Mismatched and Matched dNTP Incorporation by DNA Polymerase β Proceed via Analogous Kinetic Pathways†

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ABSTRACT: While matched nucleotide incorporation by DNA polymerase β (Pol β) has been well-studied, a true understanding of polymerase fidelity requires comparison of both matched and mismatched dNTP incorporation pathways. Here we examine the mechanism of misincorporation for wild-type (WT) Pol β and an error-prone I260Q variant using stopped-flow fluorescence assays and steady-state fluorescence spectroscopy. In stopped-flow, a biphasic fluorescence trace is observed for both enzymes during mismatched dNTP incorporation. The fluorescence transitions are in the same direction as that observed for matched dNTP, albeit with lower amplitude. Assignments of the fast and slow fluorescence phases are designated to the same mechanistic steps previously determined for matched dNTP incorporation. For both WT and I260Q mismatched dNTP incorporation, the rate of the fast phase, reflecting subdomain closing, is comparable to that induced by correct dNTP. Pre-steady-state kinetic evaluation reveals that both enzymes display similar correct dNTP insertion profiles, and the lower fidelity intrinsic to the I260Q mutant results from enhanced efficiency of mismatched incorporation. Notably, in comparison to WT, I260Q demonstrates enhanced intensity of fluorescence emission upon mismatched ternary complex formation. Both kinetic and steady-state fluorescence data suggest that relaxed discrimination against incorrect dNTP by I260Q is a consequence of a loss in ability to destabilize the mismatched ternary complex. Overall, our results provide first direct evidence that mismatched and matched dNTP incorporations proceed via analogous kinetic pathways, and support our standing hypothesis that the fidelity of Pol β originates from destabilization of the mismatched closed ternary complex and chemical transition state.

Polymerase fidelity, at its most basic level, is determined by the free energy difference between the highest energy barriers along the correct and incorrect dNTP incorporation pathways. Since fidelity requires such a comparison, an ever-increasing gap between the structural and mechanistic information available for matched dNTP incorporation compared with that concerning mismatched dNTP incorporation renders the molecular mechanism of DNA polymerase fidelity unclear (1, 2). The thermodynamic instability inherent to polymerase mismatched ternary complexes makes them difficult to crystallize. For this reason, we do not yet have available a structure representing a truly functional prechemistry closed mismatched ternary complex (E′D₃N, Scheme 1), although complexes depicting mispairs within the active site (3, 4) and mismatch extension (5, 6) for various polymerases have been resolved. Likewise, kinetic data for mismatched dNTP incorporations by many DNA polymerases have been reported (7–13), yet commonly used kinetic approaches prevent characterization of individual steps in the mismatched dNTP incorporation pathway.

The Ile260 residue of Pol β is located in the hydrophobic hinge region between the palm and the C-terminal fingers subdomain (14, 15). Several Pol β variants with alterations in residues lining this hinge region have demonstrated compromised fidelity (16–20). The mutator activity of I260Q was first identified by a genetic screen (21), and subsequent kinetic characterization showed that while this mutant possessed parameters of DNA binding similar to that of wild-type, it interestingly showed loose binding discrimination of mismatched dNTP substrates, and thus a low fidelity (22). These characteristics make the I260Q mutant of specific
interest in fidelity studies of Pol β, in order to further understand how one mutation can so drastically alter the mismatch discrimination profile of this enzyme.

Stopped-flow analyses of Pol β, using either 2-aminopurine (2-AP), a fluorescent analogue of adenine and guanine, or Pol β’s sole Trp325 residue, have been very important in advancing Pol β mechanism studies (23–26). Both probes serve as independent reporters of global enzyme–DNA conformational changes which occur during the catalysis of dNTP incorporation into DNA. While monitoring the fluorescence emission of either 2-AP or Trp325 in stopped-flow, the resultant trace observed during correct dNTP incorporation possesses two phases of fluorescence change. Intricate studies involving a variety of chemical probes have been conducted in order to determine the nature of these two phases. The collective findings of these studies support a model in which the fast fluorescence transition is attributed to dNTP-induced subdomain conformational change (step 2, Scheme 1) to form the ternary closed complex prior to chemistry (step 4, Scheme 1), and the slow fluorescence transition is attributed to subdomain reopening (step 6, Scheme 1) after the rate-limiting chemical step.

This work aims to apply our previously established stopped-flow fluorescence methods, as well as steady-state fluorescence spectroscopy to explore two specific aspects of polymerase fidelity: (i) the mechanism by which WT Pol β discriminates against incorrect dNTP substrate to maintain a moderately high fidelity, and (ii) the deviation of this mechanism in generation of a lower fidelity variant by a single I260Q mutation. Such topics are significant in developing a full understanding of the unique means by which DNA polymerases are able to choose a correct dNTP substrate, whose identity is ever-changing with each round of catalysis, from a pool four nucleotides. Moreover, while it is interesting to comprehend how a polymerase accomplishes this feat to achieve a certain fidelity in DNA replication, it is equally interesting to investigate how a single residue mutation can alter its fidelity. Here, we focus on the kinetic pathway of misincorporation, while another recent paper using small-angle X-ray scattering (SAXS) and molecular modeling lends complementary insights into structural aspects of misincorporation (27). The results of our analyses suggest that Pol β incorporates both correct and mismatched dNTP via analogous kinetic mechanisms differing at the transition state of the chemical step. Furthermore, the divergence in fidelity between WT and I260Q is consequent of differences in ability to stabilize the mismatched ternary complex and chemical transition state.

