Conformational Dynamics of a Y-Family DNA Polymerase during Substrate Binding and Catalysis As Revealed by Interdomain Förster Resonance Energy Transfer

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Supporting Information

ABSTRACT: Numerous kinetic, structural, and theoretical studies have established that DNA polymerases adjust their domain structures to enclose nucleotides in their active sites and then rearrange critical active site residues and substrates for catalysis, with the latter conformational change acting to kinetically limit the correct nucleotide incorporation rate. Additionally, structural studies have revealed a large conformational change between the apoprotein and the DNA–protein binary state for Y-family DNA polymerases. In previous studies [Xu, C., Maxwell, B. A., Brown, J. A., Zhang, L., and Suo, Z. (2009) PLoS Biol. 7, e1000225], a real-time Förster resonance energy transfer (FRET) method was developed to monitor the global conformational transitions of DNA polymerase IV from Sulfolobus solfataricus (Dpo4), a prototype Y-family enzyme, during nucleotide binding and incorporation by measuring changes in distance between locations on the enzyme and the DNA substrate. To elucidate further details of the conformational transitions of Dpo4 during substrate binding and catalysis, in this study, the real-time FRET technique was used to monitor changes in distance between various pairs of locations in the protein itself. In addition to providing new insight into the conformational changes as revealed in previous studies, the results here show that the previously described conformational change between the apo and DNA-bound states of Dpo4 occurs in a mechanistic step distinct from initial formation or dissociation of the binary complex of Dpo4 and DNA.

DNA polymerases are critical enzymes responsible for faithful replication of the genome. However, various DNA-damaging agents can lead to the formation of DNA lesions that often stall the progress of high-fidelity replicative DNA polymerases. The recently discovered Y-family DNA polymerases are able to bypass various types of DNA damage and rescue the stalled replication machinery in a process known as lesion bypass.1 However, these Y-family lesion bypass polymerases are characterized by low fidelity, poor processivity, and a lack of intrinsic proofreading activity when replicating undamaged DNA.2–7 Therefore, the switching between replicative and Y-family DNA polymerases during lesion bypass must be well regulated to prevent the introduction of unnecessary mutations by Y-family DNA polymerases. DNA polymerase IV from Sulfolobus solfataricus (Dpo4), the enzyme used in this study, is a model Y-family polymerase and has been extensively characterized through both structural and kinetic studies.7,14–17

In solution, DNA polymerases, like many enzymes, are inherently flexible molecules that can undergo a variety of motions ranging from localized vibrations of side chains to translations and rotations of loops and secondary structures and even concerted global conformational transitions.18 These conformational rearrangements are critical for the functions of enzymes, including catalysis,19 signal transduction,20 allosteric regulation,21 and structural-stability modification.22 While substrate binding and catalysis usually occur at a specific “active site” in an enzyme involving only a few residues, structural changes are often observed at residues far from the active site and may involve rigid body motions of large protein domains or secondary structure elements consisting of many residues.

All DNA polymerases share a structurally conserved polymerase core with a “right hand” architecture, consisting of Finger, Thumb, and Palm domains, while Y-family DNA polymerases contain a unique Little Finger (LF) domain in addition to the polymerase core.9,23–25 Upon binding to an incoming nucleotide, many DNA polymerases have been shown to undergo a Finger domain closing conformational change that helps bring functionally important enzyme residues in contact with the nucleotide and serve as a fidelity check point for preventing misincorporations.26 Crystal structures of Y-family DNA polymerases such as Dpo4 bound to DNA do not reveal a large domain swing upon nucleotide binding analogous to the Finger domain closing observed for other DNA polymerases.9,11 Nevertheless, stopped-flow studies monitoring

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both Trp fluorescence\textsuperscript{27} and FRET\textsuperscript{28} as well as hydrogen–deuterium exchange experiments\textsuperscript{30,31} have provided information about global conformational changes that occur throughout each of the four domains of Dpo4 during correct nucleotide binding and catalysis. Furthermore, it was shown that these conformational changes are altered by the presence of a lesion in the DNA template.\textsuperscript{30,31} Additionally, X-ray crystallography and tryptophan fluorescence studies have shown that Dpo4 undergoes a large conformational change from the apo form to the DNA-bound binary state characterized by a rotation of the LF domain relative to the polymerase core.\textsuperscript{8} In a recent study, this conformational change was proposed to be involved in polymerase switching during lesion bypass.\textsuperscript{32}

