prone TLS extension after lesion bypass and restore accurate HiFi replication.

PolY (also referred to as Dpo4) acts as the TLS Pol in the archaeon *Saccharolobus solfataricus* (Sso) (22). This model enzyme has been well studied structurally (23–26) and kinetically (24,27,28), and its legacy has provided a basis for our current understanding of the various eukaryotic TLS Pols (29). PolY adopts the canonical ‘right-handed’ Pol structure with ‘Fingers’, ‘Palm’ and ‘Thumb’ subdomains (23,30). As a Y-family polymerase, PolY also contains the unique Little-Finger (LF) domain which plays a central role in TLS activity (23,31). In response to 8-oxoguanine (8-oxoG) lesions, PolY is remarkably efficient and accurate for non-mutagenic insertion of dCTP (24,28). Other Pols are prone to incorrect insertion of dATP across 8-oxoG by formation of a Hoogsteen base pair, resulting in G-to-T transversion mutations (32). However, R332 (in the LF of PolY) stabilizes the *anti*-conformation of 8-oxoG in the PolY active site, allowing accurate Watson-Crick base pairing and efficient insertion of dCTP (24,25,33).

Downstream of 8-oxoG, PolY is capable of extension activity (24,28). However, the extent of PolY’s catalytic role beyond the lesion in the presence of other replisome components has not been investigated fully. This prompts examination into how these ‘right-handed’ Pols ‘hand-off’ the DNA in response to lesions. One previous kinetic study included PolB1 (the HiFi Pol in *Sso*) to evaluate its role in extension after PolY bypass of 8-oxoG (28), albeit in the absence of the heterotrimeric processivity clamp PCNA123 (34,35) and clamp loader complexRFC (36,37). PolB1 polymerase and exonuclease activities were unperturbed by the template 8-oxoG lesion on a DNA substrate with a primer terminus that was extended 1 base pair downstream (+1) of the lesion. This +1 position is highlighted as a kinetically favorable location for the second Pol hand-off. However, the authors ultimately conclude that inclusion of PCNA123 may alter the balance of bypass and extension activities by PolY and PolB1, and this mediation may dictate the position where the DNA is handed off between Pols after bypass of 8-oxoG (28).

Building upon this kinetic framework, we have determined that PolB1 hands-off the template to PolY to perform lesion bypass across 8-oxoG and extends to a position three base pairs beyond the lesion (+3). Additional extension by PolY from this position is inefficient, leading to accumulation of a +3 intermediate product. Within a SHE complex (composed of PolB1, PolY and PCNA123), accumulation of the +3 intermediate is reduced as a result of a second hand-off to resume rapid extension activity of PolB1. Collectively, these findings denote a specific position for the second Pol hand-off where HiFi replication resumes after TLS bypass of an 8-oxoG lesion during ‘on-the-fly’ TLS.

**MATERIALS AND METHODS**

**Protein expression and purification**

Wild type *Sso* PolB1 (38), PolY (13) and RFC (39), were expressed and purified as previously described. Wild-type PolB1 was used in these studies, as exonuclease deficient PolB1 has been shown to bypass 8-oxoG lesions (28). His-tagged subunits of heterotrimeric PCNA123 were expressed...
and purified individually and reconstituted into the heterotrimer complex as previously described (39).

The DNA sequence for PolB1 binding protein 1 (PBP1; SSO0150) and PBP2 (SSO6202) were amplified from _Sso_ genomic DNA (primer sequences in Supplemental Table S1). The amplified genes were cloned into pET30a using XhoI and NdeI restriction sites. The proteins were expressed in Rosetta2 DE3 _E. coli_ cells by autoinduction (40), followed by lysis and purification of the PBPs as previously described (41).

**Steady-state lesion bypass and extension assays**

DNA primers (Sigma) were radiolabeled at the 5′-end with 32P using the manufacturer recommended protocol for T4 PNK (New England Biolabs) and ATP [γ-32P] (Perkin Elmer). Radiolabeled primers were individually mixed with the DNA template containing an 8-oxoG lesion at the 23rd base from the 3′ end (t52_s, Figure 1A and Supplemental Table S1). Substrates were annealed at a 1:2:1 primer:template ratio in a thermocycler (BioRad) by incubation for 10 minutes at 95°C, followed by a decrease to 20°C at a rate of 1°C/min. Annealed substrates were pre-loaded with different Pol HE complexes in reaction buffer (20 mM Tris-OAc (pH 7.5), 100 mM KOAc) with PCNA123, RFC, PolB1 and/or PolY. ATP was added, followed by incubation at 60°C for 5 min to promote loading of PCNA123 to the DNA substrate. Reactions were initiated upon addition of dNTPs, Mg(OAc)2, and salmon sperm DNA (spDNA; Invitrogen) as an enzyme trap, followed by quenching with 100 mM EDTA after 3 min. The final reaction concentration of each component is indicated in Figure 1A.

