

Kinetic Mechanism of Active Site Assembly and Chemical Catalysis of DNA Polymerase β

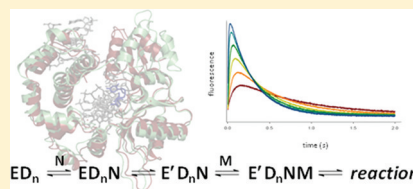
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Supporting Information

ABSTRACT: It has been inferred from structural and computational studies that the mechanism of DNA polymerases involves subtle but important discrete steps that occur between binding and recognition of the correct dNTP and chemical catalysis. These steps potentially include local conformational changes involving active site residues, reorganization of Mg^{2+} -coordinating ligands, and proton transfer. Here we address this broad issue by conducting extensive transient state kinetic analyses of DNA polymerase β (Pol β). We also performed kinetic simulations to evaluate alternative kinetic models. These studies provide some support for two-step subdomain closing and define constraints under which a kinetically significant prechemistry step can occur. To experimentally identify additional microscopic steps, we developed a stopped flow absorbance assay to measure proton formation that occurs during catalysis. These studies provide direct evidence that formation of the enzyme-bound 3'-O⁻ nucleophile is rate determining for chemistry. We additionally show that at low pH the chemical step is rate limiting for catalysis, but at high pH, a postchemistry conformational step is rate limiting due to a pH-dependent increase in the rate of nucleotidyl transfer. Finally, we performed exhaustive analyses of $[Mg^{2+}]$ and pH effects. In contrast to published studies, the results suggest an irregular pH dependence of k_{pol} , which is consistent with general base catalysis involving cooperativity between two or more protonic residues. Overall, the results represent significant advancement in the kinetic mechanism of Pol β and also reconcile some computational and experimental findings.



DNA polymerase β (Pol β), a moderate-to-high fidelity polymerase, exhibiting error frequencies of 10^{-4} – 10^{-5} ,^{1,2} is responsible for DNA synthesis in the DNA base excision repair pathway in eukaryotes.^{3,4} Specific DNA Pol β mutants with decreased fidelity have been identified in tumor cells.⁵ Moreover, Pol β is the archetypal member of the DNA Pol X family that includes DNA polymerases involved in non-homologous end joining (Pol λ , Pol σ , Pol μ), viral repair polymerases (e.g., ASFV Pol X), template-independent polymerases such as terminal deoxynucleotidyl transferase (TdT), CCA-adding enzyme, and poly(A) and poly(U) polymerases.⁶ Pol β utilizes a “two metal ion” mechanism for catalysis of nucleotidyl transfer chemistry which is conserved for all DNA and RNA polymerases.^{7,8}

On the basis of kinetic^{9–13} and structural studies^{14,15} from several laboratories, a common mechanism for DNA polymerases has been proposed (Scheme 1). This involves [1] binding of Mg -dNTP²⁻ to the enzyme–DNA binary complex to form an open ternary complex, [2] the global conformational rearrangement of subdomains to form a closed ternary complex, [3] subsequent binding of the catalytic Mg^{2+} ion, [4] the chemistry step (i.e., nucleotidyl transfer), [5] Mg^{2+} dissociation from the enzyme–product ternary complex, [6] subdomain reopening, and [7] pyrophosphate (Mg -PP_i²⁻) dissociation. [In Scheme 1, E, N, P, and M refer to Pol β (enzyme), Mg -dNTP²⁻, Mg -PP_i²⁻, and Mg^{2+} , respectively, and E' designates the enzyme in the closed subdomain

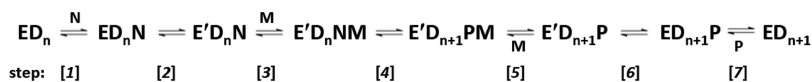
conformation.] Several studies demonstrate that, for Pol β , the open-to-closed subdomain motion [step 2] is rapid relative to the overall rate of single turnover,^{11,13,16–18} and additional kinetic studies^{19–22} support that chemistry [step 4] is rate determining.

Despite the abundance of structural and kinetic information, the chemical and biophysical basis for polymerase activity and fidelity—and of different fidelities exhibited by different polymerases—is not well understood. Several mechanistic proposals have recently been advanced based primarily on computational^{23–27} and structural^{15,28} studies. The common thread in these proposals is the purported existence of additional reaction steps, particularly conformational steps involved in organization of the active site prior to chemistry. There is strong consensus that active site assembly entails Mg^{2+} binding and consequent structural reorganization such as Mg^{2+} coordination by the 3'-OH of DNA and possibly subtle adjustment of protein side chains.²⁸ The implication is that these conformational steps may be rate determining in enzyme catalysis and that incorrect nucleotide incorporation may be characterized by altered kinetics of these steps or, alternatively, by the formation of a suboptimal active site structure.^{26,27,29} Furthermore, nucleotidyl transfer is believed to involve general

Received: June 22, 2011

Revised: September 19, 2011

Published: October 19, 2011

Scheme 1. Kinetic Mechanism of Pol β 

base catalysis, and a likely candidate for this base is the carboxylate of a conserved acid residue (D256 in Pol β), based on its proximity to the 3'-OH group, or possibly an active site bound water^{25,30,31} (Figure 1). Quantum mechanical (QM)-based

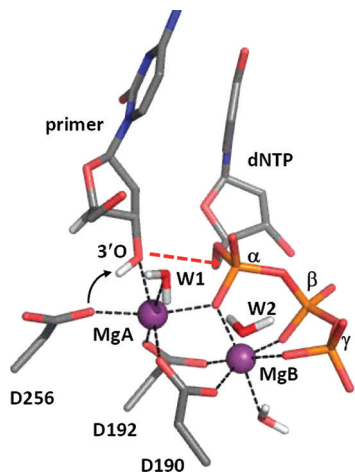


Figure 1. Active site of Pol β . The geometry of the ligands around both metals, MgA (catalytic) and MgB (dNTP-bound), is shown, including the three conserved acidic residues (D190, D192, and D256). Indicated are the hypothetical proton abstraction by D256, (black arrow), the trajectory of the 3'-O for nucleophilic attack (red dashed arrow), and metal-coordination bonds (black dashed lines). Phosphorus positions (α , β , γ) of the dNTP are labeled. Two waters are indicated: W1 is coordinated to MgA, and W2 is proximal to the triphosphate moiety of dNTP; both have been implicated in proton shuttling networks (e.g., refs 33 and 56). The model is derived from the determined structure, ref 28, of the Pol β ternary complex (pdb: 2FMS).

modeling studies^{23,24,32,33} of the Pol β active site have considered various mechanisms of proton transfer and the extent to which discrete steps—including proton abstraction from the DNA 3'-OH, nucleophilic attack by the 3'-O⁻ on the P _{α} and pyrophosphate elimination—are rate determining. It has also been proposed that correctly paired dNTPs are bound in a series of conformational steps which, together with active site assembly, constitute “kinetic checkpoints”.²⁵ To date, many of these issues have not been experimentally verified.

