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Kinetic Analysis of the Bypass of a Bulky DNA Lesion Catalyzed by Human Y-family DNA Polymerases

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Abstract

1-Nitropyrene (1-NP), a mutagen and potential carcinogen, is the most abundant nitro polyaromatic hydrocarbon in diesel exhaust, which reacts with DNA to form predominantly *N*- $(d_{\rm{coxygen}$ and -8 -yl)-1-aminopyrene (dG^{AP}) . If not repaired, this DNA lesion is presumably bypassed *in vivo* by any of human Y-family DNA polymerases kappa (hPolκ), iota (hPol*τ*), eta (hPol*η*), and Rev1 (hRev1). Our running start assays demonstrated that each of these enzymes was indeed capable of traversing a site-specifically placed dG^{AP} on a synthetic DNA template but hRev1 was stopped after lesion bypass. The time required to bypass 50% of the dG^{AP} sites (t_{50}^{bypass}) encountered by hPoln, hPolk and hPolt was determined to be 2.5 s, 4.1 s, and 106.5 s, respectively. The efficiency order of catalyzing translesion synthesis of dG^{AP} (hPoln > hPolk > hPol τ >> hRev1) is the same as the order for these human Y-family enzymes to elongate undamaged DNA. Although hPoln bypassed dG^{AP} efficiently, replication by both hPolk and hPolt was strongly stalled at the lesion site and at a site immediately downstream from dG^{AP} . By employing pre-steady state kinetic methods, a kinetic basis was established for polymerase pausing at these DNA template sites. Besides efficiency of bypass, the fidelity of those lowfidelity polymerases at these pause sites was also significantly decreased. Thus, if the translesion DNA synthesis of dGAP *in vivo* is catalyzed by a human Y-family DNA polymerase, *e.g.* hPolη, the process is certainly mutagenic.

INTRODUCTION

Both normal metabolic processes and environmental factors constantly damage cellular DNA and produce 1,000 to 1,000,000 DNA lesions per cell per day. If these DNA lesions are not repaired by cellular DNA repair machinery, they will stall replicative DNA polymerases and replication machinery.1-4 To rescue stalled DNA replication, cellular DNA repair machinery temporarily switches to a Y-family DNA polymerase that primarily

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¹Abbreviations: AAF-dG, *N*-acetyl-2-aminofluorene adduct at the C8 position of deoxyguanosine; 1-AP, 1-aminopyrene; BPDE-dG, benzocpyrene 7,8-diol 9,10-epoxide-derived adduct at the N2 position of 2′-deoxyguanosine; BSA, bovine serum albumin; Dpo4, *Sulfolobus solfataricus* DNA polymerase IV; dGAP, *N*-(deoxyguanosin-8-yl)-1-aminopyrene; dNTP, 2′-deoxynucleoside 5′ triphosphate; TLS, translesion DNA synthesis.

SUPPORTING INFORMATION AVAILABLE The structure of dG^{AP}, dG^{AP} bypass% plot, the pre-steady state kinetic parameters of each enzyme, a gel image of EMSA and corresponding plot, and the gel image displaying the efficiency of the DNA trap assay are available as supporting information. This material is available free of charge via the Internet at<http://pubs.acs.org>.

functions in the bypass of DNA lesions *in vivo*. Out of the 16 identified human DNA polymerases, four enzymes belong to the Y-family: DNA polymerases eta (hPolη), kappa (hPolκ), iota (hPolτ), and Rev1 (hRev1). These enzymes have been shown to catalyze both error-free and error-prone translesion DNA synthesis (TLS) *in vitro* and *in vivo*. 5-7 For example, hPolη catalyzes error-free bypass of *cis*-*syn* cyclobutane thymidine dimer *(cis*-syn TT ^{8, 9} and its inactivation by mutations leads to Xeroderma Pigmentosum Variant (XPV) disease that increases incidence of sunlight-induced skin cancer.8, 10 In addition, hPolη has been shown *in vitro* to bypass other lesions including apurinic/apyrimidinic (abasic) sites,¹¹⁻¹³ *N*-(deoxyguanosin-8-yl)-*N*-acetyl-2-aminofluorene (AAF-dG),¹³ and N^2 deoxyguanosine adducts of benzo[*a*]pyrene (BPDE-dG).¹² hPolk, a homolog of the wellstudied Y-family member *Sulfolobus solfataricus* DNA Polymerase IV (Dpo4), is capable of bypassing abasic sites,¹¹ AAF-dG, and BPDE-dG,¹⁴ and efficiently elongating mispaired primer termini.15 hPolτ has been shown to bypass abasic sites,¹¹ *cis*-*syn* TT,16 and AAF $dG¹⁷$ hRev1 preferentially incorporates dCTP opposite any template base¹⁸ and DNA lesions including abasic sites¹¹ and BPDE-dG.¹⁹ These biochemical studies demonstrate that there is a significant overlap in lesion bypass abilities of the four human Y-family members. At present, it is unclear which human Y-family enzyme bypasses which DNA lesion(s) *in vivo*.

Despite growing concerns about the effects of air pollution on human genomic stability, there are no comprehensive studies of the bypass of air pollutant-induced DNA adducts catalyzed by human Y-family DNA polymerases. The metabolites of 1-nitropyrene (1-NP), a product of incomplete gasoline combustion and one of the most abundant polycyclic aromatic hydrocarbons (PAH), $^{20-22}$ react with DNA to primarily form *N*-(deoxyguanosin-8yl)-1-aminopyrene (dGAP, Supplementary Figure 1).20 Because 1-NP is a potent mutagen and carcinogen in rodents,²³ it is classified as a class 2B carcinogen.^{24, 25} Moreover, the dG^{AP} lesion is shown to be mutagenic in bacterial and mammalian cells.^{24, 25} Thus, it is interesting to identify which human Y-family DNA polymerase(s) bypass(es) dGAP *in vitro* and *in vivo*. Previously, we have shown that the kinetics of nucleotide incorporation opposite dG^{AP} and the subsequent extension of the dG^{AP} bypass product catalyzed by Dpo4 are significantly altered.²⁶ A minimum kinetic mechanism for the dG^{AP} bypass has also been established via pre-steady-state kinetic methods.²⁶ Here, we employed these methods and kinetically assessed the dGAP bypass abilities of human Y-family DNA polymerases in order to shed light on which of these enzymes are better suited to bypass dGAP *in vivo*.

MATERIAL AND METHODS

Materials

Reagents were purchased from the following companies: OptiKinase from USB Corporation, [*γ* [−]32P]ATP from MP Biochemicals, and dNTPs from GE Healthcare. hPolη, hRev1, and hPolk were expressed and purified as previously described.¹¹ hPolt was expressed and purified as previously described¹¹ with one modification: the GST-tag was not removed in order to increase the protein stability of hPolτ.