Reaction products were resolved by denaturing polyacrylamide gel electrophoresis (PAGE) through either a 10 cm gel (20% polyacrylamide/6M urea/25% formamide/1x TBE) or a 38 cm DNA sequencing gel (15% polyacrylamide/6M urea/25% formamide/1x TBE). Resolved products were exposed to a phosphor screen for a minimum of 4 h and scanned with a Storm 820 phosphorimager (GE Healthcare Life Sciences). The product signals were quantified using ImageQuant software (v5.0, GE Healthcare Life Sciences).

**Pre-steady-state lesion bypass and extension kinetics assays**

For pre-steady-state lesion bypass kinetics assays, reactions were performed using a DNA substrate containing the 8-oxoG damaged (t52_s) or undamaged (t52_Und) DNA template annealed to primer p26 (+3). Where indicated, single nucleotide incorporation assays were performed by initiating extension reactions with various concentrations of dATP. PolY and/or PolB1 were introduced to the extension kinetics assays from different syringe positions as outlined in pre-loading Schemes i–iv. Products were resolved by denaturing PAGE, exposed, scanned, and quantified for total extension activity as above.

**Fluorescence anisotropy DNA binding assay**

HPLC purified and fluorescently labeled primers (p22-FAM or p26-FAM) (Sigma-Aldrich) were annealed to either damaged or undamaged templates (t52_s and t52_Und; Supplemental Table S1). Fluorescence anisotropy experiments were performed essentially as described previously (38). Fluorescence was excited at 483 nm, and the emission at 517 nm was monitored during a 2 s integration and an average of three measurements at room temperature using a Cary Eclipse (Agilent) fluorimeter with automatic polarizers. Slit widths were set at 10 nm, and the PMT was 700 V. The G-factors (G) were automatically calculated and corrected, and anisotropy (r) was calculated according to Equation (2) using the included Agilent software:

\[
r = \frac{I_{VV} - G I_{VH}}{I_{VV} + 2 I_{VH}}
\]

where \(I_{VV}\) and \(I_{VH}\) are the intensity of the vertically or horizontally polarized light, respectively. PolY was then titrated into the indicated DNA substrates (4 nM), mixed, and incubated for 90 seconds prior to measuring anisotropy values. Anisotropy values were plotted as a function of PolY concentration to obtain the binding curve and fit to Equation (3) to obtain the dissociation constant, \(K_d\):

\[
\text{Fraction Bound} = \frac{B_0 + (B_{max} * [\text{PolY}]^{n_H})}{K_d^{n_S} + [\text{PolY}]^{n_S}}
\]

where \(B_0\) is the fraction bound at 0 nM PolY, \(B_{max}\) is the fraction bound at saturation, \(n_S\) is the Hill slope, and \(K_d\) is the dissociation constant for PolY binding to DNA. Reported \(K_d\) values are the average of three independent titrations.
RESULTS

PolY and PolB1 have complementing activities for bypass of and extension from an 8-oxoG lesion

To measure lesion bypass activity, different Pol HE complexes were assembled by pre-loading PolB1 and/or PolY with PCNA123 on a damaged DNA substrate containing an 8-oxoG lesion. Primers of progressing length were utilized in order to monitor lesion bypass or extension activities from different starting positions relative to the site of the template lesion (Figure 1A). Reactions were initiated upon addition of dNTPs, Mg²⁺, and an excess amount of non-specific salmon sperm DNA (spDNA) as a trap to alter the multiequilibria of polymerase ‘exchange.’ As a pre-trapping control, spDNA was pre-incubated with the DNA substrate prior to reaction initiation (Supplemental Figure S1). At 1 mg/ml of spDNA, pre-trapped Pol complexes are not completely trapped in solution, suggesting that this concentration of spDNA does not abolish but slows the rate of distributive Pol exchange from solution. On the running start primer p18 (–5), synthesis takes place prior to encountering the 8-oxoG lesion. However, to compare lesion bypass activity, only the signal of ‘Bypassed’ products beyond the position of the lesion were quantified (Figure 1B, brackets). From the running start position, the PolB1 holoenzyme (B1HE; lanes 1–3) is completely stalled at the 8-oxoG lesion with negligible signal beyond the lesion (purple). As expected, reactions with the PolY holoenzyme (YHE, lanes 4–6) demonstrate lesion bypass activity with 39.8 ± 1.5% of products extended beyond the lesion (red). Upon examining the gel image, the distribution of lesion bypass products by the YHE appear to be enriched at an intermediate position downstream of the lesion (Figure 1B, arrow). This indicates that PolY is capable of TLS, but it exhibits poor processivity and limited extension beyond the 8-oxoG lesion. Lastly, formation of the SHE complex, containing both PolB1 and PolY (lanes 7–9), yields significantly more (52.6 ± 3.3%; blue) lesion bypass products compared to the YHE. Additionally, the intermediate observed in the YHE reactions is not apparent in the lesion bypass products for the SHE, and more products are elongated to FL.

We repeated this lesion bypass assay on a DNA substrate simulating a stalled position using primer p22 (–1; Figure 1C). Again, the B1HE (lanes 1–3) is stalled by 8-oxoG (purple), while the YHE (lanes 4–6) is capable of lesion bypass (55.0 ± 0.5%, red). As observed in Figure 1B, most of the lesion bypass products by the YHE are concentrated at a position beyond the lesion (Figure 1C, arrow). Finally, though the SHE (lanes 7–8) does not yield a significant increase in total bypassed products (64.3 ± 6.0%, blue) relative to the YHE, most of the bypassed products are extended to the end of the substrate and the observed intermediate disappears. Thus, in response to 8-oxoG, bypass and extension from the lesion are most efficient when both PolB1 and PolY are present.

We continued these lesion bypass assays for each Pol complex using DNA substrates with progressively longer primers and quantified the percentage of primer that was utilized for each substrate (i.e. extended by at least one base pair; Figure 1D, gel images in Supplemental Figure S2A–D). In this way, we can compare the activities of each Pol complex on substrates with different starting positions relative to the 8-oxoG lesion. Activity of the B1HE is negligible on primers p22 and p23, which start before (–1) or at (0) the site of the lesion. However, activity of the B1HE gradually increases on primers further downstream of the lesion (purple circles). Remarkably, products from the YHE exhibit the opposite trend; activity is elevated on primers before or at the lesion and gradually decreases on primers which require extension from the lesion (red squares). Finally, activity of
the SHE consistently yields about 65% of extension on all primers (blue diamonds). Unlike reactions with individual Pol HEs, the magnitude of primer extension by the SHE complex is independent of starting primer length.

The apparent inverse in activities relative to the 8-oxoG position by the YHE and B1HE highlights the complementing roles for PolY and PolB1. This is further supported by the SHE reactions, where the combined Pol activities yield increased levels of bypass and extension. These data also point to a specific position where the substrate is handed off between the Pols in the SHE reactions. The reciprocation of Pol activities occurs on a primer terminus positioned 3 bp downstream of the lesion. This leads us to hypothesize that the +3 position is the site of the second hand-off, from TLS PolY to HiFi PolB1, after bypass of an 8-oxoG lesion.

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Lesion bypass by PolY yields a major intermediate 3 base pairs downstream of the 8-oxoG lesion

In order to visualize discrete lesion bypass intermediates, representative products from the previous reactions were resolved by high-resolution denaturing PAGE (Figure 2A). This gel image depicts the same activity trends summarized in Figure 1D, but it also grants a more precise examination of lesion bypass and extension products formed by each Pol complex. Notably, lesion bypass products by the YHE are specifically enriched at the +3 position, regardless of the primer starting position (Figure 2A, arrow). The consistent position of this enriched YHE intermediate indicates that it is not a product of an ordinary processive step by PolY. If so, the position of the enriched intermediate would shift along the substrate in relation to the primer starting length. This observation indicates a specific perturbation in polymerase activity of PolY when positioned 3 base pairs downstream of an 8-oxoG lesion.