**MATERIALS AND METHODS**

**DNA Substrates.** The sequences of primer/template DNA substrates used in the Pol β studies are listed in Table 1. In the text, the notation X:Y indicates [template base]:[incoming dNTP]. Custom synthesized oligomers were purified by denaturing PAGE (15–18% acrylamide, 7 M urea), and desalted using a reverse phase C18 cartridge. Oligomer concentrations were determined by UV absorbance at 260 nm with the following extinction coefficients (M⁻¹ cm⁻¹): 19-mer primer, ε = 166,100; 18-mer primer, ε = 158,900; 36AP template, ε = 335,200; 35AP template, ε = 327,100;
and 15-mer downstream oligomer, $\varepsilon = 144,800$. Primer, template, and downstream oligomer (where applicable) were annealed in a 1:1:1.2 molar ratio, respectively. Downstream oligomers utilized were 5' phosphate. DNA primers used in the chemical quench experiments were $3^5$P-labeled at the 5'-end using T4 polynucleotide kinase and [$\gamma^3$P]ATP (4500 Ci/mol) according to the manufacturer’s protocol. The labeled primers were separated from unreacted ATP using a G-25 microspin column.

Reagents. Ultrapure dNTP’s and G-25 microspin columns were from GE Healthcare (Piscataway, NJ); [$\gamma^3$P]ATP from MP Biomedicals (Irvine, CA); T4 polynucleotide kinase from USB (Cleveland, OH); and reverse phase C18 cartridges from Waters (Milford, MA). All synthetic oligonucleotides used in DNA substrate preparation were provided by Integrated DNA Technologies (Coralville, IA). Materials and reagents not listed were of standard molecular biology grade.

Cloning and Purification of Full-Length Pol $\beta$ and I260Q. The plasmid pET17b-Pol $\beta$ was transformed into Escherichia coli strain BL21(DE3)pLysS (Stratagene) to overexpress and purify rat Pol $\beta$ (335 amino acids, 39 kDa) as previously described (28). To generate the I260Q mutant, site-directed mutagenesis was carried out using primers: 5'-GGGAAATCGATATCGAGTCCGACCCGAGACTACG-TACTAATGGG-3' and 5'-CCACAGTGAATCTGATCGTCTG-3' using the QuikChange method (Stratagene). The underlined region represents the site of mutagenesis. Successful mutagenesis was verified by both plasmid sequencing (OSU Plant Microbe Genomics Facility) and electrospray ionization mass spectrometry of the mutant protein (OSU Campus Chemical Instrument Center). The I260Q mutant was overexpressed and purified in the same manner as wild-type (WT) Pol $\beta$ (28).

Reaction Buffer Composition. Assay buffers consisted of 20 mM BisTrisHCl, 50 mM TrisHCl, 2 mM DTT, 10–35% (w/v) glycerol, with the ionic strength adjusted to 81 mM with KCl, free Mg$^{2+}$ adjusted to 5 mM with MgCl$_2$, at either pH 7.6 or 8.5 (adjusted at 37 °C). Standard Pol $\beta$ assay buffer contained 10% glycerol: all deviations from this are noted accordingly in the figure legends. Free Mg$^{2+}$ concentrations were calculated using the WEBMAXC STANDARD program at http://www.stanford.edu/~cp Patton/webmaxc.htm, using the appropriate pH, temperature, ionic strength, and nucleotide triphosphate concentration.

Stopped-Flow Fluorescence. Stopped-flow fluorescence assays were performed on an SX 18 MV stopped-flow apparatus (Applied Photophysics Ltd., U.K.). The excitation wavelength for 2-aminopurine (2-AP) was 312 nm with a spectral bandpass of 4 nm. Fluorescence emission was monitored using a 360 nm high pass filter (Corion, MA) on a logarithmic time scale for 1 s for correct dNTP incorporation and up to 50 s for mismatched dNTP incorporation (depending on exact conditions). A typical reaction was initiated by rapid mixing of equal volumes of two solutions in degassed Pol $\beta$ assay buffer: solution A, containing 400 nM DNA substrate preincubated with 1 $\mu$M DNA polymerase, and solution B, containing suitable correct or incorrect dNTP. In all cases, both syringes contained equal concentrations of MgCl$_2$, since it is known that this can have a significant effect on observed fluorescence (24). In experiments employing altered buffer viscosity, both stopped-flow solutions A and B contained the indicated concentration of glycerol. Typically, for all reactions a minimum of 7 runs were performed and averaged prior to data analysis.

Rapid chemical quench. Follow-up of stopped-flow fluorescence experiments was performed on a KinTek (State College, PA) rapid chemical quench instrument (29). The rapid chemical quench reactions were initiated exactly as described for stopped-flow, except that the upstream primer was 5'-radiolabeled. Reactions were quenched at various time intervals with 0.6 M EDTA, and the products visualized by denaturing PAGE followed by autoradiography using a STORM 840 PhosphorImager and quantitated using ImageQuant 5.0 software (GE Healthcare, U.K.).

Steady-State Fluorescence Spectroscopy. Steady-state fluorescence experiments were performed on an AminoBowman Series 2 luminescence spectrometer. 2-AP was excited at 312 nm, and the emission spectra recorded from 300 to 500 nm with a step size of 1 nm. In Pol $\beta$ assay buffer, reactions consisted of 200 nM deoxy-terminated DNA substrate, 500 nM DNA polymerase, and either correct (300 $\mu$M) or incorrect (5 $\mu$M) dNTP.

Product and Data Analysis. Rapid chemical quench data were fit using Sigma Plot 9.0 (Jandel Scientific, CA) to a single exponential equation: [DNAn+1] = $A_1 (1 - e^{-k_{quench}t})$, where $A_1$ is amplitude and $k_{quench}$ is the observed rate constant of single-nucleotide incorporation. Stopped-flow fluorescence traces ($>2$ ms) were fit using Applied Photophysics software to a double exponential equation: fluorescence = $A_1 e^{-t/k_{fast}} + A_2 e^{-t/k_{slow}} + C$, where $A_1$ and $A_2$ represent corresponding amplitudes, $C$ is an offset constant, and $k_{fast}$ and $k_{slow}$ are the observed rate constants for the fast and slow phases of the fluorescence transition, respectively. For fast phase analysis, $k_{fast}$ was plotted as a function of dNTP concentration and the data fit to a hyperbolic equation with a nonzero intercept: $k_{fast} = k_d[dNTP]/(K_d + [dNTP]) + y_0$. According to Scheme 1, $k_2$ is the microscopic rate constant for the forward conformational closing step, $K_d$ is the thermodynamic dissociation constant for dNTP ($K_d = k_{-L}/k_1$), and $y_0$ is a function of both microscopic constants $k_{-2}$ and $k_2$, as well as the binding constant for catalytic Mg$^{2+}$ as further elaborated in the appendix of ref 30. For slow phase analysis, $k_{slow}$ was plotted as a function of dNTP concentration and the data fit to the hyperbolic equation $k_{slow} = k_{pol}[dNTP]/(K_{d,app} + [dNTP])$, where $k_{pol}$ represents the maximum rate of nucleotide incorporation and $K_{d,app}$ represents the apparent dNTP dissociation constant.

RESULTS

Mismatched dNTP Incorporation by WT Pol $\beta$. Stopped-flow fluorescence assays and rapid quench experiments have been previously performed in our laboratory under a wide variety of conditions in order to elucidate the mechanism for correct dNTP incorporation by WT Pol $\beta$ (24–26). Here we report systematic pre-steady-state kinetic analyses of mismatched dNTP incorporation by WT Pol $\beta$. As shown in Figure 1, the stopped-flow trace of T:G mismatched incorporation catalyzed by WT follows the same biphasic pattern of fluorescence change observed for correct dNTP incorporation, albeit with lower amplitude. Follow-up of mismatched dNTP incorporation in rapid chemical quench yields rates which correspond precisely to the slow phase
The difference during the initial dNTP binding before conformational changes during the step responsible for the fast fluorescence transition. Using the same approaches previously employed for matches (26), likely originates from a step after chemistry.

Using the same approaches previously employed for matches (25), we showed that the rate of the fast conformational closing, while \( K_d \) reflects the stability of the ternary complex before closing. Therefore, WT and I260Q show little slow phases (\( k_{s,app} \)) versus dGTP concentration (after mixing) yields a hyperbolic fit (Figure 1B inset). This implies that the fast fluorescence transition results from a step before chemistry, while the slow fluorescence transition, analogous to the prior assignment for correct dNTP incorporation (26), likely originates from a step after chemistry. Using the same approaches previously employed for matches (25), we showed that the rate of the fast conformational change is slowed as a function of increasing viscosity for mismatches as well (Figure 2, red bars). Sensitivity to viscosity has been previously interpreted to be indicative of large conformational movements within the ternary complex during the step responsible for the fast fluorescence transition.

\( dNTP \) Concentration Dependence of the Fast and Slow Fluorescence Transitions during Mismatch Incorporation. Previous stopped-flow fluorescence assays investigating matched dNTP incorporation showed that both the fast and slow fluorescence transitions demonstrated a hyperbolic dependence on dNTP concentration (24, 30). Here, we analyze the dNTP dependence of both the fast and slow fluorescence phases during mismatched dNTP incorporation in stopped-flow. The observed rate constants for the fast and slow phases (\( k_{fast} \) and \( k_{slow} \)), obtained from a double exponential fit of each stopped-flow trace (see Materials and Methods), individually plotted as a function of dNTP concentration, revealed that both phases demonstrate a hyperbolic dependence on dNTP concentration (Figure 3).

| Table 2: Kinetic Rate and Binding Constants for WT versus I260Q Nucleotide Incorporation into 19/36AP(T)/15 and 19/36AP(C)/15 DNA Substrates at pH 8.5* |
|----------------|----------------|
|                 | WT             | I260Q          |
|                 | wild-type      | 1260Q          |
| \( k_2 \)       | 116 ± 5 s\(^{-1}\) | 108 ± 7 s\(^{-1}\) |
| \( K_d \)       | 30 ± 4 \( \mu \)M | 6.6 ± 2.1 \( \mu \)M |
| \( k_{pol} \)   | 42.9 ± 0.6 s\(^{-1}\) | 43.6 ± 1.4 s\(^{-1}\) |
| \( K_{d,app} \) | 6.8 ± 0.5 \( \mu \)M | 7.1 ± 1.2 \( \mu \)M |
| \( k_{quench} \) | 45.7 ± 2.8 s\(^{-1}\) | 47.6 ± 2.5 s\(^{-1}\) |
| \( k_{pol}/K_{d,app} \) | 6.3 | 6.2 |
| fidelity*       | 0.012 | 0.28 |
| \( k_{cor} \)   | 2.8 s\(^{-1}\) | 2.5 s\(^{-1}\) |
| \( k_{inc} \)   | 0.5 s\(^{-1}\) | 0.2 s\(^{-1}\) |

The \( k_2, K_d, k_{pol} \) and \( K_{d,app} \) values were obtained from hyperbolic fit of the dNTP concentration dependence of the observed rates of the fast and slow fluorescence phases (\( k_{fast} \) and \( k_{slow} \)) as described in Materials and Methods. The rate of dNTP incorporation (\( k_{quench} \)) was obtained from single exponential fit of rapid chemical quench data at a saturating dNTP concentration. The \( k_2 \) value represents the rate constant of forward conformational closing, while \( K_d \) reflects the stability of the ternary complex before closing. Therefore, WT and I260Q show little difference during the initial dNTP binding before conformational change. The \( k_{pol} \) value represents the maximum rate of dNTP incorporation, while the \( K_{d,app} \) value possesses a contribution from all steps up to the rate-limiting step and can be thought as the dissociation constant of the closed ternary complex.

Catalytic efficiency as measured in units of \( \mu \)M\(^{-1}\) s\(^{-1}\). Fidelity defined as \((k_{pol}/K_{d,app})_{cor} + (k_{pol}/K_{d,app})_{inc}/(k_{pol}/K_{d,app})_{inc}\), where the subscripts “cor” and “inc” indicate correct (matched) and incorrect (mismatched) nucleotide incorporation, respectively.

rates obtained by stopped-flow (Figure 1B and Table 2). This suggests that the slow phase is attributed to a conformational step limited by the rate of chemistry. Use of dideoxynucleotide-terminated primer, which eliminates the chemistry step, abolishes the slow phase completely (Figure 1B inset). This implies that the fast fluorescence transition results from a step before chemistry, while the slow fluorescence transition, analogous to the prior assignment for correct dNTP incorporation (26), likely originates from a step after chemistry.

In the case of the fast phase, a plot of the observed rate constant as a function of dNTP was fit to a hyperbola with a nonzero intercept to obtain values for \( k_2 \) (see Scheme 1), the rate constant of forward conformational closing, and \( K_d \), the thermodynamic dNTP dissociation constant (see Materials and Methods). If the fast phase were resultant from the thermodynamic dNTP dissociation constant (see Materials and Methods).

In the case of the fast phase, a plot of the observed rate constant as a function of dNTP was fit to a hyperbola with a nonzero intercept to obtain values for \( k_2 \) (see Scheme 1), the rate constant of forward conformational closing, and \( K_d \), the thermodynamic dNTP dissociation constant (see Materials and Methods). If the fast phase were resultant from the thermodynamic dNTP dissociation constant, one would expect a linear dependence on dNTP concentration. However, the
observed hyperbolic dependence of the fast phase on mismatched dNTP largely indicates that this phase originates from a conformational change induced by mismatched dNTP binding. Similarly, for slow phase analysis, the observed rate constant was plotted against dNTP concentration and fit to a hyperbola to obtain values for $k_{\text{pol}}$, the maximum rate of nucleotide incorporation, and $K_{d,\text{app}}$, the apparent dNTP dissociation constant (see Materials and Methods). These parameters correspond to the pre-steady-state values of $k_{\text{pol}}$ and $K_{d,\text{app}}$ traditionally obtained by rapid chemical quench. The parameters for $k_2$, $K_d$, $k_{\text{pol}}$, and $K_{d,\text{app}}$ obtained for T:A match and T:G mismatch incorporation by WT Pol β are recorded in Table 2.

Overall, all the aforementioned results support that, analogous to our prior assignments for correct dNTP incorporation, the fast and slow fluorescence changes observed for mismatched incorporation (Figure 1) can be assigned to the dNTP-induced subdomain-closing conformational change and the chemical step (which likely limits the reopening step), respectively. An important observation is that the forward rate of conformational closing ($k_2$, Scheme 1) for mismatched dNTP incorporation is comparable with that for correct dNTP incorporation (though with significant increase in $K_d$), while the maximum rate of nucleotide incorporation ($k_{\text{pol}}$) is substantially slower (also with significant increase in $K_{d,\text{app}}$) (Table 2). This suggests that overall mismatched incorporation follows a similar pathway, though both the $K_d$ and $K_{d,\text{app}}$ values are higher and the rate of the chemical step is slower.

**A Biphasic Trace Is Observed for Multiple Mismatches.** To further substantiate the conclusions drawn from the investigation of T:G mismatch incorporation in stopped-flow, several additional mismatches were investigated (T:C, C:A, A:A, A:G, G:A, G:G). A biphasic trace was observed for all mismatches examined (Figure 4), and only the fast phase was present when dideoxy-terminated primer was employed (Figure 4, insets). These results lend credence to the notion that while Pol β incorporates mismatches via the same overall mechanism, involving fast subdomain closure upon dNTP binding, each mismatch is unique in its approach to the transition state, as exemplified by the variance in slow phase rates among the mismatches examined.

**Kinetic Properties of I260Q and WT Are Similar for Correct dNTP Incorporation.** As an extension of the WT studies delineated above, further studies on the I260Q mutant were conducted in the interest of kinetic comparison. Stopped-flow analysis of correct dNTP incorporation (Figure 5 and Table 2) indicates that I260Q behaves similarly to WT in that (a) it possesses a biphasic trace with similar rate constants compared to WT, (b) use of dideoxy-terminated primer abolishes the slow phase, (c) the rapid chemical quench rate matches that of the slow phase, and (d) the viscosity dependence pattern of the fast and slow phases is identical to that of WT (Figure 6). Furthermore, dNTP concentration dependence analysis revealed that for T:A correct incorporation both enzymes have similar $k_2$ values (116 ± 5 s⁻¹ and 108 ± 7 s⁻¹ for WT and I260Q, respectively), $k_{\text{pol}}$ values (42.9 ± 0.6 s⁻¹ and 43.6 ± 1.4 s⁻¹ for WT and I260Q, respectively) and $K_{d,\text{app}}$ values (6.8 ± 0.5 μM and 7.1 ± 1.2 μM for WT and I260Q, respectively), while demonstrating a 4.5-fold difference in $K_d$ (30 ± 4 μM and 6.6 ± 2.1 μM for WT and I260Q, respectively) (Table 2). Likewise, analysis of C:G matched incorporation yielded similar values of $k_{\text{pol}}$ and $K_{d,\text{app}}$ between the two enzymes (Table 2).

**Kinetic Properties of I260Q and WT Are Different for Mismatched dNTP Incorporation.** There are clear kinetic differences between I260Q and WT during mismatched
incorporations. The similarities and differences are summarized here: (a) A comparison in Figure 7 demonstrates that the fluorescence change of I260Q mismatch incorporation follows the same pattern as that of WT, though the slow phase is faster by a factor of 2–3 (Table 2). The assignments of the two phases have again been verified by viscosity dependence (Figure 2) and rapid quench (Table 2). (b) Note that the rate of the conformational change, $k_{2}$, is not slowed down for I260Q relative to WT (Figure 8A,B and Table 2), suggesting that the dNTP-induced conformational closing is not disrupted during mismatched dNTP incorporation by I260Q. This is in contrast to what has been suggested previously (22). (c) The main difference between I260Q and WT lies in the ability of I260Q to more efficiently stabilize the mismatched ternary complex, as suggested by the 10-fold decrease in $K_{d,app}$ of I260Q (49 ± 3 μM, see Figure 8C) relative to WT (489 ± 26 μM) for T:G mismatch incorporation and the 5-fold decrease in $K_{d,app}$ of I260Q (45 ± 3 μM) relative to WT (227 ± 22 μM) for C:A mismatch incorporation (Table 2). This is in agreement with the tight mismatch dNTP binding of this mutant formerly reported (22). The combination of the lower $K_{d,app}$ and faster $k_{pol}$ values observed for I260Q mismatched incorporation yields catalytic efficiencies of 0.28 μM$^{-1}$ s$^{-1}$ for T:G mismatch and 0.21 μM$^{-1}$ s$^{-1}$ for C:A mismatch, which are significantly larger than those recorded for WT (Table 2). Note that the catalytic efficiencies for both enzymes during matched incorporation show little difference. Selective enhancement of mismatch incorporation by I260Q translates into a fidelity 24-fold lower for T:G mismatch incorporation and 16-fold lower for C:A mismatch.

The Fast Fluorescence Transition Is Also Present at Physiological pH. To ensure that the conclusions based on the results obtained at pH 8.5 are also applicable under physiological conditions, we repeated analysis of mismatched dNTP incorporation at pH 7.6. At this pH, a biphasic trace of fluorescence change is still observed in stopped-flow for both WT and I260Q during T:G misincorporation (Figure 9). Similar to that observed for pH 8.5, the observed rate of the dNTP induced conformational change is not slowed down for I260Q relative to WT at pH 7.6, again pointing out that the dNTP-induced conformational closing of I260Q is not disrupted during matched or mismatched dNTP incorporation. A rigorous dNTP titration analysis of the fast phase was not possible for mismatches at this pH due to a combination of the low amplitude of this phase at dNTP concentrations below 2 mM and a noted pattern of inhibition at high dNTP concentration. Slow phase analyses for WT
versus I260Q nucleotide incorporation at pH 7.6 are reported in Table 3. Again, for matched incorporation, little difference is observed for \( k_{\text{pol}} \) and \( K_{d,\text{app}} \) between WT and I260Q at this pH, leading to similar catalytic efficiencies (Table 3). However, a 7.4-fold decrease in the \( K_{d,\text{app}} \) of I260Q (88 ± 6 \( \mu \)M) relative to WT (654 ± 42 \( \mu \)M) for T:G mismatch incorporation, and a 4.3-fold decrease in the \( K_{d,\text{app}} \) of I260Q (69 ± 3 \( \mu \)M) relative to WT (298 ± 23 \( \mu \)M) for C:A mismatch incorporation at pH 7.6 further supports that the main difference between I260Q and WT lies in the enhanced ability of I260Q to stabilize the mismatched ternary complex.

Steady-State Fluorescence Analyses. Steady-state fluorescence emission spectra of WT versus I260Q demonstrate identical binary complex traces. For both enzymes there is an equivalent increase in 2-AP fluorescence intensity at the emission maximum (360 nm) upon addition of saturated concentrations of correct dNTP (Figure 10). Addition of mismatched dNTP also results in a fluorescence change in the same direction, though with lower amplitude. Importantly, I260Q demonstrates higher fluorescence intensity (Figure 10) and lower saturating mismatched dNTP concentration compared to WT (data not shown). The data correlate well to the observed amplitude differences in the stopped-flow fluorescence assays (Figure 1 and Figure 7). Overall, these results are consistent with the proposal that both matches and mismatches are incorporated via a similar mechanism, as both correct and mismatched dNTPs elicit the same direction of fluorescence change. In combination with conclusions from recent SAXS structural studies (27), they also qualitatively support the hypothesis that the enhanced ability of I260Q to incorporate mismatched dNTP may arise from a more “partially closed” mismatched ternary complex structure than WT (see Discussion).

**DISCUSSION**

As an extension of previous methodology used to delineate Pol β’s correct dNTP incorporation mechanism (24, 25), similar techniques were employed to examine the mechanism of mismatched dNTP incorporation by Pol β and its I260Q “mutator” mutant. Detailed studies of these two enzymes led to the following important conclusions: (i) Stopped-flow fluorescence analyses indicate that there is, in fact, a conformational closing event that occurs for mismatched dNTP incorporation, as evidenced by the existence of a fast fluorescence phase preceding chemistry. Furthermore, the rate...
of conformational change induced by mismatched dNTP is comparable to that of subdomain closing induced by correct dNTP. (ii) Comparison studies of WT and I260Q show that I260Q possesses fast conformational closing steps during both matched and mismatched dNTP incorporation as well. This argues against a hindered dNTP closing for this enzyme previously hypothesized to account for its altered fidelity (22). (iii) The main difference between the two enzymes lies in more effective stabilization of the mismatched ternary complex by I260Q. (iv) Steady-state fluorescence studies demonstrate that while both matched and mismatched dNTP elicit the same direction of fluorescence change, the I260Q mismatched ternary complex fluorescence emission spectrum bears more resemblance to that of the matched ternary complex. Recent SAXS and molecular modeling studies from our laboratory suggest that for both enzymes the mismatched ternary complex lies between the open and the closed forms (27). The combination of these conclusions with the steady-state fluorescence studies presented here provides grounds for speculation that, on the global conformational closing pathway ranging from fully open (binary complex) to fully closed (matched ternary complex), mismatches may induce a less “partially closed” (more open) conformation for WT than for I260Q.

Although our observation of an incorrect dNTP-induced conformational change by Pol β may appear to contradict the findings of two FRET-base studies monitoring the conformational motions of Klentaq (31) and Klenow (32), the differences could be reconciled. Both studies report an increase in FRET signal, upon addition of correct dNTP, yet do not observe any significant change upon addition of incorrect dNTP. In contrast to moderate and low fidelity polymerases, including Pol β, since Klentaq and Klenow are higher fidelity enzymes, it is likely that they more effectively destabilize the mismatched ternary complex to the extent that no fluorescence change is observable for mismatch binding. Our observation of differences in the amplitude of stopped-flow fluorescence traces (Figure 7) and steady-state fluorescence spectra (Figure 10) between WT Pol β and the lower fidelity I260Q variant further illustrates an extension of the correlation between fidelity and mismatch destabilization.

Our observation that the conformational change occurs with a similar rate for both correct and mismatched incorporation by Pol β differs from the conclusion of the recent single molecule kinetic analysis of T7 DNA polymerase (33), which reported a significantly reduced rate of conformational closing induced by mismatched dNTP binding. It remains to be established whether such a discrepancy reflects differences in the mismatched discrimination mechanism employed by two polymerases, or whether it results from different experimental systems and conditions. However, another study on T7 reported little difference in the forward rates of conformational closing between matches and mismatches, while noting a large difference in the reverse rates of conformational closing (34).

On the other hand, our results are in full agreement with recent reports on the linear free energy relationship (LFER) of pyrophosphate leaving group elimination for both correct and mismatched dGTP incorporations utilizing dNTP analogues in which the β,γ-bridging oxygen is substituted with various halomethylene moieties (35, 36). Based on observation that the Brønsted correlation between log $k_{pol}$ and the leaving group pK$_a$ is very similar between correct and mismatched incorporations, the authors conclude that their results support our earlier prediction that the rate-limiting step is the chemical step, not the conformational change (1).

Table 3: Kinetic Rate and Binding Constants for WT versus I260Q Nucleotide Incorporation into 19/36AP(T)/15 and 19/36AP(C)/15 DNA Substrates at pH 7.6

|          | T:A (matched) |          | T:G (mismatched) |
|----------|---------------|----------|------------------|
|          | wild-type     | I260Q    | wild-type        | I260Q              |
| $k_{pol}$ | 13.4 ± 0.5 s$^{-1}$ | 13.4 ± 0.4 s$^{-1}$ | 0.26 ± 0.1 s$^{-1}$ | 1.8 ± 0.03 s$^{-1}$ |
| $K_{d,app}$ | 2.2 ± 0.3 µM | 1.1 ± 0.2 µM | 654 ± 42 µM | 88 ± 6 µM |
| $k_{pol}/K_{d,app}$ | 6.1 | 12 | 0.00040 | 0.021 |
| fidelity* |                |          | 15100 | 590 |

|          | C:G (matched) |          | C:A (mismatched) |
|----------|---------------|----------|------------------|
|          | wild-type     | I260Q    | wild-type        | I260Q              |
| $k_{pol}$ | 7.2 ± 0.1 s$^{-1}$ | 6.6 ± 0.2 s$^{-1}$ | 0.19 ± 0.1 s$^{-1}$ | 1.1 ± 0.01 s$^{-1}$ |
| $K_{d,app}$ | 1.0 ± 0.1 µM | 0.46 ± 0.07 µM | 298 ± 23 µM | 69 ± 3 µM |
| $k_{pol}/K_{d,app}$ | 7.0 | 14 | 0.00064 | 0.016 |
| fidelity* |                |          | 10900 | 891 |

* Catalytic efficiency as measured in units of µM$^{-1}$ s$^{-1}$. * Fidelity defined as $(k_{pol}/K_{d,app})_{cor} + (k_{pol}/K_{d,app})_{inc}/(k_{pol}/K_{d,app})_{inc}$, where the subscripts “cor” and “inc” indicate correct (matched) and incorrect (mismatched) nucleotide incorporation, respectively.
The loss of fidelity of I260Q has previously been explained by a hindrance of dNTP-induced subdomain closure (21, 22). It was suggested that upon the substitution of Ile with Gln in I260Q, rotamers are generated which cause a reduction in an important cavity space near residue 260, thus perturbing subdomain motion. However, our results indicate the contrary: that the kinetic properties of I260Q, including the rate of the dNTP-induced subdomain closure, are largely unperturbed from WT Pol β for correct dNTP incorporations. Thus, for correct dNTP incorporation its free energy profile is assumed to be the same as that of WT for the purpose of discussion here. The main difference leading to the low fidelity of I260Q appears to be a loss in ability to destabilize the mismatched ternary complex (Figure 11, blue profile). This conclusion is based on the observation that the low fidelity of I260Q mutant is primarily determined by enhanced efficiency of mismatched incorporation, whereas for most low fidelity DNA polymerase variants, reduced fidelity correlates with low efficiency of correct dNTP incorporation (41). Based upon the two mismatches examined, I260Q incorporates mismatches 23- to 53-fold more efficiently than WT (see Tables 2 and 3). The observed change in $k_{pol}/K_{dapp}$ for mismatched incorporations by I260Q suggests that the “overstabilization” of the mismatched ternary complex by I260Q is further enhanced in the transition state. Molecular modeling of the I260Q mismatched ternary complex predicts the presence of additional hydrogen bonds, involving Gln260, Tyr296, Glu295, and Arg258, which may be a plausible source of this stabilization (27).