Our previous studies made use of two FRET systems to monitor changes in distance either between the DNA substrate and each domain of Dpo4 or between the Finger and LF domain.\textsuperscript{28} Here, we have further investigated the conformational changes during nucleotide binding and incorporation by utilizing a FRET system with a donor in the LF domain and an acceptor in the Palm domain as well as another FRET system with a donor in the Palm domain and an acceptor in each of the remaining domains of Dpo4. Furthermore, both intrinsic tryptophan fluorescence and FRET spectroscopy were used to investigate the conformational changes of the LF domain relative to the polymerase core of Dpo4 during DNA binding and dissociation.

\section*{Materials and Methods}

\textbf{Materials.} The cysteine reactive dye, 7-(diethylamino)-3-\((4'\text{-maleimidophenyl})\)-4-methylcoumarin (CPM), was obtained from Invitrogen. The DNA substrate consisted of two synthetic oligodeoxynucleotides synthesized by Integrated DNA Technologies (IDT), a 21-mer primer (5′-CG AGC CGT CGC ATC CTA CCG C-3′) and a 30-mer template (5′-G ATG CTG CAG CGG TAG GAT GCG ACG GCT CG-3′). The primer 21-mer in the DNA substrate 21/30-mer is either normal (DNA\textsuperscript{10}) or 3′-dideoxy-terminated (DNA\textsuperscript{10}) as synthesized by Integrated DNA Technologies. The DNA substrates were purified by denaturing polyacrylamide gel electrophoresis (17% polyacrylamide and 8 M urea), and the concentrations were determined by UV absorbance at 260 nm using the corresponding extinction coefficients.

\textbf{Mutagenesis, Purification, and Labeling of Dpo4.} The expression plasmid containing the \textit{S. solfataricus} dpo4 gene was constructed as described previously.\textsuperscript{14} Mutations were introduced by using the Quikchange kit (Stratagene) according to the manufacturer’s instructions. First, to avoid the ambiguity of labeling, the sole native cysteine of Dpo4 was mutated to serine (C31S). Second, using the C31S mutant as a template, a unique intrinsic fluorescence donor Trp residue was introduced individually into the LF (Y274W) or Palm (Y108W) domain of Dpo4. Finally, on the basis of this double mutant, a single cysteine was individually substituted on the domains of Dpo4 to serve as the specific site of attachment for the acceptor CPM. These mutations were confirmed by DNA sequencing and are shown in Figure 1. The Dpo4 mutants were expressed and purified as described for wild-type (wt) Dpo4.\textsuperscript{14} The concentration of each mutant was determined spectrophotometrically at 280 nm using a molar extinction coefficient of 28068 M\textsuperscript{−1} cm\textsuperscript{−1}, which was calculated on the basis of the sequence of wt Dpo4 containing a single Tyr to Trp mutation.

The labeling of Dpo4 mutants was performed in the labeling buffer [50 mM Tris-HCl (pH 7.5), 0.15 M NaCl, 10% glycerol, and 0.5 mM TCEP]. A 10–20-fold molar excess of CPM was added dropwise to a 50–100 \(\mu\) M protein solution. The labeling reaction was allowed to proceed while the mixture was slowly and constantly mixed at 4 °C overnight, and unreacted dye was quenched by adding an equimolar amount of glutathione. Labeled protein was separated from the unbound fraction of dye by size exclusion gel filtration (Sephadex G-25 resin). Eluted protein was extensively dialyzed against the Dpo4 storage buffer and stored as aliquots at −80 °C. The protein concentration was determined by using the Bradford assay (Bio-Rad) with wt Dpo4 as a standard, and the degree of labeling was typically ≥90%.

\textbf{Buffers.} All experiments, if not specified, were conducted in buffer R, consisting of 50 mM HEPES (pH 7.5) at 20 °C, 6 mM MgCl\textsubscript{2}, 50 mM NaCl, 0.1 mM EDTA, 1 mM DTT, and 10% glycerol.

\textbf{Stopped-Flow Assays.} Circular dichroism (CD) spectra were recorded using a model 62A DS spectrometer (Aviv) in a 1 mm path length cuvette at 37 °C. The spectra were recorded in the buffer [25 mM sodium phosphate (pH 7.5), 50 mM NaCl, and 5 mM MgCl\textsubscript{2}]. Data points from 270 to 200 nm were recorded for Dpo4 and its mutants at 1 nm intervals. Each data point was averaged for 5 s. All measurements were corrected for the background signal.

\textbf{Stopped-Flow Assays.} Stopped-flow fluorescence experiments were performed on an Applied Photophysics SX20 system at a constant temperature of 20 °C. Samples were excited at 290 nm, and the fluorescence of Trp or CPM was monitored by using a 305 or 420 nm cutoff filter, respectively. Again, the slits were set at 5 nm for both excitation and emission. All reported concentrations represent the final concentrations of components after mixing. The rate for each
fluorescent phase was determined by fitting a stopped-flow fluorescence trace using the KinTek Explorer global fitting software. The software was set to fit each trace using a single- or double-exponential equation depending on the number of observed FRET phases. Reported rates and the associated errors are reported as averages and standard deviations determined from fits to traces acquired in multiple repeat stopped-flow mixing experiments.

**RESULTS**

**Construction of an Interdomain FRET System.** To monitor changes in distance between two domains, Dpo4 mutants containing a single-trypthophan FRET donor fluorophore and a single Cys for labeling with the acceptor fluorophore CPM were generated (Figure 1 and Table 1).

**Table 1. Distances between the Trp Donor and Residues with an Attached CPM Acceptor Estimated from Crystal Structures**

| Protein Form | Distance (Å) | Palm | Thumb |
|--------------|--------------|------|-------|
| apo b        | 28.4         | 30.4 | 20.2  |
| binary c     | 38.1         | 31.0 | 44.7  |
| ternary d    | 36.6         | 31.3 | 45.8  |

| Protein Form | Distance (Å) | LF | Finger | Thumb |
|--------------|--------------|----|-------|-------|
| apo b        | 44.7         | 28.8| 38.4  | 32.7  |
| binary c     | 42.6         | 35.2| 38.3  | 31.6  |
| ternary d    | 42.6         | 35.4| 38.2  | 31.7  |

**Table 2. Observed Rates (kobs) of Incorporation of dTTP into DNA OH Catalyzed by Dpo4 or Each of Its CPM-Labeled Mutants at 20°C under Single-Turnover Reaction Conditions**

| Enzyme       | kobs (±) (s⁻¹) |
|--------------|----------------|
| wt Dpo4      | 0.8 ± 1        |
| Y274W-S96CCPM| 0.8 ± 1        |
| Y274W-S112CCPM| 0.54 ± 0.04   |
| Y108W-N70CCPM| 0.33 ± 0.03    |
| Y108W-K26CCPM| 0.39 ± 0.03    |
| Y108W-K329CCPM| 0.47 ± 0.04   |
| Y108W-S307CCPM| 0.35 ± 0.03   |
| Y108W-S207CCPM| 0.39 ± 0.03   |
| Y108W-N200CCPM| 0.36 ± 0.02   |

*Obtained from single-turnover kinetic assays in which a preincubated solution of a Dpo4 mutant (180 nM) and 5’-32P-labeled DNAOH (30 nM) was rapidly mixed with dTTP (1 mM) and subsequently quenched at various time intervals with 0.37 M EDTA and the plot of product concentration vs time was fit to the single-exponential equation [product] = A[1 - exp(-ktobs)] to yield an observed rate kobs and a reaction amplitude (A).

Because wt Dpo4 contains no native Trp residues, single Trp residues were introduced separately into the Palm (Y108W) or LF (Y274W) domain via site-directed mutagenesis. To avoid ambiguity in CPM labeling, the single native Cys (C31) of Dpo4 was mutated to Ser. Using the expression DNA plasmid encoding a Dpo4 mutant containing a single Trp and no Cys, single Cys residues were introduced into various positions throughout Dpo4 for labeling with CPM (Figure 1 and Materials and Methods). For the sake of simplicity, triple Dpo4 mutants in this study will be cited by the single Trp mutation and the introduced Cys only, e.g., Y274W-S96CCPM. Enzymatic activities of CPM-labeled, e.g., Y274W-S96CCPM, and unlabeled Dpo4 mutants, e.g., Y274W-S96CC, were measured under single-turnover conditions using 32P-based kinetic assays (Materials and Methods). The observed rates of nucleotide incorporation (kobs) for the unlabeled and CPM-labeled Dpo4 mutants were significant to that measured for the wt enzyme (Table 2), indicating that neither the mutations nor the labeling had a

**Steady-State Fluorescence of Dpo4 Mutants Y108W and Y274W.** To gain insight into the local environment of the engineered Trp residues, fluorescence emission spectra for unlabeled Dpo4 mutants Y108W and Y274W were recorded (Figure 2A). The emission maximum for Y108W was approximately 320 nm, characteristic of a buried Trp residue, while the red-shifted emission maximum (354 nm) of Y274W is consistent with a higher degree of solvent exposure. When a normal DNA substrate 21/30-mer was bound [DNA OH], a significant reduction in Trp fluorescence was observed for both Y108W (30.3%) and Y274W (21.6%). Following DNA binding, dTTP incorporation into the E-DNA complex resulted in a further decrease in Trp fluorescence. As both DNA and dTTP absorb at 290 nm, the observed decreases in Trp fluorescence upon their additions may be primarily due to inner filter effects resulting from their absorbance of the excitation light. However, the difference in fluorescence attenuation for the two mutants upon the addition of DNA (30.3 and 21.6% for Y108W and Y274W, respectively) suggests that protein conformational changes or interaction with the DNA substrate could also contribute to the changes in Trp fluorescence, as inner filter effects would be expected to decrease the Trp fluorescence of the two mutants to the same extent.

**Steady-state fluorescence spectra of mutants Y274W-S96CCPM and Y108W-K329CCPM (panels B and C of Figure 2, respectively, black traces) show that the attachment of CPM to a unique cysteine residue had a significant quenching effect on the Trp fluorescence of both Y274W and Y108W, upon excitation at 290 nm, the peak excitation wavelength of Trp. Concomitantly, an intense acceptor peak centered at 472 nm was observed, which was a strong indicator of the efficient
transfer of energy from Trp to CPM. The addition of the substrate DNA-OH resulted in a very slight increase in the acceptor CPM fluorescence for mutant Y274W-K96CCPM and a dramatic increase for mutant Y108W-K329CCPM (~30%) (Figure 2B,C, green traces). These results suggest that the Y108W-K329CCPM FRET pair is better suited for dynamically investigating the conformational change in the LF domain relative to the polymerase core. Addition of dTTP to the binary complexes of the Dpo4 mutants with DNA-OH resulted in a dramatic decrease in CPM fluorescence for both Y274W-S96CCPM (38%) and Y108W-K329CCPM (28%), which is predominantly due to inner filter effects from the absorbance of the excitation light by dTTP with a high concentration (1 mM), rather than resulting only from a change in FRET efficiency as both CPM and Trp fluorescence intensities were reduced (Figure 2B,C). However, as described above, the differences observed for the two mutants in panels B and C of Figure 2 suggest that conformational changes may also contribute to the changes in fluorescence. All other Dpo4 mutants listed in Table 2 exhibited similar trends consistent with the distance variation based upon the crystal structure analysis of apo and binary complexes (data not shown).

**Conformational Change upon Binding of Dpo4 to DNA.** To further investigate the observed conformational
change in the LF domain relative to the polymerase core during DNA binding, rapid mixing experiments were performed using the Y108W or Y108W-K329C\textsubscript{CPM} mutant in a stopped-flow apparatus upon excitation of Trp at 290 nm. Notably, all stopped-flow experiments in this study were performed at 20 °C to be consistent with published stopped-flow FRET studies with Dpo4 that showed that some FRET changes are not easily observed at higher temperatures.\textsuperscript{8,30} Mixing of Dpo4 mutant Y108W with DNA in the absence of the CPM acceptor did not result in a detectable time-dependent change in fluorescence (Figure S2A of the Supporting Information), indicating that the quenching of Trp fluorescence observed in the steady-state assay (Figure 2A) occurred within the dead time (\sim 1 ms) of the stopped-flow apparatus, as would be expected for quenching due to inner filter effects. However, mixing of Y108W-K329C\textsubscript{CPM} with DNA resulted in a measurable rapid increase in CPM fluorescence (Figure 3A), consistent with steady-state FRET measurements (Figure 2C). Interestingly, the rate of this fluorescence change (\sim 16 s\textsuperscript{-1}) did not increase as the concentration of DNA was increased from 2 to 100 nM while the enzyme concentration was held constant at 100 nM (Figure 3B). Notably, at concentrations of <2 nM, the amplitude of the fluorescence change was too low to be easily detected in the stopped-flow apparatus. The observation of a constant rate over a nearly 2 order of magnitude increase in DNA concentration suggests that the change in fluorescence may correspond to a conformational change that occurs at a step distinct from the formation of the Dpo4-DNA binary complex as the later step is expected to be a second-order process with an observed rate that increases linearly with DNA concentration. Several assays were also performed to investigate the reverse conformational change upon dissociation of the Dpo4-DNA complex. The mixing of a preincubated solution of the Y108W mutant and DNA with a 20-fold excess of tryptophan-free wt Dpo4 to sequester any DNA that dissociates from the labeled enzyme resulted in a time-dependent increase in Trp fluorescence (Figure 3C). This result is consistent with the steady-state spectra that show that the fluorescence of Y108W was decreased by 30.3% upon addition of DNA (Figure 2A). Moreover, the decrease is \sim 10% more with the Y108W mutant than with the Y274W mutant, suggesting that a conformational change upon DNA binding is partially responsible for the observed decrease in Trp fluorescence (Figure 2A). When the experiment was performed by mixing the preincubated, labeled Y108W-K329C\textsubscript{CPM}-DNA complex with excess unlabeled wt Dpo4, a large decrease in CPM fluorescence was observed (Figure 3D). Notably, the rate of the CPM fluorescence change for the labeled Y108W-K329C\textsubscript{CPM} (3.3 \pm 0.2 s\textsuperscript{-1}) was \sim 2-fold slower than the observed rate for the Y108W mutant (6 \pm 1 s\textsuperscript{-1}) (Figure 3C,D). The latter rate may be less accurate because of the low signal-to-noise ratio (Figure 3C). Using an alternative approach, a preincubated solution of unlabeled Dpo4 and DNA was mixed with an excess of the labeled Y108W-K329C\textsubscript{CPM} mutant, which resulted in a biphasic increase in CPM fluorescence (Figure 3E). The faster phase rate (15.1 \pm 0.6 s\textsuperscript{-1}) was similar to the rate measured for the forward conformational change rate (Figure 3B) and likely results from an initial binding of the small fraction of DNA that was not bound to the unlabeled Dpo4 during the preincubation prior to mixing. The slower phase likely results from the binding of DNA to the labeled Dpo4 after it has dissociated from unlabeled Dpo4 with the observed rate (1.7 \pm 0.2 s\textsuperscript{-1}), which is a function of the rates of both the forward (\sim 16 s\textsuperscript{-1}) and reverse (3.3 s\textsuperscript{-1}) conformational changes. Together, these results suggest that Dpo4 utilizes a DNA binding mechanism with a conformational change occurring at a step different from the formation and dissociation of the Dpo4-DNA binary complex.

**Conformational Dynamics during Correct Nucleotide Incorporation.** To characterize the dynamics of conformational transitions employed by Dpo4 during correct nucleotide insertion, a solution containing a CPM-labeled Dpo4 mutant preincubated with DNA\textsuperscript{OM} was rapidly mixed with the correct dTTP, and the CPM fluorescence was monitored upon excitation at 290 nm. Only the acceptor CPM fluorescence was monitored, as the Trp fluorescence was too weak to be accurately measured in the presence of the CPM acceptor. Notably, no time-dependent change in fluorescence was observed upon mixing of dTTP with labeled Dpo4 in the absence of the DNA (Figure S2B of the Supporting Information), suggesting that the fluorescence changes resulting from inner filter effects observed in the steady-state experiments (Figure 2) occurred within the mixing dead time (\sim 1 ms) of the stopped-flow instrument. Additionally, no change in Trp fluorescence was observed in control experiments in the absence of the CPM acceptor (Figure S2C of the Supporting Information), indicating that any observed changes in CPM fluorescence would be primarily due to changes in FRET efficiency rather than changes in the local environment of the Trp probe.

In a previously published study, the conformational changes of the Finger domain (N70C and K26C) relative to the LF domain (Y274W) were investigated through two labeled mutants, Y274W-N70C\textsubscript{CPM} and Y274W-K26C\textsubscript{CPM}.\textsuperscript{28} Here, similar assays were performed to investigate the conformational changes of the Palm domain at the residues of S96C and S112C relative to the LF domain. The distances between potential labeling residues in the Thumb domain, e.g., S207C and N200C, and the Y274W residue in the LF domain in the binary and ternary structures are greater than R\textsubscript{0} \pm 0.5R\textsubscript{0}, where R\textsubscript{0} is the Förster distance of 30 Å for a Trp–CPM FRET pair (Table 1). Therefore, changes in distances between these two domains during nucleotide binding and incorporation could not be accurately measured using this FRET system. When a preincubated solution of the labeled mutant Y274W-S112C\textsubscript{CPM} and DNA was mixed with the correct dTTP, a biphasic fluorescence change was observed, consisting of a rapid [17 s\textsuperscript{-1} (Table 3)] decrease phase (P\textsubscript{1}) followed by a slow [0.3 s\textsuperscript{-1} (Table 3)] fluorescence increase phase (P\textsubscript{2}) (Figure 4A, black trace). Notably, similar results were observed previously for the motion of the Finger domain relative to the LF domain.\textsuperscript{28} The signal-to-noise ratio for the Y274W-S96C\textsubscript{CPM} mutant was significantly lower than for the Y274W-S112C\textsubscript{CPM} mutant, and consequently, a second P\textsubscript{2} phase could not be clearly distinguished (Figure 4B), indicating that this FRET pair was not as well suited for measuring conformational changes. These results, along with those from previous studies,\textsuperscript{28} indicate that the LF domain moved away from both the Finger and Palm domains and then closed back in toward the polymerase core to complete one catalytic cycle.

Next, the conformational transitions of the LF, Finger, and Thumb domains relative to the Trp donor (Y108W) in the Palm domain were investigated. The fluorescence changes observed for this system were much more varied than those obtained with the Trp donor (Y274W) located in the LF domain. Finger domain residue N70C (Figure 5A and Table 3)
showed a biphasic trace similar to that observed for the same residue relative to the LF domain in published studies. Notably, the amplitude of the fluorescence change for the Finger mutant Y108W-K26C was significantly lower than that observed for the other site (N70C) on the Finger domain, and the P2 phase was less obvious (Figure 5B). In contrast, the LF residue S307C displayed a rapid decrease P1 phase while a P2 phase was not observed upon mixing with dTTP while the rapid P1 phase remained unchanged at an average rate of approximately 20 s⁻¹ (Table 3 and Figures 4 and 5). This suggests that, for these mutants, P1 represents a precatalytic conformational change while P2 corresponds to a transition taking place after nucleotide incorporation. Additionally, the rate of P2 is similar to the rate for the rate-limiting step in nucleotide incorporation under single-turnover conditions for each mutant [average kₐ of 0.5 s⁻¹ (Table 2)]. The one exception was mutant Y108W-S307C CPM (Figure 5F), for which the P2 phase was essentially identical for both DNAOH and DNAH substrates. These results indicate that Dpo4 may undergo an additional transition after the initial fast conformational change that is independent of phosphodiester bond formation. However, this conclusion needs to be verified by further studies.

Table 3. Rates of Stopped-Flow Fluorescence Changes at 20 °C

| Domain | CPM-labeled residue | phase | rate (s⁻¹) with DNAOH | rate (s⁻¹) with DNAH |
|--------|---------------------|-------|-----------------------|----------------------|
| Relative to the Trp Donor (Y274W) in the LF Domain |
| Palm   | S112C               | P1    | 17 ± 4                | 19 ± 1               |
|        |                     | P2    | 0.7 ± 0.1             | –                    |
|        | S96C                | P1    | 11.4 ± 1.7            | 12.2 ± 1.5           |
|        |                     | P2    | –                     | –                    |
| Relative to the Trp Donor (Y108W) in the Palm Domain |
| Finger | N70C                | P1    | 23 ± 4                | 20 ± 4               |
|        | K26C                | P1    | 5.2 ± 0.6             | 3.7 ± 0.6            |
|        |                     | P2    | –                     | –                    |
| Finger | N70C                | P1    | 20 ± 4                | 20 ± 5               |
|        |                     | P2    | 0.27 ± 0.07           | 0.27 ± 0.06          |
| Thumb  | N200C               | P1    | 17 ± 4                | 17 ± 3               |
|        |                     | P2    | –                     | –                    |
| Thumb  | S207C               | P1    | 16 ± 7                | 17 ± 5               |
|        |                     | P2    | 0.16 ± 0.03           | –                    |

Conformational Dynamics during Correct dNTP Binding with a Dideoxy-Terminated DNA Substrate. To explore the origins of the observed biphasic kinetics, a nonextendible substrate 21/30-mer DNAH (Materials and Methods) was employed to distinguish between events occurring before and after phosphodiester bond formation. When nucleotide incorporation was prevented, the P1 phase for all but one of the mutants was not observed upon mixing with dTTP while the rapid P1 phase remained unchanged at an average rate of approximately 20 s⁻¹ (Table 3 and Figures 4 and 5). Thus, LF residues S307C and Y274W appear to move in the opposite direction compared to the other LF residue, K329C, relative to the Palm domain. Notably, the rates of each phase were similar for all mutants except K26C (Table 3).

In summary, the stopped-flow studies presented here revealed conformational changes involved in DNA binding by Dpo4 upon nucleotide binding or incorporation. This information provides insights into the directional and rotational nature of those motions. Despite the fact that X-ray crystallography has not revealed large conformational shifts in Dpo4 upon nucleotide binding or incorporation, previous FRET studies have characterized pre- and postcatalytic conformational changes in all four domains of Dpo4 by monitoring the changes in distance between the protein and the DNA substrate. The results of this investigation uncovered additional details of these conformational changes as well as conformational changes involved in DNA binding by Dpo4 during nucleotide binding and incorporation.
monitoring changes in distance between pairs of locations on the protein itself.

Published crystallographic studies revealed that Dpo4 undergoes a significant conformational change from the apo state to the DNA-bound state with a 131° rotation of the LF domain relative to the polymerase core. The results here show that this conformational change in the LF domain occurs at a rate of \(~16\,s^{-1}\) and is independent of DNA concentration over nearly 2 orders of magnitude (Figure 3A,B). In contrast, in a previous study that monitored FRET between labeled DNA and a labeled DNA polymerases to directly monitor formation of the E·DNA binary complex, the observed rate of change in fluorescence upon mixing of the polymerase with the DNA was found to increase linearly with an increasing protein concentration, as would be expected for a second-order binding process. Thus, the results of this study are consistent with a multistep binding mode in which this conformational change occurs in a step distinct from formation of the Dpo4·DNA complex. Two potential models for the binding of Dpo4 to DNA are possible: either the conformational change occurs after the initial binding of Dpo4 to DNA (Figure 6A), or Dpo4 is in equilibrium between two conformations, only one of which is capable of binding to DNA (Figure 6B). The crystal structure of apo Dpo4 shows that the LF domain actually occupies the DNA binding cleft of the polymerase core, which would support the model shown in Figure 6B, as the LF domain would need to change its conformation to allow DNA to bind to Dpo4.

Previous kinetic studies have indicated that the rate-limiting step in steady-state catalysis is the slow dissociation of the DNA from Dpo4 (0.02 \(s^{-1}\) at 37 °C), several orders of magnitude lower than the rates of the conformational change measured here at 20 °C by Trp fluorescence \([6\,s^{-1}\) (Figure 3C)] or by FRET monitoring of CPM fluorescence \([3.3\,s^{-1}\) (Figure 3D)].

Figure 5. Conformational motions of the Finger, Thumb, and LF domains relative to the Palm domain during nucleotide binding and incorporation. Changes in CPM fluorescence upon excitation of Trp at 290 nm are shown after mixing of 1 mM dTTP with a preincubated solution of a Dpo4 mutant (200 nM) and either DNAOH (300 nM, black) or DNAH (300 nM, red) at 20 °C: (A) Y108W-N70CCPM, (B) