Fluorescence of 2-aminopurine reveals rapid conformational changes in the RB69 DNA polymerase-primer/template complexes upon binding and incorporation of matched deoxynucleoside triphosphates

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ABSTRACT

We have used 2-aminopurine (2AP) as a fluorescent probe in the template strand of a 13/20mer primer/template (D) to detect deoxynucleoside triphosphates (N)-dependent conformational changes exhibited by RB69 DNA polymerase (ED) complexes. The rates and amplitudes of fluorescence quenching depend hyperbolically on the [dTTP] when a dideoxy-primer/template (ddP/T) with 2AP as the templating base (n position) is used. No detectable fluorescence changes occur when a ddP/T with 2AP positioned 5' to the templating base (n + 1 position) is used. With a deoxy-primer/template (dP/T) with 2AP in the n position, a rapid fluorescence quenching occurs within 2 ms, followed by a second, slower fluorescence quenching with a rate constant similar to base incorporation as determined by chemical quench. With a dP/T having 2AP in the n + 1 position, there is a [dNTP]-dependent fluorescence enhancement that occurs at a rate comparable to dNMP incorporation. Collectively, the results favor a minimal kinetic scheme in which population of two distinct biochemical states of the ternary EDN complex precedes the nucleotidyl transfer reaction. Observed differences between dP/T and ddP/T ternary complexes indicate that the 3' hydroxyl group of the primer plays a critical role in determining the rate constants of transitions that lead to strong deoxynucleoside triphosphate binding prior to chemistry.

INTRODUCTION

A general feature of replicative DNA polymerases, such as RB69 pol, is their ability to select a dNTP complementary to the templating base with great accuracy during primer extension. The frequency of misincorporation errors rarely exceed $10^{-6}$ per replicated base pair because incorporated mispaired bases are usually excised by the polymerase itself or by a separate replicase subunit having 3'-5' exonuclease activity (1–3). DNA polymerases also have to move rapidly and processively along the template strand as they catalyze the formation of phosphodiester bonds (4–6). This movement during leading strand synthesis is facilitated by DNA helicases that catalyze unwinding of the duplex DNA (7–9); single-strand DNA-binding proteins that keep the template-strand accessible to the polymerase and a doughnut-shaped trimeric sliding clamp with a central cavity diameter that can accommodate dsDNA and a groove on the outer surface which provides an interface for interaction with the C-terminal tail of the polymerase (10–14). The arrangement of these proteins in relation to each other and to the template ensures that the polymerase and the sliding clamp are tethered to the DNA until an appropriate signal triggers their disengagement (15–17).

Although enormous strides have been made during the last decade in understanding how DNA polymerases function in DNA replication and repair, there are still questions that remain about certain aspects of the mechanisms used by DNA polymerases to achieve fidelity. Among them is the relationship between base discrimination and the rate-limiting step in the nucleotidyl transfer reaction (18–20). Since most DNA polymerases share...
common structural motifs as well as basic mechanisms for primer extension and nucleotide excision, it has been a general belief that a kinetic scheme determined for a well-characterized DNA pol would be universally applicable to all DNA polymerases (21–23). More recent data suggests that subtle differences in the rate and equilibrium constants of the enzymatic cycles among DNA polymerases confer specific catalytic activities that are important for their specialized tasks in the cell (21).

To gain a complete understanding of the role that nucleotide-linked conformational changes play in base discrimination and in the mechanism of nucleotidyl transfer, it is necessary to distinguish biochemical intermediates that are populated before nucleotidyl transfer from states that follow primer extension. If the prechemical transition acts as a check point for base discrimination as has been proposed (18), the conformational changes that accompany the dNTP binding-induced transition from the open to the closed state is predicted to be rate limiting for nucleoside triphosphate incorporation. This expectation appears to have been fulfilled for T7 DNA pol and the Klenow fragment (20), which are members of the A family polymerases, but Klenaq,1, also an A family polymerase, is an exception since it has been shown that the transition from the open to the closed state is much faster than chemistry (24). Chemistry appears to be rate limiting for pol β (19), however, it is not clear which mechanism applies to members of the B family which includes T4 and RB69 pol (20). To address this issue we have used 2-aminopurine (2AP) as a fluorescent probe together with a nonextendable primer/template (ddP/T) which has allowed us to estimate both the rates of domain closing and opening in the presence of an incoming dNTP.