DNA Substrates

The DNA template 26 -mer-d G^{AP} (Table 1) was synthesized and purified as previously described.26 Other DNA oligomers listed in Table 1 were purchased from Integrated DNA Technologies. The radiolabeling of the primers and the annealing of a primer and a template were performed as described previously.^{26, 27}

Buffers

All pre-steady-state kinetic assays were performed in reaction buffer R (50 mM HEPES, pH 7.5 at 37 °C, 5 mM MgCl₂, 50 mM NaCl, 0.1 mM EDTA, 5 mM DTT, 10 % glycerol, and 0.1 mg/ml BSA). All electrophoresis mobility shift assays (EMSA) were performed in buffer S (50 mM Tris-Cl, pH 7.5 at 23 °C, 5 mM MgCl₂, 50 mM NaCl, 5 mM DTT, 10 % glycerol, and 0.1 mg/ml BSA). All reported concentrations were final after mixing solutions.

Running Start Assays

The running start assay was performed similarly as described previously.^{11, 26, 28-30} Briefly, a preincubated solution of $5'-[3^2P]$ -labeled DNA (100 nM) and a human Y-family enzyme (1 μM) in buffer R was rapidly mixed with a solution containing all four dNTPs (200 μM each) at 37 °C via a rapid chemical-quench flow apparatus (KinTek). The reactions were quenched with 0.37 M EDTA after various times, and the reaction products were analyzed by denaturing polyacrylamide gel electrophoresis (PAGE, 20 % polyacrylamide, 8 M urea).

Quantitative analysis of the running start assays was performed by determining the relative lesion bypass efficiencies (dG^{AP} bypass%) as a function of reaction time. For each time point *t*, dGAP bypass% was calculated as the ratio of the bypass events to the encounter events (Equation 1):

$$
dG^{AP}bypass\% = (B/E) \times 100\% = {B / ([20 - mer] + B)} \times 100\%
$$

where the total dG^{AP} bypass events (B) was calculated from the concentration of all intermediate products with sizes greater than or equal to the 21-mer, and the total dG^{AP} "encounter" events (E) equaled the summation of the 20-mer concentration and the total dG^{AP} bypass events (B). To quantitatively define the dG^{AP} bypass efficiency, t_{50} ^{bypass} was defined as the time required to bypass 50% of the total dGAP lesions encountered.

Electrophoresis Mobility Shift Assays

hPolη (10-400 nM), hPolκ (65-950 nM), or hRev1 (15-550 nM) was titrated onto a solution containing $5'$ -[32P]-labeled DNA (10 nM) in buffer S at 23 °C. Native PAGE was used to separate the binary complex E•DNA from free DNA. After quantitation using a Typhoon Trio (GE Healthcare), the concentration of the binary complex was plotted as a function of the enzyme concentration. The data were fit to Equation 2 using Kaleidagraph (Synergy software):

$$
[E \cdot DNA] = 0.5(K_{d, DNA} + E_0 + D_0) - 0.5[(K_{d,DNA} + E_0 + D_0)^2 - 4E_0D_0]^{1/2}
$$
 (2)

where E_0 and D_0 are the initial enzyme and DNA concentrations, respectively, and K_d DNA is the equilibrium dissociation constant for E•DNA at 23 °C.

Active Site Titration Assays

A preincubated solution of 5′-[32P]-labeled DNA (10-450 nM) and hPolτ (27.5 nM, UVbased concentration) in buffer R was reacted with correct dNTP at saturating concentrations at 37 °C for various time intervals prior to be quenched by EDTA (0.37 M). The reaction products were resolved and quantitated as described above. The burst reaction amplitude (*A*) was plotted as a function of DNA concentrations and the data were then fit to Equation 3 to determine *Kd,* DNA.

(1)

 \overline{a}

$$
A=0.5\left(K_{d,\text{DNA}}+E_{o}+D_{o}\right)-0.5\left[\left(K_{d,\text{DNA}}+E_{o}+D_{o}\right)^{2}-4E_{o}D_{o}\right]^{1/2} \tag{3}
$$

Nucleotide Incorporation Efficiency and Fidelity Measurements

Single-turnover kinetic assays were employed to obtain the k_p (maximum dNTP) incorporation rate) and K_d , $dNTP$ (equilibrium dissociation constant for $dNTP$ from E•DNA•dNTP) for single nucleotide incorporation as described previously.^{26, 31} Briefly, a preincubated solution of $5'-[^{32}P]$ -labeled DNA (20 nM) and hPoln (130 nM) or hPol τ (130 nM) in buffer R was mixed with increasing concentrations of an incoming dNTP at 37 °C. For hPolk and hRev1, a preincubated solution of $5'-[3^2P]$ -labeled DNA (30 nM) and hPolk (300 nM) or hRev1 (120 nM) in buffer R was mixed with increasing concentrations of an incoming dNTP. The reactions were terminated after various times by the addition of 0.37 M EDTA. Reaction products were resolved and quantitated as described above. The time course of product formation at each dNTP concentration was fit to Equation 4:

$$
[Product] = A [1 - exp(-k_{obs}t)] \tag{4}
$$

where k_{obs} is the observed reaction rate constant. Next, the plot of the k_{obs} values as a function of dNTP concentrations was fit to Equation 5:

$$
k_{\text{obs}} = k_p \left[\text{dNTP} \right] / \left\{ \left[\text{dNTP} \right] + K_{d,\text{dNTP}} \right\} \tag{5}
$$

From this plot, the dNTP incorporation efficiency $(k_p/K_d, d_{\text{NTP}})$ was also calculated for each enzyme. The dNTP incorporation fidelity was determined using Equation 6:

$$
\text{Fidelity} = (k_p/K_{d,\text{dNTP}})_{\text{incorrect}} / \left[\left(k_p/K_{d,\text{dNTP}} \right)_{\text{correct}} + \left(k_p/K_{d,\text{dNTP}} \right)_{\text{incorrect}} \right] \tag{6}
$$

DNA Trap Assays

A preincubated solution of hPolη (130 nM) and 5′-[32P]-labeled DNA (20 nM) in buffer R was rapidly mixed with a solution of 5 μM unlabeled DNA trap D-1 (Table 1) and a correct dNTP at a saturating concentration in buffer R for various times before the reactions were quenched with 0.37 M EDTA. Reaction products were resolved and quantitated as described above. The plot of the product concentration as a function of reaction time was fit to Equation 7:

$$
[Product] = EoA1 [1 - exp(-k1t)] + EoA2 [1 - exp(-k2t)] \t(7)
$$

where E_0 is the active hPoln concentration, A_1 and A_2 the reaction amplitudes of the first and second phases, respectively, and k_1 and k_2 the rate constants of the first and second phases, respectively.

RESULTS

Bypass of dGAP catalyzed by human Y-family DNA polymerases

To determine if each of the four human Y-family DNA polymerases was able to bypass dG^{AP} (Supplementary Figure 1), we performed running start assays (Figure 1) with both a control DNA substrate 17-mer/26-mer and a damaged DNA substrate 17-mer/26-mer-dGAP containing a site-specifically placed dG^{AP} on template 26-mer-dG^{AP} (Table 1). Notably, hPolκ synthesized much more 25-mer than 26-mer with 17-mer/26-mer after long reaction

times while an intermediate product 24-mer significantly accumulated between 6 s to 60 s (Figure 1C). Because hPolκ is known to create −1 frameshift mutations11, 14, 32, 33 and the template 26-mer contains a dC-rich 5′-terminus (Table 1), the 25-mer was likely the true full-length product synthesized by hPolκ after hPolκ slipped once within the dC-rich region. The strong early accumulation of 24-mer in Figure 1C supports this possibility. If the possibility is proven to be true, the small amount of 26-mer was likely generated by hPolκ via a single blunt-end addition onto the 25-mer as we have previously observed with Dpo4.34 Interestingly, the blunt-end addition phenomenon was more significant with hPolη (Figure 1A). This enzyme has been found previously to catalyze blunt-end additions onto other DNA substrates.13, 35, 36 With the control template 26-mer, hPolη, hPolκ, and hPolτ were able to synthesize full-length products after 1 s, 6 s, and 60 s, respectively (Figures 1A, 1C, and 1E). In stark contrast, hRev1 only incorporated five nucleotides and did not synthesize full-length products even after 12 hrs (Figure 1G). The reaction was not carried out beyond 12 hrs was due to both enzyme stability and *in vivo* relevance. Interestingly, with the damaged template 26 -mer-d G^{AP} , hPoln was not significantly stalled by dG^{AP} and synthesized the full-length products (26-mer and 27-mer) after 3 s (Figure 1B). In comparison, hPolκ and hPolτ needed 60 s and 7,200 s, respectively, to achieve the same task (Figures 1D and 1F). Thus, the presence of a single dGAP slowed hPolκ and hPolτ by 10 and 120-fold, respectively. In addition, there was a significant accumulation of intermediate products 20-mer and 21-mer with both hPolκ and hPolτ, indicating that these enzymes encountered significant difficulty in bypassing the lesion and carrying out the subsequent extension steps (Figures 1D and 1F). As expected, hRev1, a dCTP polymerase, did not synthesize full-length products with 26-mer-dG^{AP} (Figure 1H). The longest product after 12 hrs was a small of 21-mer, suggesting that hRev1 was able to incorporate a nucleotide opposite dGAP but could not extend the DNA lesion bypass product.

We further analyzed these running start assays by determining both dG^{AP} bypass% using Equation 1 (Material and Methods) and bypass efficiency t_{50} ^{bypass} for each enzyme.¹¹ A plot of dG^{AP} bypass% versus reaction time for each enzyme (Supplementary Figure 2) yielded the corresponding t_{50} ^{bypass} value (Table 2), which was defined as the time required for each enzyme to bypass 50% of the total dG^{AP} sites encountered. Similar analysis was also performed with control DNA and yielded t_{50} values for each enzyme to traverse the corresponding dG site (Table 2). These t_{50} ^{bypass} values (Table 2) indicate that hPoln bypassed $dG^{\bar AP}$ encountered and bypassed 50% of dG^{AP} sites 1.6- and 42.6-fold faster than hPolk and hPolt, respectively. When comparing the t_{50} ^{bypass} and t_{50} values for each enzyme, hPolη, hPolκ, and hPolτ became 3.1-, 2.7-, and 7.1-fold slower, respectively, in the presence than in the absence of dGAP. Thus, dGAP had only a slight inhibitory effect on the overall DNA polymerase activity of hPoln and hPolk but significantly slowed DNA synthesis catalyzed by hPolt. The varying effects of dG^{AP} on these three Y-family enzymes suggest that each enzyme may utilize a unique mechanism for dG^{AP} bypass.

Effects of dGAP on DNA binding affinities

The accumulation of the intermediate 21-mer during dG^{AP} bypass (Figure 1) suggests that the presence of the bulky 1-AP adduct could have weakened DNA binding to human Yfamily DNA polymerases. To evaluate this possibility, we estimated the binding affinity of human Y-family enzymes to 20-mer/26-mer, 20-mer/26-mer-dGAP, 21-mer/26-mer, 21-mer/ 26-mer-dGAP (Table 1) via gel electrophoresis mobility shift assays (EMSA). An example of gel image for EMSA using hPoln and 20-mer/26-mer-d G^{AP} is shown in Supplementary Figure 3. The K_d DNA values for the binding of hPoln to other DNA substrates were measured similarly and are listed in Table 3. Interestingly, the calculated ratios in Table 3 demonstrate that the presence of dGAP increased the binding affinity of DNA to hPolη. The structures of Dpo4 with BPDE-d G^{37} and yeast Poln with AAF-d G^{38} suggest that the

hydrophobic 1-AP adduct in complexes hPolη•20-mer/26-mer-dGAP and hPolη•21-mer/26 mer-dGAP likely interacted with the amino acid residues in the Little Finger domain of hPolη.

When EMSA was performed for each of the other human Y-family enzymes, we were only able to estimate the K_d _{DNA} values for hRev1 (Table 3). There was no obvious mobility shift with hPolκ and hPolτ during EMSA, likely due to their weak binding to DNA. For hPolτ, active site titration assays (Materials and Methods) were employed to estimate the K_d , DNA values (Table 3). These values were similar to a previously reported value of 44 ± 7 nM with an undamaged DNA substrate.³⁹ We did not measure the K_d , _{DNA} of hPol τ •21-mer/ 26 -mer-d G^{AP} because catalysis (see below) was slower than the equilibration of the enzyme and DNA ($E + DNA \leftrightarrow E\bullet DNA$). For hPolk, it is not proper to use active site titration assays to estimate the K_d , $_{\text{DNA}}$ values because only 15% of recombinant hPolk has been previously found to be active in such an assay.⁴⁰ With undamaged DNA, the K_d , _{DNA} of a similar hPolk construct used in this paper (Materials and Methods) has been estimated to be 96 ± 21 nM at 25 \degree C by using a fluorescence anisotropy assay.⁴¹ Based on the above published and measured K_d , $_{\text{DNA}}$ values as well as calculated affinity ratios in Table 3, we concluded that hPolη bound to both damaged and undamaged DNA with the highest affinity and the impact of the 1-AP adduct on DNA binding to human Y-family DNA polymerases was not significant enough to account for the strong polymerase pausing during TLS shown in Figure 1. As we have discovered previously with other DNA lesions, $2\overline{6}$, 29 , 42 it is likely that nucleotide incorporation kinetics was dramatically influenced by dG^{AP} .