As previously noted, lesion bypass with the SHE yields an apparent decrease in the intermediate observed in bypass by the YHE complexes and an apparent increase in products extended to FL. In order to compare the relative amount of +3 intermediate formed on each substrate, the signal of the +3 intermediate product bands (Figure 2A, arrow) from the YHE and SHE reactions were normalized to their respective lanes (Figure 2B). When compared, the magnitude of the +3 intermediate signals from the SHE reactions (blue) is reduced by approximately half relative to the YHE reactions (red) on corresponding DNA substrates. Additionally, upon comparing the normalized fraction of extended products (Figure 2A, bracket), the SHE reactions (blue) yield about twice as much product that is extended beyond the +3 position relative to the YHE (red; Figure 2C). Altogether, these data indicate that PolY conducts bypass of an 8-oxoG lesion and is disrupted at a position 3 base pairs downstream, and the presence of PolB1 promotes extension from this intermediate position.

Kinetics of lesion bypass intermediates demonstrate reduced accumulation of the +3 intermediate upon formation of the SHE

After recognizing the +3 position as a major intermediate for PolY, we examined the direct kinetics of the intermediate without altering the equilibria by spDNA. In order to acquire faster time points, reactions were operated through an RQF-3 (KinTek) rapid quench instrument at 40°C. PolY was introduced to the reaction by either syringe A or B (Figure 3A, Scheme i or ii). By changing the position where PolY is introduced to the reaction, we can evaluate how the observed rate of lesion bypass changes under different pre-loading conditions. Any products which extend from this primer were quantified as ‘Total Bypass’ (Figure 3B, left bracket), as the 8-oxoG lesion is the first templating base from the stalled primer p22 (−1) terminus. Further, all extension products were classified into three categories of intermediates: ‘0 to +2’, ‘+3’ and ‘Extended.’ (Figure 3B, right brackets). The signal of each intermediate category was then quantified and fitted to observe these intermediate signals over time.

In reactions where PolY is pre-loaded with the DNA substrate and accessory factors (Figure 3C, Scheme i), total bypass proceeds with a $k_{obs}$ of 0.81 ± 0.11 s$^{-1}$. This rate is representative of the catalytic rate of pre-loaded PolY across 8-oxoG. When PolY must first be recruited to the stalled DNA substrate upon initiation (Figure 3C, Scheme ii), the apparent rate of bypass is slowed 4-fold, with a $k_{obs}$ of 0.22 ± 0.02 s$^{-1}$. This observed rate is likely limited by the rate of association of PolY to DNA; PCNA123 prior to the catalytic step. In these YHE schemes, the kinetic trace of the +3 intermediate accumulates (Figure 3C, dashed traces) and reaches a maximal normalized product of over 0.20 (at 10 s), suggesting that extension by the YHE complexes is impaired at the
+3 position. When lesion bypass reactions are performed by Scheme i using abasic and undamaged DNA templates, this intermediate is not apparent (Supplemental Figure S3A). Also, on an alternative 8-oxoG template containing a different sequence context beyond the lesion (8TAC vs 8CGC), the YHE still yields an enriched +3 intermediate after 10 seconds (Supplemental Figure S3B). Taken together, these results indicate that the +3 intermediate is characteristic of PolY bypass of 8-oxoG and is not an artifact of sequence context.

Considering the disappearance of the +3 intermediate and increased extension observed in the steady-state SHE reactions (Figure 2B and C), it follows that addition of PolB1 into the kinetic lesion bypass assay would reduce accumulation of the +3 intermediate. To examine this, PolB1 was introduced to the pre-steady-state assay schemes through syringe B or A, opposite the location of PolY (Figure 4A, Scheme iii, or Figure 4B, Scheme iv), then initiated and quenched as before. When PolY is pre-loaded in syringe A, and PolB1 is introduced upon initiation in syringe B (Figure 4A), lesion bypass proceeds at a similar rate ($k_{obs} = 0.72 \pm 0.08 \text{ s}^{-1}$) compared to when PolY is pre-loaded alone (Figure 3C, Scheme i). Schemes i & iii are both initiated with a pre-loaded PolY, thus quantification of 'Total Bypass' by these schemes is representative of the catalytic step of lesion bypass by PolY. However, in response to DNA lesions, the TLS Pol must access the primer terminus after stalling of the HiFi Pol. To investigate this step, pre-steady-state assays were performed by Scheme iv, which simulates a
stalled B1HE that has encountered an 8-oxoG lesion. Wild-type PolB1 is completely stalled by the 8-oxoG lesion (Figure 1C), so ‘Total Bypass’ products by this scheme require catalysis by PolY. Therefore, in Scheme iv, the observed rate of synthesis would be limited by rate of the first hand-off from PolB1 to PolY. Indeed, the observed rate of lesion bypass by Scheme iv ($k_{obs} = 0.077 \pm 0.009 \text{ s}^{-1}$) is about 10-fold slower than Schemes i & iii, which contain a pre-loaded PolY. This rate is also 3-fold slower than the rate of lesion bypass when PolY is recruited to an unoccupied clamp (Figure 4B, Scheme iv versus ii), indicating that the presence of a pre-loaded PolB1 slows PolY access to the stalled primer terminus. Observed rates of lesion bypass for Schemes i–iv are compared in Figure 4C.

Recently, two novel accessory subunit proteins, PBP1 and PBP2, were characterized and shown to influence the activity of PolB1 (41). PBP1 contains an acidic tail which limits strand displacement activity by PolB1 during synthesis of Okazaki fragments, whereas PBP2 moderately improves catalytic activity. Given their effects on PolB1, these PBPs were included in pre-steady-state kinetic assays on undamaged DNA templates to determine whether they have an effect in regulating or displacing PolB1 upon encountering a lesion. When both PBP1 and PBP2 (400 nM) are pre-loaded with PolY in Scheme i, there is no apparent change in the rate of PolY catalysis (Supplemental Figure S4A). This suggests that the PBPs have no observable effect on PolY activity. Similar kinetic assays were performed using PolB1 in place of PolY (later referred to as Scheme v in Figure 6B). By this scheme, reactions containing PBP1 demonstrate an inhibitory effect on the rate of catalysis by PolB1, while PBP2 alone improves PolB1 activity, as previously noted (Supplemental Figure S4B).

From its apparent inhibition of PolB1 activity, we hypothesized that PBP1 may destabilize PolB1 affinity to DNA. Upon PolB1 encountering a template lesion, PBP1-mediated displacement of PolB1 would allow faster access of PolY to the stalled primer. To investigate this, lesion bypass kinetic assays were performed by Scheme iv in the presence of PBP1 pre-loaded with PolB1 in syringe A. However, this did not yield any apparent change in the observed rate of exchange from PolB1 to PolY for lesion bypass (Supplemental Figure S4C and D). The similarity in these rates suggests that PBP1 does not regulate PolB1 activity by promoting its displacement from DNA, and these accessory subunits are not directly involved in the Pol hand-off during ‘on-the-fly’ TLS.

To compare the accumulation of the +3 intermediate in all lesion bypass kinetics schemes (Figures 3C, 4A and B, dashed traces), we plotted the maximal signal of the +3 intermediate in the kinetic traces from each Pol complex (Figure 4D). In schemes containing PolB1 (iii & iv), the maximal fraction of the +3 intermediate (0.11 ± 0.004 and 0.04 ± 0.01, respectively) is significantly less than the intermediate signal from reactions containing PolY alone (Schemes i & ii; 0.26 ± 0.03 and 0.23 ± 0.03, respectively). Despite the slower overall rate of lesion bypass in Scheme iv, the reduced fraction of +3 intermediate indicates that extension from the intermediate position is still fast. Again, this shows that PolY is capable of rapid lesion bypass up to the +3 position, but additional extension is impaired and results in accumulation of the +3 intermediate which is relieved by the presence of PolB1.

**Slow extension by the YHE complexes from the +3 intermediate indicates a perturbation in PolY catalytic activity at this position**

The peculiar features of this +3 lesion bypass intermediate prompted us to investigate the kinetics of extension from this position by the different Pol complexes. Reactions were assembled in the RQF-3 as described above, except a DNA substrate containing primer p26 (+3), which corresponds with the observed intermediate, was utilized. All products beyond the primer were quantified as ‘Extended’ products (Figure 5A). For reactions containing PolY alone (Schemes i & ii), the apparent rates of extension are equivalent for Schemes i & ii, regardless of PolY syringe position ($k_{obs} = 0.080 \pm 0.008 \text{ s}^{-1}$ and $0.078 \pm 0.008 \text{ s}^{-1}$, respectively). The similar rates from these pre-loading schemes suggest that extension by PolY from the +3 position is not limited by its rate of association to the DNA substrate. This contrasts with the observed rates obtained for lesion bypass for Schemes i & ii (Figure 3C). These YHE extension rates are about 2.8-fold slower than the apparent rate of associ-
atation for PolY (represented by lesion bypass in Scheme ii), indicating that the rate of PolY catalysis is limiting when extending from the +3 intermediate position after 8-oxoG. We considered the reduced rate of extension to be limited by a decrease in DNA binding affinity of PolY to the +3 extension substrate. This circumstance would effectively reduce the formation of active YHE complexes and yield a slower apparent rate of extension. However, as determined by fluorescence anisotropy, there was no apparent change in PolY binding affinity for the 8-oxoG substrate relative to the undamaged substrate with primers aligned at the –1 or +3 positions (Supplemental Figure S5). This suggests that slow extension of the YHE from the +3 intermediate is not a result of destabilized DNA binding, rather it is limited by a slow elementary step which follows PolY association with the DNA substrate.

To investigate the kinetic parameter of PolY that is affected at this position, we measured the observed rates of single nucleotide incorporation by Scheme i as a function of dATP concentration, the next nucleotide for extension from the +3 position. On an undamaged DNA template, $k_{obs}$ increases with elevated [dATP], as expected (Figure 5B). However, the rate of dATP incorporation from this position on the 8-oxoG damaged template is significantly slower (Figure 5C). The second-order rate constant of extension from the +3 position is 2.1 ± 0.3 s$^{-1}$. The rate of dATP incorporation for the two nucleotides past bypass of 8-oxoG (Supplemental Figure S5) increases with elevated [dATP], as expected (Figure 5B). However, the rate of dATP incorporation from this position on the 8-oxoG damaged template is significantly slower (Figure 5C). The second-order rate constant of extension from the +3 position is 2.1 ± 0.3 s$^{-1}$ relative to 0.43 ± 0.01 nM$^{-1}$s$^{-1}$, indicating a perturbation of PolY extension three nucleotides past bypass of 8-oxoG.

PolB1 performs rapid extension from the +3 intermediate position

As observed in Figure 1D, PolB1 extension activity is highest when starting from the +3 position (purple circles), indicating kinetic efficiency for extension by PolB1. In order to investigate the role of PolB1 in extension from this intermediate position, we evaluated the kinetics of extension from the +3 position by Schemes iii & iv. Extension by Scheme iii simulates a condition that examines a switch to PolB1 after PolY has bypassed an 8-oxoG lesion, which results from a perturbation in polymerase activity at that position. In the presence of PolB1, accumulation of the +3 intermediate is reduced and rapidly extended into longer products. Altogether, this points to an ‘on-the-fly’ TLS mechanism where the first hand-off (from stalled PolB1 to PolY) occurs at the -1 position (Figure 7A to B). PolY then performs TLS and extends to a position 3 base pairs beyond the lesion, where it becomes catalytically inefficient for extension to the +4 position (Figure 7C). From this +3 position, the catalytic inefficiency PolY allows for the second hand-off to restore PolB1 activity (Figure 7D to E). This marks a distinct position beyond an 8-oxoG lesion for the second Pol hand-off which resumes HiFi replication and effectively limits error-prone TLS activity.

Previous kinetic and structural studies of PolY bypass and extension across 8-oxoG did not identify the +3 intermediate (24,25,28). Accumulation of this intermediate was first apparent to us upon inclusion of spDNA trap in steady-state lesion bypass assays, which effectively amplifies sites of catalytic inefficiency during replication. In prior studies without trap DNA, multiple enzyme turnovers and distributive extension from the +3 position may have proceeded too quickly for its accumulation to be apparent. These studies also primarily focus on the kinetic steps of lesion bypass, translocation, and extension up to the +1 position by PolY; its activity was not evaluated further downstream of 8-oxoG. Finally, prior evaluations were also performed without a processivity clamp, which is a critical accessory factor that optimizes and mediates Pol and other enzymatic activities (39,42,43). Our study suggests that PCNA123 mediates the complementing activities of PolY and PolB1 not only upon encountering the lesion, but also after the lesion has been bypassed. In response to other DNA lesions, the kinetics of lesion bypass by PolY have already been characterized (44–49). In some of the previous analyses, ‘pause sites’ were noted during PolY lesion bypass, indicating points of inefficient catalysis and extension from these lesions. However, in the absence of RFC, PCNA123 and PolB1, these pause sites cannot be directly correlated with the position of the second hand-off after lesion bypass. This prompts additional investigation into how PolY and other TLS Pols are limited in extension after bypass of various DNA lesions, especially in the presence of a processivity clamp and HiFi Pols.