Overall, kinetic behavioral comparisons between WT Pol β and I260Q allow us to conclude that infidelity of I260Q originates from enhanced stabilization of the mismatched ternary complex and the chemical transition state. Regarding the mechanism of mismatched discrimination, our laboratory previously suggested that DNA polymerase fidelity originates primarily from differential substrate binding at the highest energy transition state and suggested a most likely model (Figure 1D in ref 41), which is highly consistent with the results and conclusions of this work. Of future interest to enhance our knowledge of the mechanism of polymerase fidelity would be crystallization of the I260Q mismatched ternary complex. This would provide insight into the extent of this variant’s conformational closing profile and delineate a structural basis for understanding which specific residues might contribute to this variant’s unique capability to support mismatched dNTP incorporation more efficiently.

The insights into the mechanism of misincorporation presented here add to the growing body of knowledge ultimately desired to consist of the microscopic rates of all steps in Pol β’s matched and mismatched incorporation mechanism, in order to fully and confidently assess which step is the most important contributor to this enzyme’s fidelity. Recent illumination of the importance of the reverse rate of the dNTP-induced conformational closing step and its effect on enzyme specificity further stresses the necessity for complete evaluation of all microscopic rate constants (34). Ongoing studies in our laboratory are examining this step for Pol β during both matched and mismatched incorporation. On a broader scale, through the use of Pol β we continue to develop a more precise understanding of the basis of DNA polymerase fidelity by establishing our techniques as a systematic framework of investigation that can be applied to multiple polymerases and additional site-specific mutants (42) during both matched and mismatched nucleotide incorporations.

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REFERENCES

1. Showalter, A. K., and Tsai, M. D. (2002) A reexamination of the nucleotide incorporation fidelity of DNA polymerases. Biochemistry 41, 10571–10576.
2. Showalter, A. K., Lamarche, B. J., Bakhtina, M., Su, M. I., Tang, K. H., and Tsai, M. D. (2006) Mechanistic comparison of high-fidelity and error-prone DNA polymerases and ligases involved in DNA repair. Chem. Rev. 106, 340–360.
3. Krahn, J. M., Beard, W. A., and Wilson, S. H. (2004) Structural insights into DNA polymerase beta deterrents for misincorporation support an induced-fit mechanism for fidelity. Structure 12, 1823–1832.
4. Johnson, S. J., and Beese, L. S. (2004) Structures of mismatch replication errors observed in a DNA polymerase. Cell 116, 803–816.
5. Trincao, J., Johnson, R. E., Wolfe, W. T., Escalante, C. R., Prakash, S., Prakash, L., and Aggarwal, A. K. (2004) Dpo4 is hindered in extending a G.T mismatch by a reverse wobble. Nat. Struct. Mol. Biol. 11, 457–462.
6. Batra, V. K., Beard, W. A., Shock, D. D., Pedersen, L. C., and Wilson, S. H. (2005) Nucleotide-induced DNA polymerase active site motions accommodating a mutagenic DNA intermediate. Structure 13, 1225–1233.

FIGURE 11: Qualitative free energy profile of matched and mismatched dNTP incorporation by Pol β versus I260Q. E = DNA polymerase in open conformation; E′ = closed conformation; Dn = DNA; N = Mg·dNTP; M = catalytic Mg$^{2+}$; P = Mg·PP. Our observation of a fast conformational change for mismatched dNTP incorporations supports the interpretation of this work, and additionally argues against the notion that conformational closing is a major contributor to fidelity. Figure 11 illustrates the current knowledge of the kinetic scheme of Pol β and its free energy diagram for matched dNTP incorporation (black profile) (26). What remain to be established are the free energy diagram (i.e., kinetic mechanism) and the intermediate structures (i.e., structural mechanism) for mismatched incorporations. Multiple and recent proposals contemplate a perturbed conformation of the ternary complex and/or a different reaction pathway may be engaged by a mismatched dNTP incorporation in DNA polymerases (3, 34, 37–40). Our steady-state and stopped-flow fluorescence studies strongly support that both correct and mismatches are incorporated via a similar mechanism, and that the fidelity of Pol β is controlled, at least partly, by destabilization of the mismatched ternary complex (higher $K_{dapp}$ values) and the chemical transition state (smaller $k_{pol}$ and $k_{quench}$ values) in the same reaction pathway (Figure 11, red profile).