The purpose of this study is to evaluate alternative kinetic mechanisms of Pol β with a particular focus on possible additional kinetic steps involved in dNTP binding, correct base-pair recognition, and assembly of the active site prior to chemical catalysis. We have employed a combination of transient state kinetic analyses and kinetic simulation studies. Our kinetic results are consistent with—but do not demand—a mechanism having a rate-determining prechemistry step. However, because various models involving prechemistry pathways having alleged importance in polymerase activity and fidelity have been strongly advocated in the literature,^{24,25,28,33} we performed kinetic simulations to evaluate these hypotheses. These simulations define a range of rate constants for a putative prechemistry step that are consistent with the data, thus providing insight into possible mechanisms. In addition, the rate

of enzyme-catalyzed proton formation was directly measured using a newly developed stopped flow absorbance assay and compared with the rate of nucleotidyl transfer as measured by both stopped flow fluorescence and rapid quench. These studies provide direct evidence that formation of the nucleophile (3'-O⁻) is a rate-determining step in single nucleotide incorporation and further suggest a sequential mechanism whereby proton transfer precedes nucleotidyl transfer. Finally, our exhaustive pH- and [Mg]-dependence studies have led to new insight into the mechanism of nucleotidyl transfer.

MATERIALS AND METHODS

Materials. All chemicals were of analytical quality. Recombinant wild-type rat Pol β was expressed and purified as described.³⁴ HPLC-purified DNAs were from IDT (Coralville, IA). Stopped flow kinetic and other experiments monitoring 2AP fluorescence utilized either a 28-mer: 5'-(p)-GTGGCGAAGC-CACX_TTGCTGCGAAGCAGC-3', or 29-mer DNA: 5'-(p)-GTGGCGAAGCCACX_TTGCTGCGAAGCAGCA-3', where (p) is a 5'-phosphoryl group, T is the template deoxythymidine, and X is deoxy-2-aminopurine, designated in the text as substrate (D_{*n*}) or product (D_{*n+1*}) DNA, respectively. Stopped flow absorbance, related stopped flow kinetic (monitoring intrinsic Trp fluorescence emission), and rapid quench experiments utilized a 29-mer, 1-nucleotide gapped DNA: 5'-(p)-GGTGGCGAAGCCACCTGCTGCGAAGCAGC-3'. Some experiments, as indicated in the text, utilized a 2',3'-dideoxy-DNA (ddD_{*n*}), identical in sequence to the DNAs above. DNAs were dissolved in 10 mM KCl and stored at -20 °C. Because of their extensive secondary structure, DNA concentrations were determined by UV/vis spectroscopy after complete digestion with P1 nuclease using the ϵ_{260} calculated from their constituent deoxyribonucleotides. dAMPCPP was purchased from Jena Bioscience (Jena, Germany).

Stopped Flow Kinetic Assays. Stopped flow experiments were performed on an Applied Photophysics (Leatherhead, U.K.) SX20 stopped flow apparatus having a dead time of 1.3 ms. For 2AP fluorescence, the excitation wavelength was 312 nm, the spectral bandpass was 4 nm, and fluorescence emission was measured using a 360 nm cutoff filter (Corion). For Trp-fluorescence, the excitation wavelength was 295 nm, the spectral bandpass was 4 nm, and emission was measured using a 320 nm cutoff filter (Corion).

Stopped flow fluorescence assays were performed at 37 °C in 50 mM each bisTris/AMPSO buffer, adjusted to a specific pH (range 6.1–8.9; see text). The ionic strength of the buffer was adjusted using KCl to the equivalent of 100 mM KCl as determined by ion conductivity measurement. Buffers also contained 30.4% glycerol (w/w), 1 mM DTT, and MgCl₂ [0–20 mM], as indicated in the text. In single turnover nucleotide (dT:dATP) incorporation (i.e., nucleotidyl transfer) experiments, solution A (0.5 μ M Pol β ; all concentrations are postmixing) was combined with solution B (0.2 μ M DNA and a variable concentration of Mg-dATP²⁻ in the range 2.4–600 μ M). In our previous studies,^{19,35} Pol β kinetic experiments were performed by premixing polymerase and DNA and then initiating the reaction by addition of dNTP. Using that

method, much of the data pertaining to fast subdomain motion ($k_{\text{obs}} = 10^2\text{--}10^3 \text{ s}^{-1}$) is lost during the ~ 4 ms of instrument dead time and mixing. The method employed in this study, whereby enzyme was mixed with a solution containing both DNA and dNTP, imposes a kinetic lag phase (~ 6 ms) due to DNA binding prior to dNTP-induced subdomain motion, which enables observation of the entire kinetic phase covering the conformational step.

Sequential-mixing stopped flow experiments to measure the kinetics of the reverse pathway upon quenching of the reaction with EDTA were performed similarly as before.³⁶ In these experiments, solution A (0.5 μM Pol β) was initially mixed with solution B (0.2 μM DNA and 131 μM Mg-dATP²⁻), both with 0.2 mM Mg²⁺, and allowed to age for 15 ms, during which time transient enzyme intermediates accumulate. After this aging time, the reaction was mixed with solution C (buffer which included 20 mM EDTA and no MgCl₂).

Stopped flow absorbance assays were performed in weakly buffered solutions in the presence of pH indicator comprised of either 0.5 mM MES/0.05 mM bromocresol purple (pH 6.2) or 0.5 mM bicine/0.05 mM cresol red (pH 8.6). Solutions also included 100 mM KCl, 1 mM DTT, and 30.4% glycerol. Dissolved CO₂ was removed from stock solutions of buffer/indicator by N₂ sparging, and reaction solutions were prepared using vacuum degassed H₂O. For these measurements and the supporting intrinsic Trp fluorescence measurements, solution A (9.2 μM of Pol β , *no indicator*) was combined with solution B (8.5 μM DNA, 100 μM Mg-dATP²⁻, and indicator); the final concentration of indicator was 0.05 mM after mixing. The solutions also contained a variable concentration of MgCl₂ (indicated in the text). Proton binding by the pH indicator was monitored by absorbance at 590 nm (bromocresol purple) or 574 nm (cresol red), and proton equivalents were calculated from specific color changes determined in separate calibration experiments using standard solutions of HCl.

Rapid Chemical Quench Assays. Single turnover experiments (37 °C in bisTris/AMPSO buffer, pH 6.2 or 6.8, identical to those described above) were done using a KinTek (Austin, TX) RQF-3 instrument. Reactions were performed by combining a solution containing (final concentrations) 2 μM Pol β with a solution containing 1 μM DNA and 100 μM Mg- $[\alpha\text{-}^{32}\text{P}]\text{-dATP}^{2-}$ (>98% saturating); both solutions contained a variable concentration of Mg²⁺ (0.07–20 mM). The reactions were quenched using 0.2 M EDTA. The reaction was followed by the incorporation of $[\alpha\text{-}^{32}\text{P}]\text{-dAMP}$ in the DNA by acid precipitation. Briefly, an aliquot of each sample was transferred to a tube containing 10 μL of 50 μM carrier DNA (random oligos approximately 50–100 nucleotides in length) and 1 mg/mL BSA and precipitated with 750 μL of 15% cold trichloroacetic acid (TCA). The DNA was precipitated by centrifugation (12000g, 20 min) and then washed several times with cold 15% TCA. The precipitates were dissolved in 10 mM NaOH and 0.1% SDS, and the radioactivity was measured by liquid scintillation counting. The total radioactivity (per volume) in the original RQF sample before precipitation was also determined and used to calculate the normalized amount of radioactivity incorporated into the DNA.

Conventional Data Analysis. Nonlinear regression analysis was performed by the program gnuplot (<http://www.gnuplot.info/>). Both stopped flow and rapid quench kinetic traces were analyzed according to eq 1, where where A is the total amplitude change, t is time in seconds, λ is the rate term (which approximates the observed k_{pol}), and C is a

baseline term. [Note: For stopped flow traces, only the portion of the curve corresponding to the slow, rate-limiting portion of the reaction was analyzed. For rapid quench experiments, the normalized radioactivity incorporated, $f(t)$, divided by A is equal to the fraction of product D_{n+1} formed.]

$$f(t) = A(1 - e^{-t\lambda}) + C \quad (1)$$

To determine pH-dependent rate constants (k_{pol}) and Mg²⁺ dissociation constants (K_{d}^{Mg}), observed rate constants (k_{obs}) were analyzed as a function of $[\text{Mg}^{2+}]$ according to eq 2a or, in cases where substrate-like inhibition by Mg²⁺ was observed, eq 2b:

$$k_{\text{pol}(\text{app})} = \frac{k_{\text{pol}}[\text{Mg}]}{[\text{Mg}] + K_{\text{d}}^{\text{Mg}}} \quad (2a)$$

$$k_{\text{pol}(\text{app})} = \frac{k_{\text{pol}}[\text{Mg}]}{[\text{Mg}] + K_{\text{d}}^{\text{Mg}} + \frac{[\text{Mg}]^2}{K_{\text{II}}}} \quad (2b)$$

Global Analysis of Kinetic Data. Global kinetic analysis was performed using KinTek Explorer and FitSpace Explorer.^{37,38} The following strategy was used to collect stopped flow fluorescence data for Pol β intended for global analysis: (i) First, background fluorescence of buffer, free Pol β , and free DNA were measured under the same conditions (pH, temperature, etc.) as the kinetic assays. (ii) The kinetics of Pol β -DNA (ED_n) binary complex formation was independently measured to determine amplitudes and rate constants involved in the formation of the ED_n complex and other intermediates. (iii) Finally, stopped flow fluorescence assays to measure nucleotidyl transfer were performed. A complete description of DNA-binding kinetics and further details pertaining to analysis of kinetic data are provided as Supporting Information.

Complex Formation of Mg²⁺ with dATP and Tri(poly)-phosphate. The equilibrium concentrations of Mg-dATP²⁻ (or Mg-PPP₁²⁻) and Mg²⁺ in experiments were computed using literature values of the pK_a and Mg²⁺ dissociation constants for Mg-dATP²⁻ (28 μM) and Mg₂-dATP⁰ (25.1 mM),³⁹ and these calculations are provided in the Supporting Information. For tri(poly)phosphate, the active form is presumed to be Mg-X-PPP₁²⁻, where X = Na⁺, K⁺, or H⁺; binding of X was not considered in the calculation.

RESULTS AND ANALYSIS

Pol β Stopped Flow Fluorescence Kinetics. Pre-steady-state kinetic experiments measuring single nucleotide incorporation catalyzed by Pol β were performed at 37 °C, pH 7.1, over a range of both $[\text{Mg}^{2+}]$ and $[\text{Mg-dATP}^{2-}]$. As described previously, the stopped flow kinetics of Pol β -catalyzed nucleotidyl transfer monitored by fluorescence using either intrinsic Trp or 2AP-labeled DNA is dominated by two phases that correspond to large-scale subdomain motions that occur during catalysis.^{19,35} The rapid fluorescence increase seen at early time points corresponds to the subdomain open-to-close transition induced upon nucleotide (Mg-dNTP²⁻) binding. The subsequent decrease in fluorescence corresponds to subdomain reopening after nucleotidyl transfer (i.e., chemistry), and the rate of this step has been shown by us to be limited by chemistry under these conditions.³⁵ All the data together were analyzed globally using KinTek Explorer^{37,38} according to two alternative models for nucleotide induced subdomain motion involving either one or two conformational steps. The results

Table 1. Kinetics of dT:dATP Insertion by Pol β^a

		one-step	two-step
K_d^{dATP} ($1/K_1$)	μM	300 ^b	724 ^b
$k_{+2(a)}$	s^{-1}	1365 \pm 155	3190 ^c
$k_{-2(a)}$	s^{-1}	11.9 \pm 2.8	32.1 ^c
$k_{+2(b)}$	s^{-1}		122 ^c
$k_{-2(b)}$	s^{-1}		11.6 ^c
K_d^{Mg} ($1/K_3$)	mM	1.76 \pm 0.38	1.56 \pm 0.30
k_{+4}	s^{-1}	6.43 \pm 1.15	6.38 \pm 0.93
(ED)		1 ^b	1 ^b
(EDN)		1.83 ^d	2.26 ^d
(E'DNM)		2.47 \pm 0.03	2.46 \pm 0.01
χ^2/ndf		82.8	77.6

^aKinetic parameters were derived by global fitting to Scheme 1 or an analogous scheme having two conformational steps [2a and 2b]. Kinetic and binding parameters for steps 2–4 and fluorescence amplitude parameters (relative units) for ED, EDN, and E'DNM species are reported. Error limits were derived from contour confidence analysis (Figure 3) reflecting values within 1.2 \times the chi squared (χ^2/ndf) minimum. ^bValue fixed during global fit. ^cLower limit defined only. ^dUpper limit defined only.

from both analyses are shown in Table 1, and the best fit results to the two-step model are shown in Figure 2. Complete details describing the mechanisms used in global analysis are provided as Supporting Information. Particular aspects of the evaluated models are described below.

Kinetics of Nucleotide-Induced Subdomain Closing. The conventional, one-step conformational model is described in Scheme 1. In the two-step model, step 2 is divided into {EDN \rightleftharpoons E_iDN \rightleftharpoons E'DN}, steps 2a and 2b, respectively, where E_iDN is a partially closed intermediate complex. On the basis of the relative chi squared (χ^2/ndf) for each fit, the data were better described by the two-step ($\chi^2/\text{ndf} = 77.6$) than by the one-step subdomain closing mechanism ($\chi^2/\text{ndf} = 82.8$).

The information content of the data pertaining to subdomain motion is unfortunately limited both by the very rapid kinetics and because the proposed intermediate in the two-step mechanism, E_iDN, is not predicted to accumulate in the pre-steady state; thus, the conclusion of biphasic subdomain motion should be considered tentative. In the two-step mechanism, a very rapid (10^3 – 10^4 s^{-1}) conformational change occurs upon nucleotide binding, followed by a slower (10^1 – 10^2 s^{-1}) relaxation that results in full subdomain closing. Both models were evaluated by confidence contour analysis (Figure 3) as described by Johnson et al.,³⁸ which indicated that the one-step mechanism is well determined, while only lower limits can be determined for rates pertaining to biphasic conformational closure [steps 2a and 2b].

Kinetics of Mg·dATP²⁻ Binding. Global analysis of the one-step conformational change mechanism yielded a lower limit for K_d^{dATP} on the order of 10–100 μM and of the two-step mechanism, a K_d^{dATP} on the order of 100 μM –1 mM (Table 1). This difference can be rationalized in that the two-step mechanism explicitly accounts for very rapid and exothermic isomerization of the EDN complex [step 2a] to form the intermediate, EDN_i, whereas the one-step mechanism does not. The $K_{d(\text{app})}$ for Mg·dATP²⁻ ($\sim 2.7 \pm \mu\text{M}$), defined here as $1/(K_1K_2)$ or $1/(K_1K_{2a}K_{2b})$, is similar for both models and consistent with a value (1.2 μM) determined in independent titration experiments measuring the equilibrium binding of Mg·dATP²⁻ to form a closed ternary complex with Pol β and 2',3'-dideoxy-DNA (Figure 4A).

We reasoned that initial binding of dNTP is dominated by interactions involving the Mg–triphosphate moiety with the enzyme; this is consistent with structure–function studies of Pol β .^{15,40} We therefore performed a titration of the E·D_n complex with Mg-tri(poly)phosphate, Mg·PPP_i (Figure 4B). Interestingly, there was a small fluorescence increase upon Mg·PPP_i binding, suggesting that the E·D_n·Mg·PPP_i complex is partially closed. Analysis of the titration yielded a K_d for

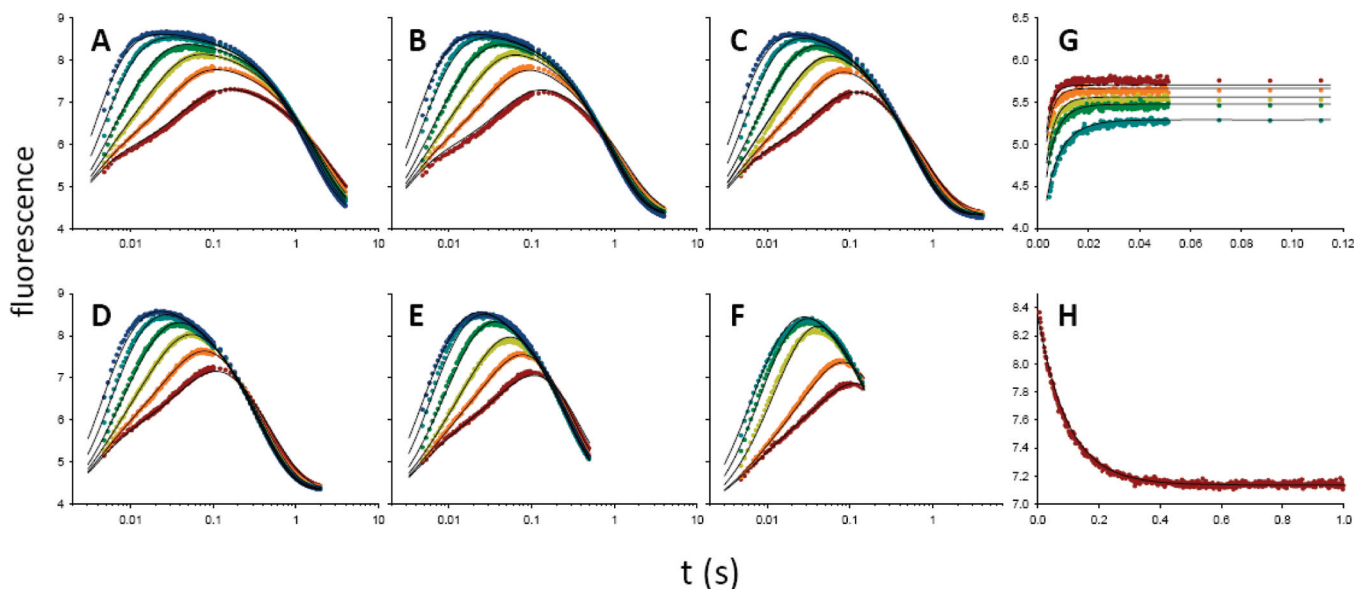


Figure 2. Global analysis of stopped flow fluorescence kinetic data. In all experiments, Pol β was 500 nM and DNA was 200 nM. (A–F) Single turnover nucleotidyl transfer reactions: dT:dATP insertion (plotted on a log time scale). The solid lines are simulated results after global fit to the mechanism involving biphasic subdomain motion. In each panel, the curves are reactions performed at different $[\text{Mg}\cdot\text{dATP}^{2-}]$ in the range of 3–350 μM (red to blue). Each panel shows reaction sets performed at a given $[\text{Mg}^{2+}]$: (A) 0.2, (B) 0.4, (C) 0.75, (D) 1.5, (E) 2.5, and (F) 5.0 mM. (G) Stopped flow kinetics of ED_n binary complex formation. The solid lines show simulated result after global fit to *only* the DNA binding portion of the mechanism. (H) Kinetics of subdomain reverse opening; $[\text{Mg}^{2+}]$ after first mix was 0.2 mM (trace after second mix with EDTA is shown).

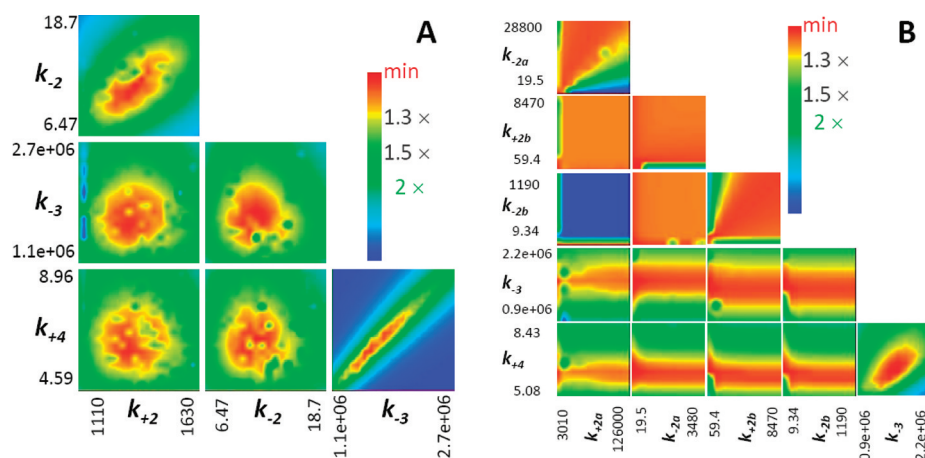


Figure 3. Confidence contour analysis. FitSpace Explorer³⁸ was used to determine upper and lower boundaries of the indicated rate constants for mechanisms involving (A) one- and (B) two-step subdomain conformational change. In this analysis, amplitude terms are held constant, each rate constant is varied and held constant in pairs, and all other rate constants are refit by global analysis. The ratio of the resulting χ^2/ndf value to that of the original best-fit solution is calculated and used to generate the contour plots; the red zone defines a region of good fit corresponding to maximum deviation of 20% ($1.2\times$) in χ^2/ndf value. For the two-step mechanism (B), confidence analysis reveals that the upper limits for the rates of subdomain conformational steps (2a and 2b) are not defined.

Mg-PPP_i of 280 μM , similar to the estimates of $K_{\text{d}}^{\text{dATP}}$ from global analysis of nucleotidyl transfer kinetic data (Table S2). The results support that weak initial nucleotide binding, primarily mediated by Mg^{2+} -triphosphate interactions, is followed by an energetically favorable conformational closing step upon recognition of a correct base pair.

Kinetics of Mg^{2+} Binding and Chemistry and Evaluation of a “Prechemistry” Step by Kinetic Simulation. The kinetic data are consistent with a mechanism involving the rapid equilibrium binding of Mg^{2+} [Scheme 1, step 3], followed by a single rate-determining step for nucleotide insertion chemistry [step 4]. Analysis using both one- and two-step conformational mechanisms (Table 1) gave essentially identical solutions for k_{pol} ($\sim 6.4 \text{ s}^{-1}$) and K_{d}^{Mg} ($\sim 1.7 \text{ mM}$). Others have proposed mechanisms that involve an additional kinetic step before chemistry.^{23,24,33} To investigate this mechanism by kinetic simulation, we deliberately constructed a more complex mechanism that includes a prechemistry step: the isomerization of E'DNM to form an activated enzyme complex, E'DNM (Figure 5A). The purpose of the simulation was to determine if the kinetic data are consistent with such a mechanism and to investigate quantitatively how the rate constants for the prechemistry and chemistry steps can vary, given the constraints imposed by the data, thus providing insight into possible mechanisms.

Here we considered a general case (Figure 5A) where step 1' is reversible and step 2' is irreversible. Only the rate constants involved in Mg^{2+} binding, steps 1' and 2', and the amplitude terms for E'DNM and E'DNM (which were constrained to be equal) were fit, and all other parameters were held constant at their best fit values. Since we are interested in mechanisms that include two rate-determining steps, we constrained the forward rate constants, $k_{1'}$ and $k_{2'}$, at various fixed ratios between 0.01 and 100 and then refit the experiment; the best-fitted parameters are reported in Figure 5B. This analysis shows there are multiple solutions to the prechemistry model that satisfy the data, indicating, as expected, that the data contain insufficient information to determine a prechemistry step. Figure 5B demonstrates that as the $k_{1'}:k_{2'}$ ratio varies between 0.1 and 10, $k_{-1'}$ is optimized so that the overall flux through

both steps can account for the observed rate of nucleotide insertion (k_{pol}). Note at the extreme ends of the x-axis two situations occur, both of which represent kinetic collapse of the mechanism to a single rate-limiting step. When $k_{1'} \ll k_{2'}$, the $K_{1'}$ equilibrium is unfavorable, but a very fast and exothermic step 2' drives the reaction forward. In this situation, the observed rate (k_{pol}) establishes a lower limit for $k_{1'}$. When $k_{1'} \gg k_{2'}$, the equilibrium steps K_{d}^{Mg} and $K_{1'}$ become linked, and the value for K_{d}^{Mg} increases to compensate for a very fast and exothermic step 1'. In this situation, the observed rate (k_{pol}) establishes a lower limit for $k_{2'}$. Interestingly, when both steps 1 and 2 are made irreversible (Supporting Information, Figure S5), the simulation indicates a poor fit when $k_{1'} \geq k_{2'}$, indicating that a mechanism with two irreversible rate-determining steps is not supported by the data. Overall, the simulations demonstrate that a reversible, nonequilibrium prechemistry step is consistent with kinetic data.

pH- and $[\text{Mg}^{2+}]$ -Dependence Studies. Proper active site assembly is a prerequisite for enzyme catalysis,⁴¹ and for DNA polymerases, this entails Mg^{2+} binding and subsequent structural reorganization around the metal, including proton transfer.^{25,27,28} The kinetics and mechanism of this process is poorly understood; thus, we performed additional kinetic studies. In a pH range of (6.2–8.9), the observed rate of nucleotidyl transfer (k_{obs}) was plotted against $[\text{Mg}^{2+}]$ to determine apparent (pH-dependent) K_{d}^{Mg} and k_{pol} values (Supporting Information, Figure S6A,B; replots for pH 6.2 and 8.6 are also shown in Figure 7C,D for comparison to rapid quench results, described below). At pH > 8, inhibition at high $[\text{Mg}^{2+}]$ was evident; this was analyzed by an uncompetitive inhibition model to extract K_{d}^{Mg} and k_{pol} values (Supporting Information, Figure S6C). Figure 6 shows the pH dependence of $\text{p}K_{\text{d}}^{\text{Mg}}$ and $\log(k_{\text{pol}})$. The $\text{p}K_{\text{d}}^{\text{Mg}}$ varies between two limiting values, with stronger binding at high pH (Figure 6A), consistent with a one-proton binding/dissociation event. The S-shape of the curve indicates that Mg^{2+} is capable of binding to a protonated form of the active site with lower affinity. Therefore, the data were analyzed according to eq 3. Limiting values (Y_{limit}) for K_{d}^{Mg} are 37.5 and 0.32 mM at low and high pH conditions, respectively. The macroscopic $\text{p}K_{\text{a}}$ value of 7.13

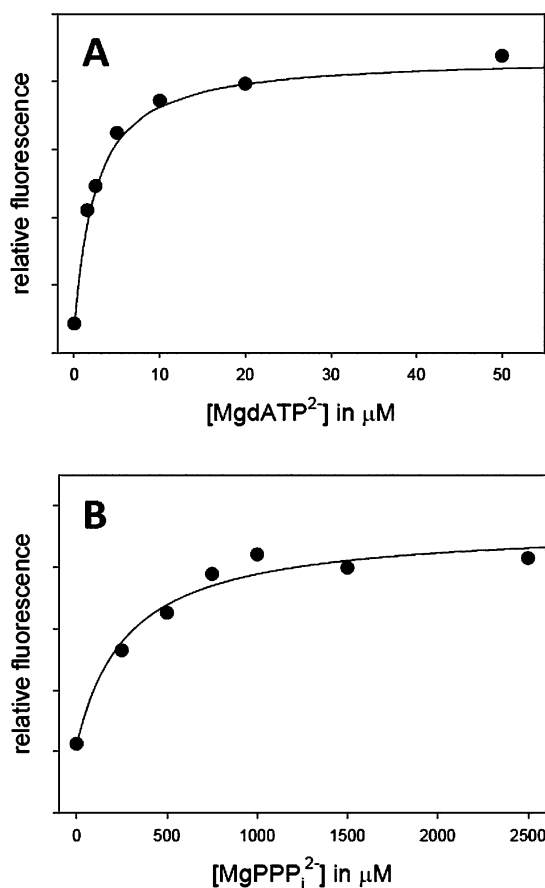


Figure 4. Equilibrium binding of Mg:dATP²⁻ and Mg-tri(poly)phosphate to the Pol β–DNA complex. Conditions were pH 7.1, 0.2 mM MgCl₂, 37 °C. Pol β and DNA concentrations were 500 and 200 nM, respectively. The solid curve shows the best fit to the binding equation: $f(x) = a[L]/(K_d + [L]) + bkg$, where a is the amplitude change, L is the ligand (Mg:dATP²⁻ or Mg-tri(poly)phosphate), K_d is an apparent binding constant, and bkg is the fluorescence of the enzyme–DNA complex without added ligand. Solutions for K_d were (A) 1.2 μM (Mg:dATP²⁻) and (B) 281 μM (Mg-tri(poly)phosphate).

(Figure 6A) corresponds to ionization of E'DN (i.e., the enzyme form that binds Mg²⁺).

$$pK_d^{Mg} = \frac{Y_{\text{limit}1} + (Y_{\text{limit}2} \times 10^{pK_a - \text{pH}})}{1 + 10^{pK_a - \text{pH}}} \quad (3)$$

Our data demonstrate an increase in k_{pol} at high pH (Figure 6B), similar to previous pH dependence studies of Pol β.^{20,22} This is consistent with general base catalysis involved in the formation of the 3'-O⁻ nucleophile on the enzyme. However, the pH dependence of $\log(k_{\text{pol}})$ does not conform to a simple Henderson–Hasselbalch description, but instead increases gradually over at least 3 pH units with a Hill parameter of 0.47 (slope in the log–log plot, Figure 6B). This type of irregular pH titration is often an indication of an electrostatic interaction (cooperativity) between two or more protonic groups involved in catalysis.^{42,43} Alternatively, as Cleland notes in his classical work,⁴⁴ a flat pH–rate profile could result if proton dissociation from the Michaelis complex [$\{E'D_n\text{NM}\cdot\text{H}^+ \rightleftharpoons E'D_n\text{NM} + \text{H}^+\}$; $pK_a = 5.56$, calculated from the pK_a of E'D_nN (7.13) and the two limiting K_d^{Mg} values, Figure 6A] is slow relative to substrate dissociation and the central step (chemistry). Because of the limited pH range accessible by

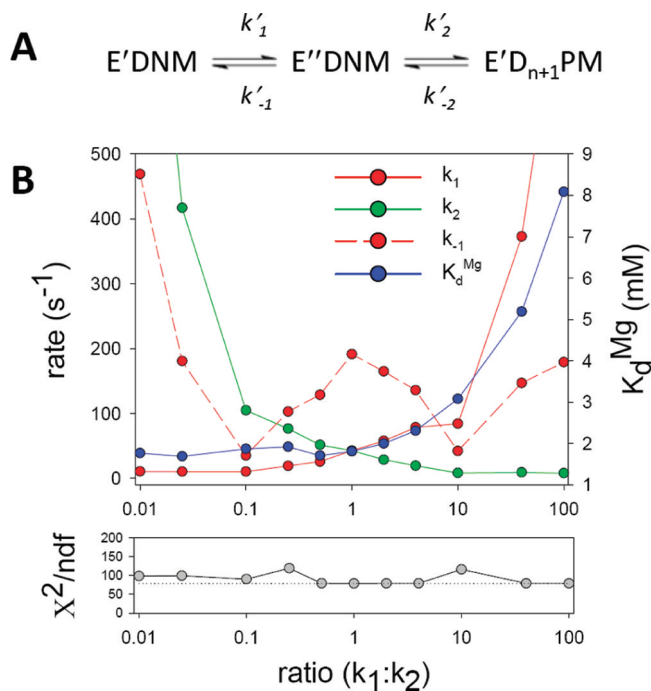


Figure 5. Kinetic simulation of a hypothetical prechemistry step. (A) Simulated kinetic model involving prechemistry isomerization [1'] and chemistry [2'] steps. (B) The fitted parameters k_1' , k_{-1}' , k_2' , and K_d^{Mg} from multiple global fits of the data (same data set as in Figure 2) and the corresponding χ^2/ndf values are plotted versus the ratios of $k_1':k_{-1}'$, as described in the text; k_{-2}' was held constant at zero. The dotted reference line in (B) indicates the minimum χ^2/ndf from the original data fit.

experiment, as well as the irregular pH–rate profile, the maximum k_{pol} and the associated apparent pK_a value could not be defined. This conclusion differs substantially from previous studies,^{20,22} although the general pH trend is consistent (see Supporting Information).

To corroborate the results obtained by the stopped flow experiments (above) and to further investigate the observed uncompetitive inhibition by Mg²⁺, additional rapid quench experiments were performed to directly measure the rate of nucleotidyl transfer as a function of $[\text{Mg}^{2+}]$ at pH 6.2 and 8.6. Figure 7A,B shows the time courses of product D_{n+1} formation at pH 6.2 and 8.6, respectively, and these data were analyzed by nonlinear regression (eq 1) to obtain observed rate constants (k_{obs}). The replots (k_{obs} versus $[\text{Mg}^{2+}]$) are shown in Figure 7C,D and compared with those from stopped flow studies (described above) performed under identical pH conditions. At pH 6.2 (Figure 7C), there is excellent agreement between the results obtained by the two methods, supporting the previous conclusion²⁰ that chemistry is overall rate limiting at low pH. At pH 8.6 (Figure 7D), the inhibition observed at high $[\text{Mg}^{2+}]$ in the stopped flow experiment is not observed in the rapid quench experiment. This indicates that the inhibition is not due to enzyme inactivation, but instead due to the decrease in rate of a step after chemistry. Note the results obtained by analysis of the stopped flow data accurately predict the results of the rapid quench experiment—absent uncompetitive inhibition (Figure 7D, dotted line). This difference seen in k_{pol} values obtained by the two assay methods at high pH has previously been attributed to a pH-dependent increase in the rate of chemistry such that a later step (most likely subdomain

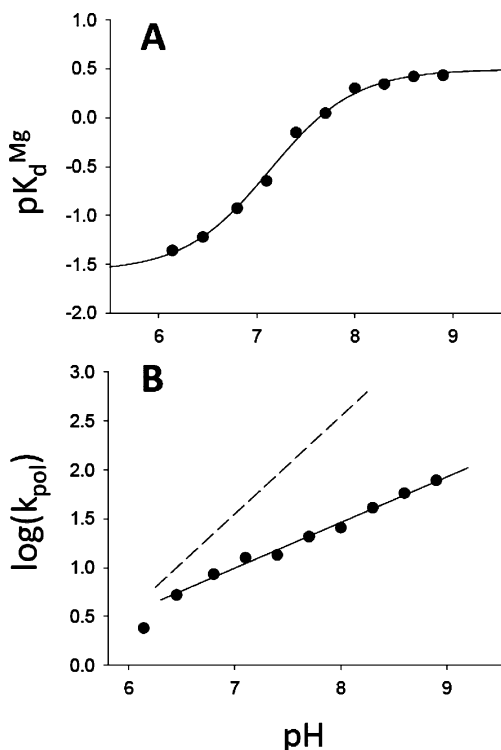


Figure 6. pH dependence of kinetic parameters of Pol β . (A) Variance of pK_d^{Mg} (K_d in mM) as a function of pH. (B) Variance of $\log(k_{pol})$ (k_{pol} in s^{-1}) with pH. The solid line shows the linear regression fit of the data between pH 6.4 and 8.9. The slope defines the Hill parameter (0.47); an arbitrary line (dashed) with slope = 1 is shown for reference.

opening after chemistry) becomes rate limiting.²⁰ According to Scheme 1, subdomain opening after chemistry [step 6] occurs after Mg^{2+} dissociation from the enzyme [step 5]; thus, we conclude that Mg^{2+} inhibition is attributed to accumulation of the intermediate, $E'D_{n+1}PM$, resulting in a decrease in the rate of flux through these two steps.

Stopped Flow Absorption Kinetics of Enzymatic Proton Generation. The following experiments were performed to measure the kinetics and stoichiometry of proton release during enzyme catalysis. The formation of protons was detected by absorption change of the pH indicator present in the reaction solution. In Figure 8, the absorbance data are superimposed by data from stopped flow experiments measuring intrinsic Trp fluorescence performed under identical conditions. The results show that at pH 6.2 (Figure 8A) the rate of H^+ formation is coincidental with k_{pol} ($1.0 \pm 0.1 s^{-1}$). Under this condition, subdomain opening [step 6], which is reported by the change in Trp fluorescence, is fast relative to chemistry. At higher pH, the rate of the chemical step increases. At pH 8.6 (Figure 8B), the rate of H^+ formation ($70 s^{-1}$) is approximately 4-fold faster than the Trp fluorescence change ($17 s^{-1}$). It is remarkable that at high pH proton release occurs prior to subdomain opening [step 6]. This result indicates, in accordance with microscopic reversibility, that the reverse reaction (pyrophosphorolysis) can occur by binding of a proton to the closed $E'D_{n+1}PM$ complex.

The rate of proton formation corresponds either to formation of the $3'-O^-$ nucleophile in the active site or, alternatively, to deprotonation of the catalytic base prior to proton abstraction from $3'-OH$. It is interesting to compare this

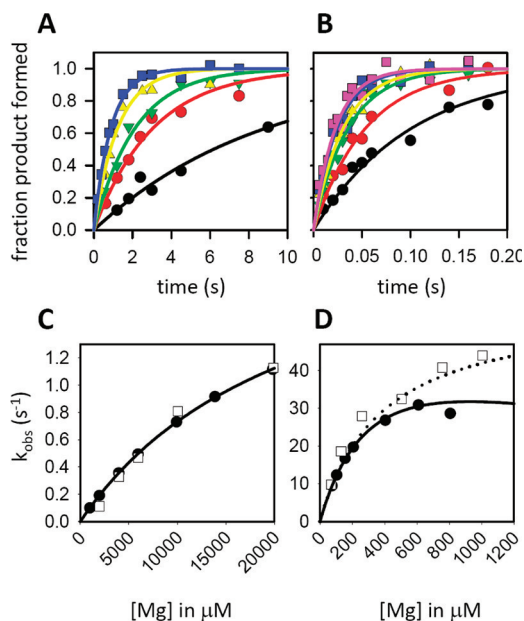


Figure 7. Rapid quench kinetics of Pol β and comparison to stopped flow results. Reaction time courses are shown for experiments performed at (A) pH 6.2, $[Mg^{2+}] =$ (ascending order) 1990, 4003, 5980, 10 060, and 19 920 μM and (B) pH 8.6, $[Mg^{2+}] =$ 70, 132, 261, 506, 757, and 1006 μM . All experiments were performed at 37 $^{\circ}C$. The solid curves (A, B) are individual fits of each data set to eq 1, and the replots of k_{obs} as a function of $[Mg^{2+}]$ are shown in (C) pH 6.2 and (D) pH 8.6; the results from stopped flow (\bullet) and rapid quench (\square) experiments are plotted together for comparison. The solid curves (C, D) are best fits of the stopped flow data to eqs 2a and 2b, respectively. At pH 6.2 (C), $k_{pol} = 2.41 s^{-1}$ and $K_d^{Mg} = 22.8 mM$. At pH 8.6 (D), $k_{pol} = 57.86 s^{-1}$, $K_d^{Mg} = 0.38 mM$, and K_{II}^{Mg} (uncompetitive inhibition constant) = 2.25 mM; the dotted curve is the same solution, only in the absence of Mg inhibition (eq 2a).

rate ($70 s^{-1}$) at pH 8.6 with the rate of nucleotidyl transfer (dotted line, Figure 7D) at this condition ($\sim 30 s^{-1}$). These data suggest that deprotonation of the $3'-OH$ is a rate-determining step in the overall nucleotide insertion reaction. It is often assumed that deprotonation of the $3'-OH$ occurs as a concerted step with nucleotidyl transfer. However, our data suggest an alternative sequential mechanism involving two discrete *rate determining* steps—here, (i) $3'-OH$ deprotonation and (ii) nucleotidyl transfer. This mechanism predicts, similar to the results presented here (Figures 8A and 7D), that the *observed rate* of proton formation would be faster than that of the final product (D_{n+1}), given identical rates of the two microscopic rate steps (kinetic simulation, Figure S7; also see ref 45).

DISCUSSION

We combined pre-steady-state kinetics, $[Mg]$ - and pH-variation studies, and kinetic simulation in an effort to characterize transient enzyme intermediates important in the Pol β reaction pathway. Our particular interest is in possible conformational steps that constitute prechemistry organization of the active site, including local conformational steps related to Mg binding and proton transfer. We report several major conclusions. First, we show that the kinetic mechanism of Pol β is adequately described with a single rate-limiting step corresponding to nucleotidyl transfer but that a rate-determining prechemistry step *is possible* given the constraints defined in our simulation

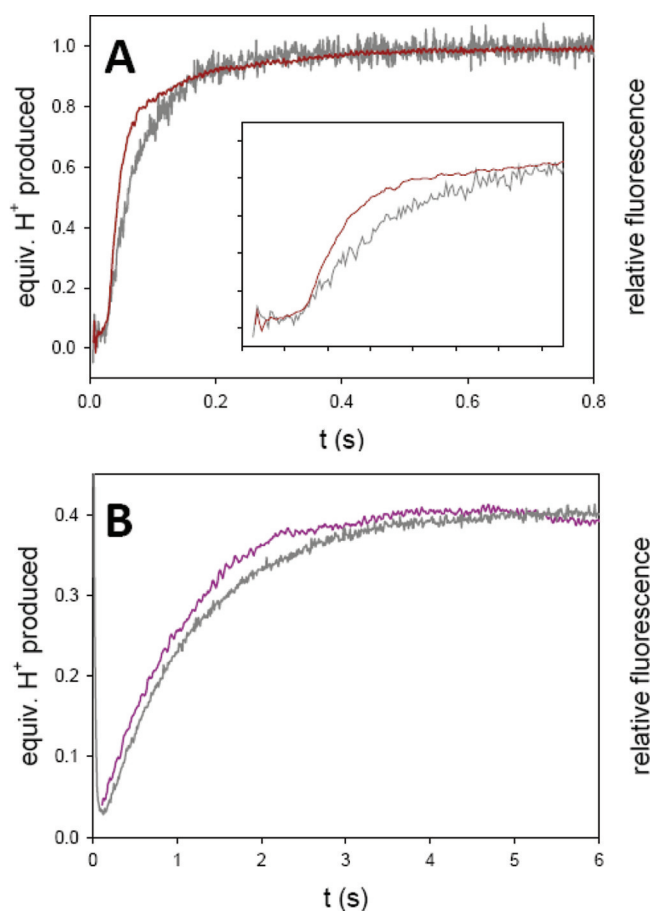


Figure 8. Stopped flow absorbance kinetics of enzymatic H⁺-formation. Equivalents of H⁺ produced by Pol β , colored, were determined from stopped flow absorbance traces. The stopped flow (intrinsic Trp-) fluorescence traces, gray, were normalized and scaled for superimposition. (A) Bicine, cresol red, pH 8.6, 0.5 mM MgCl₂; the Trp-fluorescence emission trace exhibits a single-component exponential change in fluorescence ($\lambda = 17.4 \text{ s}^{-1}$) following a ~ 20 ms lag time that corresponds to formation of a closed EDN ternary complex. The H⁺-formation kinetic plot, red, is biphasic, dominated by a large amplitude phase ($\lambda = 70.5 \text{ s}^{-1}$). A replot between 0 and 0.15 s is shown in the inset. (B) MES, bromocresol purple, pH 6.2, 20 mM MgCl₂; both Trp-fluorescence and H⁺-formation plots exhibit single-component exponential kinetics with similar rates ($\lambda = 0.90$ and 1.10 s^{-1} , respectively). The observation of less than 1 equiv of protons formed can be explained by either formation of H·Mg·PP₁⁻ or by net proton binding to the product (E·D_{n+1}) enzyme species.

study. We provide evidence to support that proton abstraction from the 3'-OH is rate determining for single nucleotide incorporation. A recent computational study proposed that proton transfer occurs concertedly with rate limiting nucleophilic attack.²⁵ Our observation that proton release occurs faster than nucleotidyl transfer (at pH 8.6) seems to be in conflict with a strictly concerted mechanism. Lin et al.²⁴ alternatively proposed that the deprotonation of the 3'-OH occurs as an equilibrium step, prior to the formation of a transient enzyme intermediate in which the Mg-coordinated 3'-O⁻ is poised for nucleophilic attack on the α P. Our data are consistent with this mechanism but further suggest that the rate of proton transfer from 3'-OH is similar to that of nucleotidyl transfer.

It has long been understood that binding of a correct dNTP induces a conformational change in DNA polymerases that

activates the enzyme for nucleotidyl transfer.^{7,14} Here we present experimental evidence in support of two discrete steps in the nucleotide-induced subdomain closing of Pol β . Our data, of course, do not address the structural mechanism of this conformational change. Previously, however, computational studies from the Schlick laboratory²³ suggested that subdomain closing occurs by a biphasic process involving the sequential rearrangement of side chains D192, R258, and F272. It was proposed that a steric clash between R258 and F272 delays the enzyme in a partially open conformation. Mutation studies with R258A Pol β indicated that R258 does not impede the forward reaction rate but that subdomain opening in the reverse direction was faster for the mutant.^{20,36} This is consistent with R258 forming a stabilizing interaction in the closed ternary complex.

It has been proposed that having multiple kinetic steps prior to chemistry could affect polymerase fidelity by imposing so-called "kinetic checkpoints".^{46–48} These proposed steps include both nucleotide-induced global conformational changes and more subtle conformational changes associated with the preparation of the active site for chemical catalysis. While structural⁴⁹ and computational studies^{23,50} have given insight into the molecular mechanisms for coupling substrate binding and recognition with enzyme activation, the impact of these individual steps on the kinetics of correct versus incorrect dNTP incorporation (and, hence, fidelity) is controversial. It has been pointed out that multiple prechemistry steps would not be expected to affect fidelity, provided that these steps occur faster than the rate-limiting chemical step.^{26,51} Our data are consistent with this idea, showing that formation of an active E·DNM complex (for Pol β) occurs essentially as a rapid equilibrium step. Moreover, binding of an incorrect dNTP is known to destabilize the closed enzyme conformation leading to infrequent formation or malformation of the active site, resulting in inefficient catalysis. Together, these data support that ground-state dNTP binding and equilibration of subsequent conformational steps are important for polymerase fidelity: for correct dNTP binding, these steps are favorable, allowing for efficient catalysis, whereas for incorrect dNTP binding, these steps are unfavorable, allowing for facile dNTP dissociation rather than chemistry, consistent with a large body of structural^{14,15,28,49} and experimental^{29,52–54} studies.

The mechanism of general base and general acid catalysis in enzyme-catalyzed nucleotidyl transfer is a subject of intense interest. Proton migration within the active site would logically be expected to stabilize negative charge that develops at the transition state of nucleotidyl transfer.⁸ The pH dependence of k_{pol} reported here supports a role for general base catalysis. In Pol β , the same residue—D256—has been implicated in both ground-state Mg²⁺-binding and general base catalysis, and the observed pH dependence of $\text{p}K_{\text{d}}^{\text{Mg}}$ and k_{pol} likely reflects ionization of this residue in the absence or presence of the bound Mg²⁺ ion, respectively. The pH dependence of k_{pol} is irregular, implicating multiple protonic groups involved in catalysis, and thus could reflect ionization of a separate group, perhaps an intervening species involved in proton transfer between the 3'-OH of DNA and bulk solvent (Figure 1). In general, the irregular pH dependence of an enzyme activity suggests electrostatic interaction between multiple protonic sites, a phenomenon which is now understood to be common for enzymes.^{42,43,55} Electrostatic interactions in proteins can occur either directly (via Coulombic interactions) or by conformational effects if the ionization of a residue structurally

alters the microenvironment (and thus the microscopic pK_a) of another group. In summary, our kinetic data in total support that deprotonation of the 3'-OH is a rate-determining step in the Pol β reaction. We propose this proton is transferred to other groups in the active site, thus providing important charge stabilization during catalysis. Finally, proton dissociation from the enzyme can occur prior to—and is possibly the event that triggers—subdomain opening [step 6] prior to release of products.

Our presented evidence for the involvement of multiple protonic groups in proton transfer and general base catalysis is in accordance with a number of recent studies. For example, independent computational studies^{33,56} have proposed mechanisms that involve multiple, discrete proton transfer events between the 3'-OH, catalytic protein residues, and active site waters that are coupled to nucleotidyl transfer. Experimental studies of Pol β by Sucato et al.^{21,22} provided evidence that pyrophosphate formation was rate determining for chemistry, and Castro et al.^{57,58} concluded that multiple proton transfer events were involved in catalysis based on solvent deuterium kinetic isotope effects of several polymerases (not including Pol β). Our proposed model of electrostatic cooperativity provides a similar, but differently nuanced, explanation for their proton inventory study. Castro et al.^{57,58} proposed that general acid catalysis was required to promote pyrophosphate elimination based on their evidence for a second protonic group and their pH-dependence studies. Our pH-dependence studies are consistent with proton transfer between the 3'-OH and multiple active site groups, including to the pyrophosphate leaving group. Our kinetic data are also consistent with a rate-determining step corresponding to pyrophosphate elimination and/or subsequent active site disassembly. We caution that there are substantial differences among residues that line the dNTP binding pockets of the active sites of DNA polymerases in general and between Family X and other DNA polymerase families in particular, which suggests the possibility of alternative pathways for proton transfer in different systems.

■ ASSOCIATED CONTENT

● Supporting Information

Tables S1 and S2 providing complete solutions from global analysis to the kinetic mechanisms for ED binary complex formation and nucleotidyl transfer, respectively; Table S3 giving calculated concentrations of free Mg^{2+} and $Mg \cdot dATP^{2-}$ used in stopped flow studies; Figure S1 describing the exact kinetic models used in global analysis; Figures S2–S4 describing experiments characterizing the kinetics and mechanism of ED binary complex formation; Figure S5 describing additional kinetic simulations of the prechemistry step mechanism; Figure S6 describing details of the pH- and $[Mg^{2+}]$ -dependence study, the results of which are reported in Figure 6; Figure S7 describing kinetic simulation of sequential proton transfer and nucleotidyl transfer steps; additional analytical methods. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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Funding

This work was supported by funds from Academia Sinica.

■ ACKNOWLEDGMENTS

We thank Dr. Marina Bakhtina for critical reading of the manuscript. We also thank New England BioLabs (Ipswich, MA) for use of their RQF3 rapid quench instrument.

■ ABBREVIATIONS

Pol, DNA polymerase; ASFV, African swine fever virus; PP_i , inorganic pyrophosphate; PPP_i , inorganic tri(poly)phosphate; bisTris, 1,3-propanediol, 2-(bis(2-hydroxyethyl)amino)-2-(hydroxymethyl); AMPSO, *N*-(1,1-dimethyl-2-hydroxyethyl)-3-amino-2-hydroxypropanesulfonic acid; MES, 2-(*N*-morpholino)ethanesulfonic acid; TCA, trichloroacetic acid; dAMPCPP, 2'-deoxyriboadenosine-5'-[α,β -methylene] triphosphate.

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