DISCUSSION

In this study, we have identified a novel +3 intermediate for PolY after bypass of an 8-oxoG lesion, which results from a perturbation in polymerase activity at that position. In the presence of PolB1, accumulation of the +3 intermediate is reduced and rapidly extended into longer products. Altogether, this points to an ‘on-the-fly’ TLS mechanism where the first hand-off (from stalled PolB1 to PolY) occurs at the -1 position (Figure 7A to B). PolY then performs TLS and extends to a position 3 base pairs beyond the lesion, where it becomes catalytically inefficient for extension to the +4 position (Figure 7C). From this +3 position, the catalytic inefficiency PolY allows for the second hand-off to restore PolB1 activity (Figure 7D to E). This marks a distinct position beyond an 8-oxoG lesion for the second Pol hand-off which resumes HiFi replication and effectively limits error-prone TLS activity.

Previous kinetic and structural studies of PolY bypass and extension across 8-oxoG did not identify the +3 intermediate (24,25,28). Accumulation of this intermediate was first apparent to us upon inclusion of spDNA trap in steady-state lesion bypass assays, which effectively amplifies sites of catalytic inefficiency during replication. In prior studies without trap DNA, multiple enzyme turnovers and distributive extension from the +3 position may have proceeded too quickly for its accumulation to be apparent. These studies also primarily focus on the kinetic steps of lesion bypass, translocation, and extension up to the +1 position by PolY; its activity was not evaluated further downstream of 8-oxoG. Finally, prior evaluations were also performed without a processivity clamp, which is a critical accessory factor that optimizes and mediates Pol and other enzymatic activities (39,42,43). Our study suggests that PCNA123 mediates the complementing activities of PolY and PolB1 not only upon encountering the lesion, but also after the lesion has been bypassed. In response to other DNA lesions, the kinetics of lesion bypass by PolY have already been characterized (44–49). In some of the previous analyses, ‘pause sites’ were noted during PolY lesion bypass, indicating points of inefficient catalysis and extension from these lesions. However, in the absence of RFC, PCNA123 and PolB1, these pause sites cannot be directly correlated with the position of the second hand-off after lesion bypass. This prompts additional investigation into how PolY and other TLS Pols are limited in extension after bypass of various DNA lesions, especially in the presence of a processivity clamp and HiFi Pols.
Figure 6. PolB1 conducts rapid extension from the +3 intermediate position. Pre-steady-state extension kinetics were performed on a DNA substrate containing primer p26 (+3) by (A) Schemes iii (yellow, •→) and iv (green, •▲) containing PolB1 and PolY, or (B) Schemes v (blue, •→) and vi (purple, •▲) containing PolB1 alone. Error bars represent standard deviation from three independent replicates of each time point. Quantification of 'Extended' products were fit to Equation (1) to obtain the observed rate constant for extension. Insets show the trace of 'Extended' product at a shorter time scale. (C) Observed rate constants for extension from +3 were plotted; error bars indicate error of the regression fit.

Figure 7. Proposed mechanism for DNA hand-offs between PolB1 and PolY in response to an 8-oxoG lesion. (A) Upon encountering 8-oxoG on the template strand, PolB1 stalls. (B) Individual contacts with PCNA123 mediate the first Pol hand-off from PolB1 to PolY. (C) PolY then performs translesion synthesis to the template position three base pairs beyond the 8-oxoG lesion, where it becomes catalytically inefficient for additional synthesis. (D) The second Pol hand-off from PolY to PolB1 is then mediated through PCNA123 to re-establish HiFi PolB1 for (E) extension and effective bypass of the lesion. The shaded step highlights the second transient polymerase handoff to restore high fidelity synthesis.

Interestingly, E. coli Pol IV (or DinB, a homolog of PolY) (22) demonstrates a similar +3 intermediate following bypass of N2-dG-peptide adducts (50). A Pol IV variant led to identification of a +3 intermediate in response to an N2-furfuryl-dG lesion, with a drastic reduction in catalytic efficiency from the intermediate position (51). Additionally, human Polη was observed to become less processive at template positions one or two base pairs beyond a thymine-thymine dimer (52,53). These studies suggest the potential for structural impediments, or communication, between a bypassed template lesion and the TLS Pol active site once it has translocated to a certain position downstream, but the basis of this disruption has not been explored. We hypothesize that these observed inefficiencies correspond with a disruption in translocation. A series of positively charged residues of the PolY LF domain interact with the DNA backbone and are implicated in facilitating PolY translocation (25). During extension from a bypassed lesion, an altered base with modifications that are oriented toward the LF may sterically disrupt interactions between template DNA and the residues involved in translocation. A disruption in translocation would prevent the Pol active site from accessing the next template base and interrupt binding of an incoming nucleotide, ultimately weakening the apparent efficiency of successive catalytic extensions. The apparent conservation of the +3 intermediate for TLS Pols from E. coli and Sso and similar intermediates observed in humans strengthens the hypothesis that the LF domain can sense bypassed lesions after translocating to a certain position downstream. Activation of this ‘pinky trigger,’ along with complementation of extension activity by a HiFi Pol and mediation through a processivity clamp may serve as the molecular switch which restores HiFi Pol activity. Though the various TLS Pols have been characterized in their bypass of a diverse set of lesions, further investigation is needed to understand the mechanisms which limit the extent of their error-prone activities.

Previously, under conditions similar to the current study, we measured the rate of association of PolY to a pre-loaded B1HE using pre-steady-state FRET (13). This apparent rate
(0.8 ± 0.1 s⁻¹) is over 10-fold faster than the apparent rate of lesion bypass by the same pre-loading approach evaluated in Scheme iv (0.077 ± 0.009 s⁻¹). In that same study, we also observed that PolB1 was not displaced upon PolY association within the measured timescale, indicating that both Pols were simultaneously bound to the clamp and formed a concerted PCNA tool belt (13). Combined with our current findings, this suggests that PolY may rapidly associate with the stalled BIHE to form a SHE complex, but the rate of the first Pol switch within the complex (involving transient dissociation of PolB1 and PolY binding to the stalled substrate) is slower and rate limiting for lesion bypass by PolY.

The apparent rate for the first Pol hand-off to PolY for TLS (0.077 ± 0.009 s⁻¹) is 6.4-fold slower than the second switch to PolB1 for extension (0.49 ± 0.05 s⁻¹; Figure 4A, Scheme iv versus Figure 6A, Scheme iii). This suggests that PolB1 is more stably bound to DNA relative to PolY, which is consistent with previous Pol:DNA binding affinity studies (54), yet these binding affinities have not been evaluated in the presence of PCNA123. The apparent difference in hand-off rates observed in this study indicates that the fundamental basis for each hand-off is molecularly unique, and additional investigation is also required to obtain direct insight into the kinetic and structural mechanisms of these Pol hand-off events.

After an ‘on-the-fly’ TLS event and resumption of HiFi Pol activity, the TLS Pol may remain bound to the clamp during replicative extension. This configuration allows TLS to be transiently available for subsequent encounters with DNA lesions. Yet, in the context of a more complex multi-equilibria system, dynamic exchange of HiFi Pols and other DNA processing enzymes has been observed (55–57). PCNA123 has also been shown to coordinate an alternate arrangement of enzymes, optimizing their combined activities for Okazaki fragment maturation (58). This indicates that an effective cellular response would require dynamic configurations of various cellular tools in order to faithfully maintain the genome. Therefore, the mechanisms of Pol hand-offs may include aspects of distributive Pol exchange. However, this does not eliminate the possibility of switching through direct contacts between Pols by way of a transient hand-off through a PCNA tool belt scaffold, especially considering the network of direct interactions among TLS Pols (59,60) and the sharing of accessory subunits between eukaryotic TLS Pol and HiFi Pol (61–63).

Characterizing the intricacies of ‘on-the-fly’ TLS and its hand-off mechanisms is integral to our understanding of this early DNA damage response. Though this damage tolerance pathway benefits the cell by relieving stalled replication, unregulated TLS activity can detrimentally contribute to mutagenesis and resistance to DNA-damaging cancer therapies. Recent advances have addressed these drawbacks through identification of small molecule inhibitors which target Pol–Pol interactions and/or disrupt mutagenic TLS activity, supplementing the effectiveness of current chemotherapeutic drugs (64–70). These studies provide a promising new approach for mitigating the role of TLS in chemoresistance and provide an exciting incentive for furthering our understanding of the kinetic and structural foundations of TLS.

### SUPPLEMENTAL DATA

Supplementary Data are available at NAR Online.

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### Conflict of interest statement

None declared.

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