The nucleotide analog, 2AP, does not change the standard B-type conformation when it is part of a DNA duplex and is therefore a good adenosine analog (25). 2AP has been widely used as a fluorescent probe because its spectral properties respond to changes in its immediate environment that occur during nucleotidyl transfer. 2AP fluorescence is quenched when it stacks against adjacent bases or aromatic residues in proteins (25,26). Fluorescence enhancement occurs when stacking of 2AP with aromatic residues is perturbed (27,28). The fluorescence of 2AP is insensitive to base-pairing or other hydrogen bonding interactions (29). Furthermore, fluctuations in stacking of 2AP with an adjacent base does not interfere with the standard Watson–Crick hydrogen bonds between 2AP and thymidine (28). Thus, 2AP can be used as the templating base (n position) and changes in its fluorescence, when deoxynucleoside triphosphate (dTTP) is added to a DNA pol:ddP/T complex, allow monitoring of conformational alterations in the complex independent of fluorescence changes that arise as a consequence of chemistry. 2AP can also be placed just 5’ to the templating base (n + 1 position) so that conformational changes in the environment around the 2AP can potentially be monitored by fluorescence change in RB69:ddP/T complexes with any dNTP as long as it is complementary to the templating base (26–28). This extends the range of this approach since the incoming dNTP does not have to be restricted to dTTP as is the case when 2AP is in the n position.

In this report, we have determined the kinetic parameters for nucleotide binding to the RB69 DNA polymerase primer/template complex and subsequent base incorporation by monitoring the nucleotide-dependent fluorescence changes of 2AP that has been incorporated into the n or n + 1 position of a primer/template. To distinguish conformational changes preceding chemistry from rates of phosphoryl transfer and subsequent steps, we have employed a ddP/T (with 2AP in the n position) that is chemically inert with respect to nucleotidyl transfer reaction (26,30–35).

We selected RB69 pol, a member of B family DNA polymerases, because crystal structures have been determined for the apo enzyme (36) as well as for complexes of the enzyme with ddP/Ts and complementary dNTPs (37–40). Our results provide evidence for the existence of a transient conformational state of a RB69 pol:ddP/T:dTTP complex (2AP in the n position) that is populated after dTTP binding but before chemistry. We have evaluated the relative contribution of the rate of this conformational change to the rate-limiting step for incorporation of a correct base. We find no evidence for the existence of three distinct intermediates populated before the chemical step as proposed for T4 DNA polymerase (35), a close relative of RB69 pol. Rather, the experimental data is reliably accounted for by a mechanism in which two distinct biochemical states of the ternary complex, one that binds incoming nucleotide weakly and one that binds strongly, precedes nucleotidyl transfer. This information is crucial for understanding RB69 fidelity at the base selection stage.

EXPERIMENTAL PROCEDURES

Reagents

Escherichia coli strain BL21(DE3) was obtained from Stratagene Corp. DH5α cells were from Invitrogen. dNTPs, and T4 polynucleotide kinase were purchased from New England Biolabs; γ-[32P]-ATP was from Perkin Elmer Life Sciences Inc; and Ni-NTA resin was obtained from Qiagen. Other chemicals were analytical grade.

RB69 pol cDNA was a generous gift from J. Karam (Tulane University). Oligonucleotides were provided by W. M. Keck Foundation Biotechnology Resource Laboratory (Yale University). The sequences of the DNA primer/template substrates (P/Ts) used in this study are shown in Figure 1. The fluorescent nucleotide analog 2AP is in the n or n + 1 position of the template.

Protein expression and purification

Site-directed mutagenesis was used to substitute Ala for Asp at residues 222 and 327 to create the exonuclease-deficient (exo−) RB69 pol which we refer to throughout the article as RB69 pol. Expression of the (exo−) RB69 pol was carried out as previously described (41). For ease in subsequent isolation, the cDNA for RB69 pol was subcloned into the pSP72 vector (Promega) so that the expressed protein would have six histidine residues.
appended to its C-terminus. This permitted facile purification of the (exo-) RB69 pol on a Ni-NTA column (42). For the final purification RB69 pol was loaded on a Source Q column (Pharmacia) and eluted with a linear gradient of increasing sodium chloride. Purified RB69 pol migrated as a single band on SDS-PAGE, was free of imidazole as judged by a 280/260 nm ratio of 1.91 and had pre-steady-state kinetic parameters for the pol reaction that corresponded closely to those obtained by Capson et al. (43) for T4 pol. After elution from the Source Q column, RB69 pol was dialyzed against pol storage buffer (20 mM Tris–HCl, pH 7.5; 0.1 mM EDTA; 5 mM 2-mercaptoethanol; 25% glycerol) and stored in small buffer (20 mM Tris–HCl, pH 7.5; 0.1 mM EDTA; 5 mM imidazole as judged by a 280/260 nm ratio of 1.91 and had pre-steady-state kinetic parameters for the pol reaction that corresponded closely to those obtained by Capson et al. (43) for T4 pol. After elution from the Source Q column, RB69 pol was dialyzed against pol storage buffer (20 mM Tris–HCl, pH 7.5; 0.1 mM EDTA; 5 mM 2-mercaptoethanol; 25% glycerol) and stored in small aliquots at –80°C. Protein concentrations were determined by the Bradford assay and by absorbance at 280 nm using prepared standards.

**Equilibrium binding titrations**

Fluorescence emission spectra of the dD/P/T (2AP in the n position) (200 nm) together with RB69 pol (1 or 2 μM), 10 mM MgCl₂, 66 mM Tris–HCl, pH 7.5, 0.5 mM EDTA and varying [dTTP] were acquired at 25°C with a Photon Technology International Alphascan scanning spectrometer. Samples were excited at 310 nm to minimize inner filter effects. Fluorescence emission spectra were collected from 320 to 460 nm and the intensities were corrected for the intrinsic fluorescence of RB69 pol. Peak intensities at 365 nm were plotted as a function of [dTTP] and fit to Equation (1) to obtain the overall dissociation equilibrium constants ($K_{d}^{overall}$) for dNTP binding:

$$FI = F_{I0} + \frac{(F_{I∞} - F_{I0})}{K_{d}^{overall} + [dTTP]}$$

where $FI$ is the observed fluorescence intensity, $F_{I0}$ is the fluorescence in the absence of dTTP, $F_{I∞}$ is the fluorescence at saturating dTTP and $K_{d}^{overall}$ is the overall dissociation constant for dTTP binding. This expression assumes binding to a single site. The same procedure was used for the dD/P/T with 2AP in the n + 1 position.

**Kinetic measurements and analysis**

Transient kinetic fluorescence experiments were performed using an Applied Photophysics SX18MV-R stopped-flow apparatus (Leatherhead, UK) thermostatted at the indicated temperatures ±0.1°C. The excitation wavelength for 2AP was 313 nm. Fluorescence emission was monitored at 90° through a 345 nm long pass colored glass filter. The final concentrations of the reaction components after mixing were 66 mM Tris–HCl (pH 7.5), 200 nM dP/T or dD/P/T, 1 μM RB69 pol, 10 mM MgCl₂ and varying concentrations of dNTPs. For some experiments the final concentrations of RB69 pol and the P/T were 2 μM and 400 nM, respectively.

Time courses of fluorescence changes were fitted to a sum of exponentials [Equation. (2)] using Pro-K software provided with the instrument or with Kaleidagraph (Synergy Software, Reading, PA, USA):

$$F(t) = F_{∞} + \sum_{i=1}^{n} A_{i}e^{-k_{i}t}$$

where $F(t)$ is the fluorescence at time $t$, $F_{∞}$ is the final fluorescence intensity at equilibrium, $A_{i}$ is the amplitude and $k_{i}$ is the observed pseudo-first order rate constant characterizing the $i$th relaxation process and $n$ is the total number of observed relaxations. The value of $n$ was one (single exponential) or two (double exponential). Most time courses shown are the average of five to seven traces.

**Chemical quench experiments**

All chemical quench reactions were performed using a KinTek RQF-3 quench flow instrument (KinTek Corp., Austin, TX, USA). Single turnover experiments were carried out at 22 or 4°C. The complex of 5’-[32P] labeled primer/template (dP/T) and RB69 pol was mixed with various concentrations of dNTP. The final concentrations after mixing were 200 nM dP/T, 1 μM RB69 pol, 10 mM MgCl₂ and the indicated [dNTP]. Reactions were quenched with 0.5 M EDTA. The disappearance of substrate and the formation of product were monitored after gel electrophoresis (20% polyacrylamide/50% urea) by phosphorimager and quantified using NIH imaging software. Time courses for product formation were fitted to single exponentials. The [dNTP] dependence of the observed rate constant ($k_{obs}$) was fitted to a rectangular hyperbola [Equation. (3)] where the maximum observed rate constant is the rate constant for dNMP incorporation ($k_{pol}$) and $K_{d}^{app}$ represents the apparent overall dissociation equilibrium constant for dNTP binding to the RB69 pol:dP/T complex prior to dNMP incorporation:

$$k_{obs} = \frac{k_{pol}[dNTP]}{K_{d}^{app} + [dNTP]}$$

**Kinetic simulations**

Kinetic simulations were performed with Tenua (provided by Dr D. Wachsstock; available free online at: http://billilite.com/tenua/), which is based on the kinetic simulation program KINSIM (44).

**RESULTS**

Our goal in this study was to use the fluorescent nucleotide analog (2AP) at either the templating base (n position) or just 5’ to the templating base (n + 1 position) to detect transition intermediates during the RB69 polymerase

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**Figure 1.** Primer/template sequences used in the fluorescence studies. The template is underlined. **A** stands for 2-aminopurine (2AP). (A) When 2AP is the templating base it is in the n position. (B) When 2AP is located 5’ to the templating base, it is in the n + 1 position. The dD/P/Ts are identical to the dP/Ts except that for dD/P/Ts the last nucleotide at the 3’ end of the primer is 2’, 3’-dideoxy-C. 2AP is located 5’ to the templating base, it is in the n + 1 position. The dD/P/Ts are identical to the dP/Ts except that for dD/P/Ts the last nucleotide at the 3’ end of the primer is 2’, 3’-dideoxy-C.
primer extension reaction, and to determine the rate constants for their formation and loss both prior to and after phosphoryl transfer. We used a ddP/T to monitor fluorescence changes associated with ternary complex formation in the absence of base incorporation. A dP/T was used to monitor changes in 2AP fluorescence when nucleotidyl transfer reaction takes place.