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REFERENCES

1. Showalter, A. K., and Tsai, M. D. (2002) A reexamination of the nucleotide incorporation fidelity of DNA polymerases. Biochemistry 41, 10571–10576.
2. Showalter, A. K., Lamarche, B. J., Bakhtina, M., Su, M. I., Tang, K. H., and Tsai, M. D. (2006) Mechanistic comparison of high-fidelity and error-prone DNA polymerases and ligases involved in DNA repair. Chem. Rev. 106, 340–360.
3. Krahn, J. M., Beard, W. A., and Wilson, S. H. (2004) Structural insights into DNA polymerase beta deterrents for misincorporation support an induced-fit mechanism for fidelity. Structure 12, 1823–1832.
4. Johnson, S. J., and Beese, L. S. (2004) Structures of mismatch replication errors observed in a DNA polymerase. Cell 116, 803–816.
5. Trincao, J., Johnson, R. E., Wolfe, W. T., Escalante, C. R., Prakash, S., Prakash, L., and Aggarwal, A. K. (2004) Dpo4 is hindered in extending a G.T mismatch by a reverse wobble. Nat. Struct. Mol. Biol. 11, 457–462.
6. Batra, V. K., Beard, W. A., Shock, D. D., Pedersen, L. C., and Wilson, S. H. (2005) Nucleotide-induced DNA polymerase active site motions accommodating a mutagenic DNA intermediate. Structure 13, 1225–1233.
25. Bakhtina, M., Roettger, M. P., Kumar, S., and Tsai, M. D. (2007) A unified kinetic mechanism applicable to multiple DNA polymerases. Biochemistry 46, 5463–5472.

26. Bakhtina, M., Lee, S., Wang, Y., Dunlap, C., Lamarche, B., and Tsai, M. D. (2005) Use of viscosimeters, dNTPalphas, and rhodium(III) as probes in stopped-flow experiments to obtain new evidence for the mechanism of catalysis by DNA polymerase beta. Biochemistry 44, 5177–5187.

27. Tang, K. H., Niebuhr, M., Tang, C. S., Chan, C. H., Chou, C. C., and Tsai, M. D. (2008) Mismatched dNTP incorporation by DNA polymerase beta does not proceed via globally different conformational pathways. Nucleic Acids Res. 36, 2948–2957.

28. Werneburg, B. G., Ahn, J., Zhong, X., Hondal, R. J., Kraynov, V. S., and Tsai, M. D. (1996) DNA polymerase beta: pre-steady-state kinetic analysis and roles of arginine-283 in catalysis and fidelity. Biochemistry 35, 7041–7050.

29. Johnson, K. A. (1986) Rapid kinetic analysis of mecanochemical adenosinetriphosphatases. Methods Enzymol. 134, 677–705.

30. Zhong, X., Patel, S. S., Werneburg, B. G., and Tsai, M. D. (1997) DNA polymerase beta: multiple conformational changes in the mechanism of catalysis. Biochemistry 36, 11891–11900.

31. Rothwell, P. J., Mitaksov, V., and Waksman, G. (2005) Motions of the fingers subdomain of kendalq are fast and not rate limiting: implications for the molecular basis of fidelity in DNA polymerases. Mol. Cell 19, 345–355.

32. Stengel, G., Gill, J. P., Sandin, P., Wilhelmsson, L. M., Albinsson, B., Norden, B., and Millar, D. (2007) Conformational dynamics of DNA polymerase probed with a novel fluorescent DNA base analogue. Biochemistry 46, 12269–12279.

33. Luo, G., Wang, M., Raag, S. H., and Xie, X. S. (2007) Single-molecule and ensemble fluorescence assays for a functionally important conformational change in T7 DNA polymerase. Proc. Natl. Acad. Sci. U.S.A. 104, 12610–12615.

34. Tsai, Y. C., and Johnson, K. A. (2006) A new paradigm for DNA polymerase specificity. Biochemistry 45, 9675–9687.

35. Ducat, C. A., Upton, T. G., Kashemirov, B. A., Osuna, J., Oertell, K., Beard, W. A., Wilson, S. H., Flurian, J., Warshel, A., McKenna, C. E., and Goodman, M. F. (2008) DNA polymerase beta fidelity: halomethylene-modified leaving groups in pre-steady-state kinetic analysis reveal differences at the chemical transition state. Biochemistry 47, 870–879.

36. Ducat, C. A., Upton, T. G., Kashemirov, B. A., Batra, V. K., Martinek, V., Xiang, Y., Beard, W. A., Pedersen, L. C., Wilson, S. H., McKenna, C. E., Flurian, J., Warshel, A., and Goodman, M. F. (2007) Modifying the beta,gamma leaving-group bridging oxygen alters nucleotide incorporation efficiency, fidelity, and the catalytic mechanism of DNA polymerase beta. Biochemistry 46, 461–471.

37. Joyce, C. M., and Benkovic, S. J. (2004) DNA polymerase fidelity: kinetics, structure, and checkpoints. Biochemistry 43, 14317–14324.

38. Purvis, V., Grindley, N. D., and Joyce, C. M. (2003) Use of 2-aminopurine fluorescence to examine conformational changes during nucleotide incorporation by DNA polymerase I (Klenow fragment). Biochemistry 42, 10200–10211.

39. Florian, J., Goodman, M. F., and Warshel, A. (2005) Computer simulations of protein functions: searching for the molecular origin of the replication fidelity of DNA polymerases. Proc. Natl. Acad. Sci. U.S.A. 102, 6819–6824.

40. Xiang, Y., Oelschlaeger, P., Flurian, J., Goodman, M. F., and Warshel, A. (2006) Simulating the effect of DNA polymerase mutations on transition-state energetics and fidelity: evaluating amino acid group contribution and allosteric coupling for ionized residues in human pol beta. Biochemistry 45, 7036–7048.

41. Beard, W. A., Shock, D. D., Vande Berg, B. J., and Wilson, S. H. (2002) Efficiency of correct nucleotide insertion governs DNA polymerase fidelity. J. Biol. Chem. 277, 47393–47398.

42. Kraynov, V. S., Showalter, A. K., Liu, J., Zhong, X., and Tsai, M. D. (2000) DNA polymerase beta: contributions of template-positioning and dNTP triphosphate-binding residues to catalysis and fidelity. Biochemistry 39, 16008–16015.