Effect of dGAP on the kinetics of nucleotide incorporation

hPoln—To determine the effect of dG^{AP} on nucleotide incorporation efficiency at or near the lesion site, we performed single nucleotide incorporation assays with each human Yfamily DNA polymerase under single-turnover conditions. For example, a preincubated solution of 5'-[$32P$]-labeled DNA 20-mer/26-mer-dG^{AP} (20 nM) and hPoln (130 nM) at 37 °C was rapidly mixed with varying dCTP concentrations for various reaction times before being quenched with EDTA (0.37 M). The reaction mixtures were analyzed by denaturing PAGE and quantitated by using a Typhoon Trio (Materials and Methods). Subsequently, the product concentrations were plotted against reaction times and each time course was fit to Equation 4 to obtain an observed reaction rate constant (k_{obs}) (Figure 2A). The plot of the *k*obs values versus dCTP concentrations was then fit to Equation 5 to obtain an apparent maximum dCTP incorporation rate constant (k_p) of 8.1 \pm 0.2 s⁻¹ and an apparent equilibrium dissociation constant (K_d , _{dNTP}) of 287 \pm 24 μM opposite dG^{AP} (Figure 2B). The efficiency $(k_p/K_d, dNTP)$ of correct dCTP incorporation opposite dG^{AP} was then calculated to be $2.8 \times 10^{-2} \mu M^{-1} s^{-1}$ (Table 4).

To determine if nucleotide incorporation efficiency and fidelity were affected within the vicinity of the bulky 1-AP adduct, we performed similar single-turnover kinetic assays for each correct or incorrect nucleotide with control or damaged DNA substrates and the resulting kinetic parameters are listed in Table 4 and Supplementary Table 1. In the presence of either 20-mer/26-mer or 21-mer/26-mer (Table 1), hPolη incorporated correct nucleotides with 50-500 fold higher k_p/K_d , dNTP values than incorrect nucleotides, which are mainly derived from the *kp* differences (Supplementary Table 1). Relative to corresponding control DNA substrates, hPolη incorporated correct nucleotides onto 20-mer/26-mer-dGAP and 21 mer/26-mer-dG^{AP} with 6-18 fold lower k_p values and 3-fold higher $K_{d, dNTP}$ values, leading to 20-48 fold lower k_p/K_d , $dNTP$ values (Table 4). The large decreases in nucleotide incorporation efficiency with the damaged DNA appear to be contradictory to the conclusion of weak polymerase pausing as revealed by Figure 1B. However, the overall polymerase efficiency at the lesion site and the immediate downstream site only dropped by

7-16 fold if the 3-fold tighter DNA binding affinity (Table 3) was taken into consideration. Moreover, the relative high k_p values (2-8 s⁻¹) for correct nucleotide incorporation further diminished intermediate product accumulation near the lesion site in Figure 1B. Interestingly, the nucleotide incorporation fidelity of hPolη was 16-fold lower on average with the damaged DNA substrates than undamaged ones and thus, significant percentages of incorporated nucleotides opposite dG^{AP} and the downstream template base dC were incorrect (Table 4). Remarkably, hPolη misincorporated dCTP onto 21-mer/26-mer-dGAP with 3-fold higher k_p and k_p/K_d dNTP values than it incorporated correct dGTP. Taken together, hPolη was highly error-prone although it catalyzed efficient TLS of dG^{AP}.

hPolk—The kinetic parameters for correct and incorrect nucleotide incorporations with control (Supplementary Table 2) and damaged (Table 5) DNA substrates were determined under similar single-turnover kinetic conditions as described for hPolη (Supplementary Figure 4). Like hPolη, hPolκ incorporated correct dNTPs onto damaged DNA with decreased efficiency than onto undamaged DNA. Notably, the decrease is less dramatic during the dGAP bypass step (5.7-fold) than during the extension step (208-fold) and hPolκ was 123-fold less efficient during the later step than during the former step. These efficiency differences and the low k_p value $(2.3 \times 10^{-3} \text{ s}^{-1})$ for correct dGTP incorporation onto 21mer/26-mer-dGAP contributed to the strong accumulation of the intermediate product 21 mer (Figure 1D).

With control DNA substrates, the nucleotide incorporation fidelity of hPolκ was calculated to be in the range of 10^{-3} to 10^{-2} (Supplementary Table 2). The low nucleotide incorporation fidelity of hPolκ was further worsened by 2-94 folds and was in the range of 10⁻² to 10⁻¹ with the damaged DNA substrates (Table 5). The probability for hPolk to incorporate incorrect nucleotides during TLS of dGAP was estimated to be as high as 30% (Table 5).

hPolτ—Like hPolη and hPolκ, the kinetic parameters for nucleotide incorporations with undamaged (Supplementary Table 3) and damaged (Table 6) DNA substrates were determined by employing single-turnover kinetic assays (Supplementary Figure 5). Since misincorporation of either dATP or dCTP onto 21-mer/26-mer-dGAP catalyzed by hPolτ was not detected even after eight hours, the corresponding kinetic parameters with either 21 mer/26-mer-dGAP or 21-mer/26-mer were not determined. Like hPolκ, hPolτ incorporated correct nucleotides onto damaged DNA with lower efficiency than onto undamaged DNA and the impact was determined to be 7- and 4,545-fold with 20-mer/26-mer-dGAP and 21 mer/26-mer-dGAP, respectively (Table 6). Accordingly, 21-mer/26-mer-dGAP was extended by hPolt with more than 1,000-fold lower k_p and k_p/K_d , dNTP values than 20-mer/26-merdGAP. These kinetic differences led to the strong accumulation of the intermediate product 21-mer in Figure 1F.

With 20-mer/26-mer-dG^{AP}, all three incorrect nucleotides were incorporated onto 20-mer/ 26-mer-dGAP by hPolτ with comparable low efficiencies and probabilities (Table 6). In comparison, incorrect dTTP was incorporated onto 21-mer/26-mer-dGAP with similar efficiency and probability to those of correct dGTP (Table 6). These kinetic data suggest that hPol τ was more error-prone during the subsequent extension step than during dG^{AP} bypass.

hRev1—Consistent with Figure 1H, hRev1 could not incorporate any nucleotide onto 21 mer/26-mer-dG^{AP} after three hours. Thus, we only determined the kinetic parameters for nucleotide incorporations onto 20-mer/26-mer-dGAP (Table 7) and the corresponding control substrate 20-mer/26-mer (Supplementary Table 4) under single turnover conditions (Supplementary Figure 6). Like the other three human Y-family enzymes, hRev1 incorporated dCTP opposite dGAP with 22-fold lower efficiency than opposite undamaged

dG, which was predominantly contributed by the k_p difference. In addition, all three incorrect dNTPs were also incorporated with lower efficiency when the template base dG was damaged, leading to a higher fidelity with 20-mer/26-mer-dGAP. Thus, hRev1 was slightly more error-free but catalytically slower as it bypassed dGAP.

Biphasic kinetics of nucleotide incorporation catalyzed by hPolη

Previously, we have shown that correct nucleotide incorporation opposite dG^{AP} and the downstream template base dC catalyzed by Dpo4 in the presence of a DNA trap follows biphasic kinetics and that the amplitudes of both the fast and slow phases are higher during the dG^{AP} bypass step.²⁶ To explore if human Y-family DNA polymerases also follow similar biphasic kinetics during dG^{AP} bypass in the presence of a DNA trap, hPoln was chosen as a model enzyme because it bound DNA tightly (Table 3), a condition for the DNA trap assay.26, 29, 42-45 Here, we performed single dGTP incorporation onto either 21-mer/26 mer or 21-mer/26-mer-dGAP in the presence of an unlabeled DNA trap which was in large excess over 5′-radiolabeled DNA. First, we determined the effectiveness of 5 μM D-1 (Table 1) as a DNA trap. This trap was effective to prevent correct dGTP incorporation onto radiolabeled 21-mer/26-mer (20 nM) by hPolη within 33 s (Supplementary Figure 7). Next, we conducted the DNA trap assay for correct dGTP incorporation onto either 21-mer/26 mer or 21-mer/26-mer-dG^{AP} (Material and Methods). The concentration of products was plotted as a function of reaction time (Figure 3) and the plot was fit to Equation 6 to yield kinetic parameters listed in Supplementary Table 5. Unlike Dpo4,²⁶ hPoln followed biphasic kinetics with both control and damaged DNA substrates. With 21-mer/26-mer, the amplitude of the fast kinetic phase $(A_1 = 39\%)$ is smaller than that of the slow kinetic phase $(A_2 = 47\%)$ while the former $(k_1 = 72.2 \text{ s}^{-1})$ occurs with a higher reaction rate constant than the latter $(k_2$ $= 4.4 \text{ s}^{-1}$). Interestingly, the sum of the fast and slow reaction rate constants $(A_1k_1 + A_2k_2 =$ 30 s⁻¹) for 21-mer/26-mer is close to the k_p value (39 s⁻¹, Supplementary Table 1) determined by using the single-turnover kinetic assay. For dGTP incorporation onto 21-mer/ 26-mer-dG^{AP}, both A_1 and k_1 were larger than corresponding A_2 and k_2 (Supplementary Table 5). The sum of reaction rate constants for 21-mer/26-mer-dG^{AP} ($A_1k_1 + A_2k_2 = 1.5$ s⁻¹) is also approximately equal to the *k_p* value in Table 4 (2.2 s⁻¹). Similar relationship has been previously observed with other lesions and Dpo4.26, 29, 42-45 This is not surprising because both k_p and the sum $(A_1k_1 + A_2k_2)$ reflect the rate constant of nucleotide incorporation during single binding event between an enzyme and DNA. Furthermore, all four biphasic kinetic parameters with 21-mer/26-mer-dGAP are smaller than the counterparts with 21-mer/26-mer, suggesting that the presence of dG^{AP} negatively affected the biphasic kinetics of nucleotide incorporation in the presence of a DNA trap.

DISCUSSION

In this paper, we have compared the abilities of four human Y-family DNA polymerases to catalyze TLS of dG^{AP} in the same reaction conditions. All of them except hRev1 bypassed a single site-specifically placed lesion dGAP and synthesized full-length products with different efficiency (Figure 1, Table 2). There was more accumulation of the intermediate product 21-mer than 20-mer synthesized by hPolκ (Figure 1D) and hPolτ (Figure 1F). This suggests that for these enzymes, the dG^{AP} bypass step was less problematic than the subsequent extension step. Similar strong polymerase pausing immediately downstream from a bulky DNA lesion site has also been observed for the bypass of $dG^{AP\,26}$ and AAF dG^{46} catalyzed by Dpo4 and for the bypass of BPDE- dG^{47} and AAF-dG catalyzed by yeast Poln.³⁸ To understand the polymerization patterns in Figure 1, it is necessary to establish a kinetic basis for how each human Y-family DNA polymerase catalyzes TLS of dG^{AP}.

Comparison of efficiency of human Y-family DNA polymerases with undamaged DNA

For efficient DNA synthesis, a DNA polymerase must bind to DNA tightly and incorporate nucleotides with high substrate specificity (k_p/K_d) dNTP). Since the K_d DNA values do not change significantly within the temperature range of 23 to 37 °C, we ranked the DNAbinding affinity order of human Y-family enzymes as $hPoln > hPoln > hPolk \approx hRev1$ based on the K_d , DNA values (Table 3) and the affinity of undamaged DNA to hPolk (96 \pm 21 nM) measured by fluorescence anisotropy at 25 °C.⁴¹ Notably, the K_d , _{DNA} values of human Yfamily DNA polymerases were within 5-fold and are not the major factor to differentiate their relative polymerization efficiency. In contrast, the k_p/K_d dNTP values for correct nucleotide incorporation onto undamaged DNA by these enzymes varied by several orders of magnitude (Supplementary Tables 1-4) and likely governed the polymerization efficiency order. For example, the order of k_p/K_d , Δ_{MTP} values of dCTP incorporation opposite dG in 20-mer/26-mer was as followed: hPolη (0.56 μM⁻¹s⁻¹) > hRev1 (0.14 μM⁻¹s⁻¹) > hPolκ (0.037 μM⁻¹s⁻¹) > hPolτ (0.0016 μM⁻¹s⁻¹). Similarly, the *k_p*/*K_{d, dGTP} values for correct* dGTP incorporation onto 21-mer/26-mer were ranked as hPoln $(0.84 \mu M^{-1} s^{-1})$ > hPolk (0.011 μM⁻¹s⁻¹) > hPolτ (0.0016 μM⁻¹s⁻¹). Although we did not measure the *k_p*/*K_{d, dNTP}* value for dGTP incorporation onto 21-mer/26-mer by hRev1, our previous work has shown that this deoxycytidyl transferase preferentially incorporates dCTP, rather than a correct incoming nucleotide, opposite dC, dA, or dT with 10^3 -10⁴ fold lower $k_p/K_{d, dNTP}$ than incorporating dCTP opposite dG.¹⁸ Accordingly, the $k_p/K_{d, \text{ dNTP}}$ values for dGTP incorporation onto 21-mer/26-mer follow the order of hPolη > hPolκ > hPolτ > hRev1. Although we did not determine the k_p/K_d , dNTP values for correct dNTP incorporations at other positions along template 26-mer, the k_p/K_d dNTP order was expected to follow hPoln > hRev1 > hPolκ > hPolτ at each of the two template dG positions and hPolη > hPolκ > hPolτ > hRev1 at each of the other five template positions. Taken together, the overall efficiency for the elongation of primer 17-mer to 26-mer along the template 26-mer is expected to follow the order of $hPol\eta > hPol\kappa > hPol\tau >> hRev1$. Interestingly, this prediction was consistent with the speed order of elongating 17-mer/26-mer to full-length products by human Y-family enzymes: hPol η (1 s) > hPol κ (6 s) > hPol τ (60 s) >> hRev1 (> 12 hrs) (Figure 1).

Comparison of the TLS of dGAP efficiency catalyzed by human Y-family DNA polymerases

When human Y-family enzymes elongated 17-mer along 26-mer-d G^{AP} (Figure 1) the most troubled steps were the $dG^{\dot{A}P}$ bypass step and the subsequent extension step. Opposite dG^{AP} , $dCTP$ was incorporated onto 20-mer/26-mer-d G^{AP} with the following efficiency order of hPol η (2.8×10⁻² μ M⁻¹s⁻¹) > hRev1 = hPolk (6.5×10⁻³ μ M⁻¹s⁻¹) > hPolt $(2.3 \times 10^{-4} \,\mu\text{M}^{-1}\text{s}^{-1})$ (Tables 4-6), which is consistent with the t₅₀bypass order of hPoln (2.5) s) < hPol κ (4.1 s) < hPol τ (107 s) (Table 2). During the subsequent extension step, dGTP was incorporated opposite dC in 21-mer/26-mer-dG^{AP} with the k_p/K_d , Δ_{NTP} order of hPoln $(1.8×10⁻² μM⁻¹s⁻¹ > hPolκ (5.3×10⁻⁵ μM⁻¹s⁻¹) > hPolτ (2.2×10⁻⁷ μM⁻¹s⁻¹). Since$ hRev1 did not elongate 21-mer/26-mer-dG^{AP} after several hours, its k_p/K_d dNTP value should be much lower than that of hPolτ. Thus, the order of nucleotide incorporation efficiency during the extension step is hPolη > hPolκ > hPolτ >> hRev1. On the basis of these k_p/K_d , Δ_{MTP} values and the small impact of dG^{AP} on the K_d , DNA values (Results), we predicted that the overall efficiency of TLS of dG^{AP} and possibly the elongation of 17-mer/ 26-mer-dG^{AP} to full-length products follow the order of hPol η > hPol κ > hPol τ >> hRev1. Consistently, this order matched the speed order for creating full-length product 26-mer along template 26-mer-d G^{AP} (Figure 1). Thus, hPoln is the most efficient human Y-family DNA polymerase to catalyze TLS of dG^{AP}. Interestingly, the same conclusion has been reached by us for the TLS of an abasic site. 11

A kinetic basis for polymerase pausing during TLS of dGAP

In the presence of dG^{AP} , the accumulation of the intermediate products 20-mer and 21-mer was more obvious with hPolκ (Figure 1D) and hPolτ (Figure 1F) than with hPolη (Figure 1B). These data indicated that these former enzymes had different extents of difficulty in either binding damaged DNA or incorporating nucleotides opposite both dG^{AP} and the downstream template base dC on template 26-mer- dG^{AP} . Table 3 displays the small impact of dG^{AP} on the K_d , DNA values of human Y-family DNA polymerases. In contrast, the $k_p/$ K_d , d _{dNTP} values were drastically affected by the presence of dG^{AP} (Tables 4-6). For example, hPolη, hPolκ, and hPolτ were 20-, 6-, and 208-fold, respectively, less efficient when incorporating dCTP opposite dG^{AP} than opposite dG in undamaged DNA. During the subsequent extension step, the embedded dG^{AP} affected the k_p/K_d , ∂_{NTP} values of hPoln, hPolκ, and hPolτ by 48-, 208-, and 4,565-fold, respectively. Thus, the large decrease in correct nucleotide incorporation efficiency caused by dG^{AP} was the main reason why these human Y-family enzymes paused during TLS of dG^{AP} (Figure 1). Specifically, the decreases in k_p/K_d , dNTP values were contributed by 2-4 fold higher K_d , dNTP values and up to 1,000-fold lower k_p values. In addition, the larger negative effect of dG^{AP} on the k_p / K_d , d _{NTP} values of hPol_K and hPol_T than on those of hPol_T explained why both hPol_K and hPol τ paused more strongly than hPol η at the 20th and 21st template positions (Figure 1). The 3-fold higher binding affinity of hPolη to damaged DNA than to undamaged DNA (Table 3) and relatively high k_p values (2 to 8 s⁻¹) for correct nucleotide incorporations opposite both dGAP and downstream dC (Table 4) further lessened hPolη pausing at these sites.

Clearly, the intermediate product 21-mer accumulated more than 20-mer with both hPolκ (Figure 1D) and hPolτ (Figure 1F). This observation suggested that the subsequent extension step was affected by dG^{AP} more than the DNA lesion bypass step with these enzymes. Consistently, the k_p/K_d dNTP value for correct nucleotide incorporation was 123-fold lower during the extension step than during the lesion bypass step with hPolκ (Table 5) while the difference was 1,045-fold lower with hPolτ (Table 6). Additionally, these efficiency differences were mainly contributed by \sim 1,000-fold decrease in k_p . Thus, the conversion from 21-mer to 22-mer was less efficient and much slower than the conversion from 20-mer to 21-mer, leading to the stronger accumulation of 21-mer in Figures 1D and 1F.

Biphasic kinetics of nucleotide incorporation with hPolη

Surprisingly, we observed a fast phase $(A_1 = 39\%, k_1 = 72.2 \text{ s}^{-1})$ and a slow phase $(A_2 =$ 47% , $k_2 = 4.4 \text{ s}^{-1}$) during correct nucleotide incorporation onto undamaged 21-mer/26-mer catalyzed by hPolη in the presence of a DNA trap (Figure 3). In contrast, only the fast phase has been observed during nucleotide incorporation onto the same undamaged substrate catalyzed by Dpo4.26 At present, it is unclear what contributed to the slow phase observed with hPolη. Mechanistically, it is possible that 39% of the binary complex hPolη•DNA were in a productive conformation $(E\cdot DNA_n^P)$ while 47% of the complex were in a slightly nonproductive manner $(E^{\bullet}DNA_{n}^{N})$. The latter was isomerized to become the former at a rate of $4.4 s⁻¹$. Since the total reaction amplitude $(A_1 + A_2)$ of 86% is close to the calculated percentage of the binary complex (83%) based on the K_d , $_{\text{DNA}}$ value (Table 3), the remaining 14% of 21-mer/26-mer must not be bound by hPolη and were free in solution. For the unprecedented biphasic kinetics of nucleotide incorporation onto normal DNA, we proposed a mechanism shown in Figure 4A.

Like Dpo4,²⁶ biphasic kinetics of correct dNTP incorporation onto 21-mer/26-mer-dG^{AP} was observed with hPoln (Figure 3). In comparison to what was observed with 21-mer/26mer, the total reaction amplitude of the fast (14%) and slow (5.3%) phases with 21-mer/26 mer-d G^{AP} was only 19.3%, much less than the calculated reaction amplitude (93%) based

on the K_d , $_{\text{DNA}}$ value in Table 3. Besides the decrease in reaction amplitudes in both phases, the values of k_1 and k_2 were also reduced by the presence of the bulky 1-AP adduct (Supplementary Table 5). Based on the above calculation, 7% of 21-mer/26-mer-dGAP was free in solution and 73.7% were in a dead-end conformation $(E\cdot DNA_n^D)$ which was not elongated within the observation time window in Figure 3. The physical nature of $\text{E}\cdot\text{DNA}_\text{n}^{\text{D}}$ is unclear at present. It is likely that hPolη bound to DNA in a catalytically incompetent conformation. It is also possible that some hPolη molecules bound to the blunt-end of 21 mer/26-mer-dG^{AP}, rather than the staggered end, as suggested by the formation of 27-mer through a blunt-end addition (Figure 1B). On the basis of the above information, we proposed a kinetic mechanism for TLS of dGAP shown in Figure 4B, which has been previously applied to other DNA lesions and polymerases.26, 29, 42, 44 There are a few indirect structural evidences to support this mechanism. Previously, the NMR structure of a damaged 11-mer duplex reveals that the hydrophobic 1-AP adduct of an embedded dG^{AP}:dC base pair is intercalated onto the DNA helix between adjacent Watson-Crick base pairs with the sugar of the modified dG possessing a *syn* glycosidic torsion angle, and the bases of the dG^{AP} :dC base pair are displaced onto the major groove.⁴⁸ If this conformation of dG^{AP} is preserved within the active site of hPolη, the bulky 1-AP adduct would occupy the position of an incoming dNTP. Such a binary complex would not be elongated without undergoing dramatic structural changes and likely represents E•DNAⁿ ^D. Consistently, a crystal structure of the binary complex of yeast Poln and DNA containing an AAF-dG³⁸ shows that the AAF adduct stacks above the last primer-template base pair as proposed for $E\cdot DNA_n^D$. The other crystal structure of the same binary complex illustrates that the enzyme rotates along DNA and allows the DNA adduct to partially block the active site. If 1-AP is in the quasiintercalative conformation as AAF in the corresponding binary crystal structure, the binary complex hPolη•20-mer/26-mer-dGAP requires subtle to mild structural changes before elongation and thereby exists in the conformation of $E\cdot DNA_n^N$. When 1-AP is completely rotated out of the DNA duplex, hPolη•21-mer/26-mer-dGAP will be in the conformation of $E \cdot DNA_n^P$ and be rapidly elongated after nucleotide binding.

Impact of dGAP on polymerase fidelity

The fidelity of nucleotide incorporation opposite dG^{AP} (Tables 4-7) ranked as hRev1 (10^{-4}) > hPolτ (10⁻³ to 10⁻²) > hPolη (10⁻² to 10⁻¹) = hPolκ (10⁻² to 10⁻¹). The calculated fidelity ratios implied that both hPolη and hPolκ were about 10-fold more error-prone when opposite dG^{AP} than opposite undamaged dG . Interestingly, the bulky 1-AP adduct enhanced the fidelity of hRev1 significantly and hPolt slightly during dG^{AP} bypass. Such an impact of the 1-AP adduct on polymerase fidelity was more negatively significant during the subsequent extension step. Opposite 5'-dC from dG^{AP}, hPoln incorporated incorrect dCTP with 3-fold higher efficiency than correct dGTP, hPolk incorporated incorrect dATP with only 2-fold lower efficiency than correct dGTP, and hPolτ incorporated both correct dGTP and incorrect dTTP with almost equal k_p/K_d dNTP values, leading to lowered polymerase fidelity at this position (Tables 4-6). Taken together, our kinetic studies suggested that hPolη, hPolκ and hPolτ became more error-prone during TLS of dG^{AP} than during normal DNA synthesis.

The pathways for the formation of DNA lesions *in vivo* and whether the DNA lesions stall cellular DNA replication machinery have not been thoroughly explored.⁴⁹ So far, it is known that genetic markers, *e.g. Umu* gene, for mutagenesis activity in several cell lines are elevated in the presence of 1-NP or certain metabolities.²¹⁻²⁵ These results suggest that the Y-family DNA polymerases are active in the presence of lesion dG^{AP} . If so, the mutagenic bypass of dGAP catalyzed by these enzymes may lead to genetic mutations *in vivo.*25, 50-54

In summary, our kinetic data demonstrated that hPolη, hPolκ, and hPolτ were able to bypass dGAP and extend the lesion bypass product with dramatically different efficiency and

fidelity values. In contrast, hRev1 was only able to bypass the bulky DNA lesion. A kinetic basis for polymerase pausing during error-prone TLS of dGAP was established.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1.

Running start assays at 37 °C. A preincubated solution of 100 nM 5'-[$32P$]-labeled 17-mer/ 26-mer (A, C, E and G) or 17-mer/26-mer-dG^{AP} (B, D, F and H) and 1 μM hPoln (A and B), hPolκ (C and D), hPolτ (E and F), or hRev1 (G and H) was rapidly mixed with a solution of four dNTPs (200 μM each) for various times before being quenched with 0.37 M EDTA. Sizes of important products are denoted and the 21st position marks the location of dG^{AP} from the 3′-terminus of the DNA template.

Figure 2.

Kinetics of correct dCTP incorporation onto 20-mer/26-mer-dGAP catalyzed by hPolη under single-turnover conditions. (A) A preincubated solution of hPoln (130 nM) and $5'-[^{32}P]$ labeled 20-mer/26-mer-dGAP (20 nM) was rapidly mixed with increasing concentrations of dCTP (25 μM, •; 50 μM, ■; 100 μM, △; 200 μM, ▲; 400 μM, ○; 800 μM, ◆; 1200 μM, □) for various time intervals at 37 °C. Each time course was fit to Equation 4 to obtain a k_{obs} for a specific dCTP concentration. (B) The plot of k_{obs} values against dCTP concentrations was fit to Equation 5 to yield a $K_{d, dCTP}$ of 287 \pm 24 μ M and a k_p of 8.1 \pm 0.2 s⁻¹.

Figure 3.

Biphasic kinetics of correct dGTP incorporation onto 21-mer/26-mers catalyzed by hPolη in the presence of a DNA trap. A preincubated solution of hPolη (130 nM) and 5'-[32P]-labeled 21-mer/26-mer (20 nM, \bullet) or 21-mer/26-mer-dG^{AP} (20 nM, \bullet) was mixed with the solution of dGTP (0.8 mM, ■; 1.2 mM, •) and a DNA trap D-1 21/41-mer (5 μM, Table 1) for various time intervals. The plot of the product concentrations against reaction times was fit to Equation 7. For 21-mer/26-mer, the fast phase had a reaction amplitude of 7.8 ± 0.3 nM and a reaction rate constant of 72 ± 5 s⁻¹ while the slow phase had a reaction amplitude of 9.5 ± 0.3 nM and a reaction rate constant of 4.4 ± 0.5 s⁻¹. For 21-mer/26-mer-dG^{AP}, the fast phase had a reaction amplitude of 2.9 ± 0.1 nM and a reaction rate constant of 11 ± 1 s⁻¹ while the slow phase had a reaction amplitude of 1.1 ± 0.2 nM and a reaction rate constant of 0.20 ± 0.02 s⁻¹.

Figure 4.

Proposed kinetic mechanisms for nucleotide incorporation opposite undamaged dG (A) and damaged dGAP (B) catalyzed by hPolη. *A*1, fast phase amplitude; *k*1, fast phase reaction rate constant; A_2 , slow phase amplitude; D_0 , the initial concentration of DNA; k_2 , slow phase reaction rate constant; *E*, DNA polymerase; *DNAn*, DNA substrate; *E•DNA^D ⁿ*, dead-end binary complex; *E•DNA^N ⁿ*, nonproductive binary complex; *E•DNA^P ⁿ*, productive binary complex; *E•DNA^N ⁿ•dNTP*, nonproductive ternary complex; *E•DNA^P ⁿ•dNTP*, productive ternary complex; *DNAn*+1, extended DNA product by a base; *PPⁱ* , pyrophosphate.

DNA Substrates for dGAP.

 a **G** designates the 1-AP adduct on C8 position of dG (dG^{AP}).

dGAP bypass efficiencies of human Y-family DNA polymerases.

 a ^{*d*} Calculated as the time required to bypass 50% of the dG^{AP} sites.

b Calculated as the time required to bypass 50% of dG, or 20-mer.

Binding affinity of hPolη, hPolτ, and hRev1 to normal and damaged DNA at room temperature.

ND denoted 'not determined'.

All given errors were derived from data fitting.

 a ^aWith adduct refers to 26-mer-dG^{AP}.

Without adduct refers to 26-mer.

c Calculated as (*Kd*, DNA)Normal/(*Kd*, DNA)Damaged.

d Values were determined using active site titration assays.

Kinetic parameters of nucleotide incorporation onto damaged DNA catalyzed by hPol Kinetic parameters of nucleotide incorporation onto damaged DNA catalyzed by hPoln.

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 $b_{\mbox{\scriptsize The}}$ values are from Supplementary Table 1 using control 26
mer DNA template. *b*The values are from Supplementary Table 1 using control 26mer DNA template.

 c Calculated as $(k_p/K_d,$ dNTP)
incorrect/[(kp/Kd, dNTP)
correct + (kp/Kd, dNTP)
incorrect]. *c*Calculated as (*kp*/*Kd*, dNTP)incorrect/[(*kp*/*Kd*, dNTP)correct + (*kp*/*Kd*, dNTP)incorrect].

*d*Calculated as Fidelitydamaged/Fidelitynormal

.

 e Calculated as $((k_p/K_d,\mathrm{dNTP})\mathrm{damped}/\mathrm{E}(k_p/K_d,\mathrm{dNTP})\mathrm{damage}\mathrm{d}l)\times100.$ *e*Calculated as ((*kp*/*Kd*, dNTP)damaged/[Σ(*kp*/*Kd*, dNTP)damaged])×100.

Kinetic parameters of nucleotide incorporation onto damaged DNA catalyzed by hPol Kinetic parameters of nucleotide incorporation onto damaged DNA catalyzed by hPolk.

Calculated as $(k_p$ Kd, dNTP)normal/ $(k_p$ Kd, dNTP)damaged. α Calculated as $(k_p$ / K_d , dNTP)normal $/(k_p$ / K_d , dNTP)damaged.

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 $b_{\mbox{The}}$ values are from Supplementary Table 2 using control 26
mer DNA template. *b*The values are from Supplementary Table 2 using control 26mer DNA template.

Calculated as (k_pK_d, dNTP) incorrect $|(k_pK_d, \mathrm{dNTP})$ correct $+$ (k_pK_d, dNTP) incorrectles *c*Calculated as (*kp*/*Kd*, dNTP)incorrect/[(*kp*/*Kd*, dNTP)correct + (*kp*/*Kd*, dNTP)incorrect].

*d*Calculated as Fidelitydamaged/Fidelitynormal

.

 e Calculated as $((k_p/K_d,\mathrm{dNTP})\mathrm{damped}/\mathrm{E}(k_p/K_d,\mathrm{dNTP})\mathrm{damage}\mathrm{d}l)\times100.$ *e*Calculated as ((*kp*/*Kd*, dNTP)damaged/[Σ(*kp*/*Kd*, dNTP)damaged])×100.

Kinetic parameters of nucleotide incorporation onto damaged DNA catalyzed by hPolt. Kinetic parameters of nucleotide incorporation onto damaged DNA catalyzed by hPolτ.

*b*The values are from Supplementary Table 3 using control 26mer DNA template.

*c*Calculated as (*kp*/*Kd*, dNTP)incorrect/[(*kp*/*Kd*, dNTP)correct + (*kp*/*Kd*, dNTP)incorrect].

Calculated as $(k_p/K_d, \mathrm{dNTP})$ incorrect/[$(k_p/K_d, \mathrm{dNTP})$ correct + $(k_p/K_d, \mathrm{dNTP})$ incorrect].

*d*Calculated as Fidelitydamaged/Fidelitynormal

.

*e*Calculated as ((*kp*/*Kd*, dNTP)damaged/[Σ(*kp*/*Kd*, dNTP)damaged])×100.

 e Calculated as $((k_p/K_d,\,\mathrm{dNTP})\mathrm{damped}/\mathrm{E}(k_p/K_d,\,\mathrm{dNTP})\mathrm{damage}\mathrm{d}l)\times100.$

Kinetic parameters of nucleotide incorporation onto damaged DNA catalyzed by hRev1. Kinetic parameters of nucleotide incorporation onto damaged DNA catalyzed by hRev1.

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*e*Calculated as ((*kp*/*Kd*, dNTP)damaged/[Σ(*kp*/*Kd*, dNTP)damaged])×100.

 e Calculated as $((k_P/K_d,\,\mathrm{dNTP})\mathrm{damaged}/\mathrm{E}(k_P/K_d,\,\mathrm{dNTP})\mathrm{damaged}]\times100.$