Equilibrium fluorescence measurements

RB69 pol (1 μM) binding to the ddP/T (200 nM) with 2AP in the n position (Figure 1) increased the 2AP fluorescence by ~6-fold (Figure 2A). Doubling the [RB69 pol] to 2 μM yielded a comparable ~6-fold enhancement, consistent with saturation being achieved at 1 μM RB69 pol and an RB69 pol:ddP/T affinity of ~100 nM, as reported for T4 pol with the same sized P/T (13/20-mer) (43).

dTTP binding to a RB69 pol:ddP/T complex (2AP in the n position) quenches the 2AP fluorescence (Figure 2A). We interpret the fluorescence quenching with dTTP as a consequence of a conformational rearrangement during the pol reaction cycle that affects 2AP stacking (30,32–35). The best fit of the [dTTP]-dependence of the equilibrium fluorescence intensity at 365 nm to a single-site binding isotherm [Equation (1)] yields an overall dissociation equilibrium constant ($K_{d}^{app}$) of 9 (±1) μM for dTTP binding (Figure 2B), a value about seven times tighter than the observed $K_{d}^{app}$ for dTTP utilization estimated by chemical quench (63 μM, Table 1). The $K_{d}^{app}$ from chemical quench measurements is the product of all dissociation equilibrium constants preceding chemistry. The equilibrium titrations of dTTP binding to RB69 pol:ddP/T (Figure 2B) should report the same overall affinity for dTTP binding prior to base incorporation. The apparent discrepancy will be addressed in the Discussion section.

With a ddP/T (2AP in the n + 1 position, Figure 1B), addition of the complementary dCTP, even at concentrations 4-fold higher than the value of $K_{d}^{app}$ determined by chemical quench, did not change the 2AP fluorescence (data not shown). Thus, dCTP binding to the RB69 pol:ddP/T complex does not alter the base-stacking interactions of 2AP when it is in the n + 1 position with the sequence we have used (33).

Kinetic analysis using a ddP/T with 2AP in the n position

Time courses after mixing RB69 pol:ddP/T complex (2AP at the n position) with a range of [dTTP] under pseudo-first order conditions ([dTTP] → [RB69 pol:ddP/T complex]) follow single exponentials (Figure 3A) with observed rate constants (Figure 3B) and amplitudes (Figure 3C) that depend hyperbolically on the [dTTP]. The nucleotide concentration dependence of the observed pseudo-first order rate constant ($k_{obs}$) was fitted to a rectangular hyperbola in the form of Equation (4):

$$k_{obs} = \frac{k_{+2}[dNTP]}{1/K_1 + [dNTP]} + k_{-2}$$

where $K_1$ is the association equilibrium constant for the formation of collision complex (EDN)** (Scheme 1) that is in rapid equilibrium with free enzyme:P/T complex (ED*) and nucleotide (N). The forward and reverse isomerization rate constants for formation of (EDN)* are $k_{+2}$ and $k_{-2}$, respectively, as defined by the following two-step reaction scheme:

$$ED^* + N \xrightarrow{K_1} (EDN)^* \xrightarrow{k_{+2}/k_{-2}} EDN^*$$

Scheme 1

in which the ** superscript denotes a higher fluorescence than *. The best fit of the [dTTP] dependence of the observed rate constants to Equation (4) yields a $1/K_1$ of ~34 μM, $k_{+2} \sim 849$ s$^{-1}$ (Table 1), and $k_{-2} \sim 44$ s$^{-1}$ (Figure 3B). The overall affinity calculated from the rate and equilibrium constants ($K_{d}^{overall} = 1/K_1K_2 = k_{-2}/K_2k_{+2}$) is ~2 μM, comparable, given experimental uncertainty, to the value obtained by equilibrium titration (9 ±1 μM; Figure 2B) and from the best fit of the [dTTP]-dependence of the fluorescence quenching amplitudes of the kinetic time courses to
Equation (1) (14 ± 7 μM; Figure 3C; Table 1). The rate and equilibrium constants of dTTP binding to RB69 pol:ddP/T are weakly dependent on temperature (Figure S1, Supplementary Data).

### Kinetic analysis using dP/T with 2AP in the n position

Time courses of fluorescence change after mixing dTTP with the RB69 pol:dP/T complex (2AP in the n position) reveal a very fast quenching phase that was completed in the stopped-flow instrument dead time (~2 ms), as indicated by the decreasing start points of the transients (Figure 4A). The magnitude of the start point changes depends on the [dTTP] with an apparent affinity of 83 μM (Figure 4C, Table 1), comparable to the nucleotide binding affinity (K_{app} = 78 μM, Table 1) determined for the best fit of the observed rate constant of the next phase following the start point change (Figure 4B). The further fluorescence decay after the starting point change follows a double exponential at 22°C (Figure 4B and Figure S2, Supplementary Data) with both observed rate constants that depend hyperbolically on the [dTTP]. At 22°C, the maximum rate of the fast phase that follows the change in start points is 220 s⁻¹ and the apparent overall equilibrium constant (K_{app}) is 78 μM, comparable to the dTTP incorporation rate constant (k_{pol}) and nucleotide affinity determined from chemical quench experiments (k_{pol} = 293 s⁻¹; K_{app} = 63 μM) (Figure 5), suggesting that these two transitions are monitoring the same process or that the biochemical transition that coincides with the fast phase of fluorescence change precedes and limits rapid base incorporation. The Y-axis intercept of the observed fast rate constant after the starting point change versus [dTTP] (Figure 4B) is indistinguishable from the origin, indicating that the process is essentially irreversible.

These results indicate that there are at least two biochemical transitions coupled to dTTP binding and incorporation with a dP/T substrate; one is very rapid (>
1000 s⁻¹) and completed within the 2 ms instrument dead time, and the second occurs at ~220 s⁻¹ when [dTTP] is saturating. This overall behavior, including the comparable values of the rate and equilibrium constants measured by fluorescence and chemical quench (Table 1) were similar at 4°C except that the fluorescence and the chemical quench were 4- to 6-fold slower. The magnitude of the rapid ‘missing’ phase (i.e. reduction in starting point) was smaller at 4°C than at 22°C (Figure S2, Supplementary Data) suggesting that we were following a process that occurred after formation of the collision complex, since formation of the collision complex should be only weakly affected by temperature. Finally, there is also a slow phase which saturates at 24 s⁻¹ at 22°C and must reflect a biochemical transition after base incorporation, which occurs at ~293 s⁻¹ (Figure 5, Table 1). This slow phase was reduced to 10 s⁻¹ at 4°C (Figure S2, Supplementary Data).

Chemical quench experiments using dTTP versus templating 2AP with the dP/T sequence shown in Figure 1A gave a k_{pol} of 293 s⁻¹ and a K_{app} of 63 μM (Figure 5A), which differs from previous measurements yielding values of 165 s⁻¹ for k_{pol} and 367 μM for K_{app} for dTTP binding and incorporation to 2AP at template n position by T4 pol (32). However, the degree of discrimination between dTTP versus dA and dTTP versus 2AP with the dP/T we have used in this study (Figure 1A, Table 1) was considerably lower than what has been reported in T4 pol (35), indicating the importance of sequence context on the kinetic parameters in this system.

### Kinetic analysis using a dP/T with 2AP at the n + 1 position

With a dP/T where 2AP is in the n position we were restricted in our choice of substrates to dTTP, as it is the only nucleoside triphosphate that can form a Watson–Crick (W–C) base pair with 2AP. To determine if the multiple phases in the time courses of fluorescence change could be observed with base pairs other than dTTP:2AP, we used 2AP in the n + 1 position with dG as the templating base (Figure 1B). Time courses of fluorescence change after mixing RB69 pol:dP/T (2AP at the n + 1 position) with increasing [dCTP] were best fit to single exponentials (Figure 6A) with the k_{obs} depending on 26 July 2018
hyberbolically on the [dCTP] (Figure 6B). Curve fitting yielded $K_{\text{app}}^{d}$ for dCTP of 10–12 $\mu$M (Table 1), comparable to the value derived from chemical quench experiments (16 $\mu$M; Table 1). The maximum rate constant is $\sim$200 s$^{-1}$ (Table 1) and is in the same range as the $k_{\text{pol}}$ value obtained from chemical quench experiments ($\sim$256 s$^{-1}$; Figure 5B, Table 1).

Kinetic simulations

Kinetic simulations were carried out based on Scheme 2 (see Discussion section) and our experimentally
determined rates and equilibrium constants for dP/T’s (with 2AP in the n position). The simulations are in excellent agreement with the observed time courses of 2AP fluorescence decay (Figure 7), indicating that our Scheme 2 is consistent with the kinetic mechanism of nucleotidyl transfer catalyzed by RB69 pol.

**DISCUSSION**

The minimal kinetic scheme for the nucleotidyl transfer reaction catalyzed by T4 pol, a close homolog of RB69 pol, was determined first by Capson et al. (43) using rapid chemical quench methods. Subsequent work using stopped-flow fluorescence provided evidence for a conformational change prior to chemistry induced by binding of the noncomplementary dATP aS to the binary complex (45). This work was extended further with T4 pol using 2AP at different positions in the template strand by Reha-Krantz and colleagues (32–35), who showed that the pre-chemistry conformational change was partially rate limiting. In our study, only one detectable conformational change preceding phosphoryl transfer induced by correct nucleotide binding to RB69 pol was observed (discussed subsequently), which differs from the results obtained with T4 pol.

**Figure 5.** Kinetics of dNMP incorporation determined by rapid chemical quench. (A) The dependence of $k_{obs}$ on [dTTP] using a dP/T (2AP in the n position). The solid line is the best fit to a hyperbola yielding a $k_{pol}$ of 293 ($\pm$20) s$^{-1}$ and $K_{a}^{app}$ of 63 ($\pm$11)µM. (B) The dependence of $k_{obs}$ on [dCTP] using a dP/T (2AP in the n + 1 position). The solid line is the best fit to a hyperbola yielding a $k_{pol}$ of 256 ($\pm$7) s$^{-1}$ and a $K_{a}^{app}$ of 16 ($\pm$2)µM.

**Figure 6.** Kinetics of dCTP binding to the dP/T (2AP in the n + 1 position) in complex with RB69 pol. (A) Time course of fluorescence changes after mixing (bottom to top) 0, 10, 20, 40, 80 and 160 µM dCTP with 200 nM RB69 pol:dP/T complex. The solid lines through the data best fits a single exponential. (B) The dependence of $k_{obs}$ on [dCTP]. The solid lines through the data (as shown by filled circle) best fits a hyperbola, yielding the maximum rate for fluorescence enhancement of 200 ($\pm$18) s$^{-1}$ and $K_{a}^{app}$ of 28 ($\pm$3)µM.

**Figure 7.** Comparison between stopped-flow data (dP/T with 2AP in the n position) and the result from kinetic simulation. As described in Experimental Procedures, simulations were carried out using the kinetic mechanism shown in Scheme 2. The simulated curve is represented by the solid line in red and is superimposed onto the data from stopped-flow fluorescence experiments shown in black. The three dTTP concentrations included for comparison are 40, 100 and 160µM, (top to bottom) respectively.
Evidence for the existence of a conformational change when these B family polymerases form a catalytically competent complex with DNA and a dNTP was based on a comparison of the crystal structures of the RB69 pol apo-enzyme (36) and the corresponding ternary complex (38), which show that fingers closing is the most prominent structural rearrangement involved in the formation of the ternary complex. The available crystal structures of binary and ternary RB69 pol complexes indicate that changes in base stacking occur and these could contribute to changes in 2AP fluorescence (38–40). It has been suggested that this conformational change provides a major checkpoint that is partly responsible for the base selectivity exhibited by many DNA polymerases (18).

Kinetic mechanism of nucleotide-dependent conformational changes in the RB69 pol complexes

Fluorescence quenching is observed both by equilibrium fluorescence titrations (Figure 2) and by stopped-flow fluorescence experiments (Figure 3), when a correct dTTP is added to an RB69 pol:ddP/T binary complex. We have assumed that the RB69 pol:ddP/T:dNTP collision complex formation does not perturb the 2AP environment, thus, no alteration in 2AP fluorescence would be expected at this point. The rapid change in fluorescence is interpreted to be a consequence of the subsequent isomerization of the open state to a closed state with lower fluorescence (Scheme 1).

The hyperbolic [dTTP]-dependence of the observed rate constant and amplitude change with an RB69 pol:ddP/T complex indicates that correct nucleotide binding is a two-step process in which a collision complex isomerizes to a partially quenched conformational state ‘preceding’ and ‘independent’ of base incorporation (Figure 3B and C and Scheme 1). The isomerization likely represents closing of the fingers, which favors 2AP stacking with the adjacent 3’ base in the primer/template duplex.

With a dP/T, the conformational change induced by correct dNTP binding was too fast to measure (>1000 s⁻¹), as it occurs within the 2 ms dead time of the instrument (Figure 4A). Evidence for this transition comes from the [dTTP]-dependent decrease in start points. In contrast, when [dTTP] is added to the RB69 pol:ddP/T complex the start points do not change and the time course of the conformational change can be determined. The amplitude of quench observed with the ddP/T (Figure 3C) is approximately equal to the magnitude of decrease in start points (i.e. missing amplitudes) observed with the dP/T substrate (Figure 4C), suggesting these changes are monitoring the same process, albeit with different rates (i.e. the process is somewhat slower and observable with ddP/T).

Time courses of fluorescence change with a dP/T after the initial rapid quench follow double exponentials. The rate of the first phase is comparable to base incorporation measured by chemical quench. The second phase is much slower than incorporation and must reflect a step after chemistry, such as switching of the EDₙ₊₁ from the pol to the exo mode, which may alter 2AP stacking interactions.

This second phase represents a process distinct from the pathway involving repeated steps in primer extension since incorporation of the next base is faster (200s⁻¹, data not shown) than the 24 s⁻¹ slow phase (Table 1 and Scheme 2).

The minimum kinetic scheme for the ddP/T and the dP/T before base incorporation are identical and consistent with Scheme 2, in which three fluorescence levels exist (as indicated by the number of superscript stars).

\[
\begin{align*}
    EDₙ^{**} + N & \xrightarrow{k_{-2}} (EDₙN)^{**} \xrightarrow{k_{-2}} (EDₙN)^* \\
    \left(\frac{k_{+3}}{k_{-3}}\right) [EDₙ₊₁•PPI] & \xrightarrow{k_{+4}} (EDₙ₊₁) + PPI \\
    k_{-8} \uparrow & k_{+8} \\
    [EDₙ₊₁•PPI]
\end{align*}
\]

Scheme 2

According to Scheme 2, the \(K_d^{overall}\) for nucleotide binding (i.e. all states preceding chemistry) and the apparent \(K_d^{app}\) values measured by stopped-flow fluorescence and by chemical quench, represent the product of all equilibria preceding chemistry (\(1/K_1K_2\)), and is supported by the agreement among: (i) the \((K_d^{overall})\) of the decrease in start points which represent the missing phase (83 μM, Figure 4C); (ii) the \((K_d^{app})\) from the chemical quench (63 μM, Figure 5A) and (iii) the \((K_d^{app})\) value from the plot of \(k_{obs}\) versus [dTTP] (78 μM, Figure 4B and Table 1) when dTMP incorporation by RB69 pol is assayed by stopped-flow fluorescence using a dP/T (2AP in the n position). Scheme 2 is consistent with either the chemical step or another conformation transition, not detectable by changes in 2AP fluorescence that precedes and limits the rate of base incorporation (data not shown in Scheme 2), as being the rate-determining step in the nucleotidyl transfer reaction catalyzed by RB69 pol. The inclusion of \([EDₙ₊₁•PPI]\) in Scheme 2 is based on the assumption that the fluorescent state of the complex just prior to chemistry is identical to that of the complex immediately after the formation of the phosphodiester bond.

It should be noted that the maximum forward rate constant \((k_{+2})\) for isomerization of the collision complex, determined with a ddP/T pseudo-substrate is slower than the rate of the very rapid fluorescence quench seen with the dP/T (>1000 s⁻¹), indicating that the ddP/T does not precisely mimic the dP/T substrate. This could be due to the absence of a 3’ terminal hydroxyl group on the ddP/T which serves as one of the ligands for the metal ion in the A site (46).

The slow rate of fluorescence quenching \(k_{+5}\) (24 s⁻¹) subsequent to chemistry leads to a state with slightly lower fluorescence that we have not defined, but have represented as \([EDₙ₊₁•PPI]\) in Scheme 2. This step competes with the pathway that leads to the \((EDₙ₊₁)\) complex that can be extended by the next correct incoming dNTP with a rate far exceeding 24 s⁻¹.
When 2AP is in the \( n + 1 \) position, fluorescence enhancement occurs after base incorporation

In contrast to the 6-fold fluorescence enhancement observed when RB69 pol forms a binary complex with a P/T (2AP in the \( n \) position), there is only a small enhancement observed for the binary complex when 2AP is in the \( n + 1 \) position, indicating that binding of the enzyme has a minor effect on 2AP stacking in this case. Addition of saturating [dCTP] to a RB69 pol:ddP/T binary complex (2AP in the \( n + 1 \) position), where G is the templating base, did not produce any change in 2AP fluorescence (data not shown). Since binding of a correct dNTP to the templating base induces a conformational change in the polymerase ternary complex, it was surprising that dCTP binding did not affect 2AP stacking (when it is in the \( n + 1 \) position) sufficiently to give a change in the fluorescence signal. The same observation with T4 pol was reported by another group (33). Only with the corresponding P/T, and after incorporation of the correct dNMP, did fluorescence enhancement occur. The maximum rate \( (k_{\text{max}}) \) for fluorescence enhancement was comparable to \( k_{\text{pol}} \) for dCMP incorporation (Table 1), suggesting that the enhancement occurs concurrently, or subsequent to, nucleotidyl transfer when the 2AP translocates from the \( n + 1 \) to the \( n \) position where it would be unstacked. We conclude that when 2AP is in the \( n + 1 \) position of a P/T with the sequence flanking the 2AP, shown in Figure 1B, it is not sensitive to conformational changes associated with the closed ternary complex formation and cannot distinguish conformational changes from base incorporation. We found that the behavior of the RB69 pol:dP/T complex (2AP in the \( n + 1 \) position) differs from that of T4 DNA polymerase (35), a close relative of RB69 pol. Hariharan et al. (35) observed a rapid 2AP fluorescence enhancement rate (314 s\(^{-1}\)) that preceded a slow rate (164 s\(^{-1}\)) of dAMP incorporation opposite a templating T. In contrast, we found that, with RB69 pol, the rates of enhancement and dCMP incorporation were nearly equivalent. In addition, the mechanism proposed by these authors, in which the conformational change associated with base unstacking (2AP in the \( n + 1 \) position) occurs before base incorporation, predicts a fluorescence enhancement with both T4 and RB69 pols when a non-extendable primer/template with 2AP in the \( n + 1 \) position is employed. This was not observed with a non-extendable primer either with T4 pol (33) or RB69 pol.

Comparison of the stopped-flow fluorescence behavior of RB69 pol complexes with other DNA pol complexes

A very rapid initial phase of fluorescence quenching with the dP/T substrate (2AP in the \( n \) position) that is nearly complete within the instrument dead time has also been observed with the Klenow fragment (KF) (30). However significant differences in the time courses for fluorescence quenching exist between KF and RB69 pol. With KF, the change in start points observed with a ddP/T, upon addition of increasing [dTTP] [Figure 3B in (30)], was not seen with the RB69 pol:ddP/T complex (Figure 3A). Another difference between KF and RB69 pol is the behavior of the enzyme:ddP/T complex (2AP in the \( n + 1 \) position) where, upon addition of the correct dNTP, fluorescence enhancement was observed with KF [Figure 4B in (30)] but not with RB69 pol. Also the behavior of the KF:dP/T complex (2AP in the \( n + 1 \) position) [Figure 4A in (30)] differed from the behavior of the corresponding RB69 pol:dP/T complex (Figure 6A). In the case of KF there was fluorescence enhancement followed by quenching upon incorporation of the correct dNMP. For RB69 pol, only fluorescence enhancement was observed (Figure 6A). The explanation for these distinctions rests on the structural differences among the various biochemical intermediates with respect to the stacking environment of 2AP during the nucleotide addition cycle.

In experiments with pol β, using a ddP/T with 2AP in either the \( n \) or \( n + 1 \) positions, a fast phase for fluorescence change was observed (26). It was suggested that this rapid phase corresponds to finger closing and that chemistry, rather than a conformational change, is rate limiting for the pol reaction (19). Recent FRET experiments with Klentaq1 also support the idea that fingers closing are faster than the kinetically determined rate-limiting step. The authors propose that the rate-limiting step, which presumably involves rearrangement of residues in the nucleotide binding pocket, occurs after the fingers close (24).

Changes in 2AP fluorescence correlate with crystal structures of RB69 pol complexes

In an attempt to understand the structural basis for the change in 2AP fluorescence, we have relied first on the commonly accepted idea that stacking of 2AP with adjacent bases or aromatic residues in the protein results in quenching of 2AP fluorescence and that enhancement...
appear to be partially stacked against the adjacent base 5' + 1 position is 11.5 Å away from the templating base in the P/T duplex even though the nucleotidyl transfer reaction has not yet occurred (Figure 8). The base in the n + 1 position is 11.5 Å away from the templating base and is displaced by ~180˚ from the P/T duplex (38). It appears to be partially stacked against the adjacent base 5' to it in the single-stranded template overhang (Figure 8). The position of the templating base before binding of the incoming dNTP, in terms of stacking, can only be inferred from the fluorescence behavior of 2AP and by analogy with other DNA pol:P/T binary complexes (47), as there is no crystal structure of binary RB69 pol:P/T complex in the pol mode, except one that has a furan moiety in the n position instead of a templating base (40). When a P/T (2AP in the n position) forms a binary complex with RB69 pol, 2AP fluorescence is also enhanced as observed with T4 pol (33). The observed bending of the template strand at the P/T junction, as seen in the crystal structure of the binary RB69 pol complex with an abasic site in place of the templating base, suggests that 2AP in the n position is less stacked in the binary complex than in the P/T alone. Our studies suggest that binding of the correct dNTP causes only the templating base (n position) to stack against its 3' neighbor, however, unstacking of the base in the n + 1 position from its 5' neighbor occurs either concurrently or subsequent to nucleotidyl transfer.

Several of our experimental observations suggest that the change in 2AP fluorescence is monitoring a conformational change, most likely fingers domain closing, because the amplitudes of the decrease in start point with increasing [dTTTP] using a dp/T is comparable to the amplitudes of quenching observed with a ddP/T (Figures 3A and 4A). In addition, the amplitude changes seen from fluorescence equilibrium titrations indicate that a structural rearrangement affecting 2AP stacking occurs after addition of a correct dNTP (dTTTP) to the highly fluorescent binary complex. This results from isomerization of an open collision complex to a closed ternary complex with lower fluorescence in the absence of chemistry.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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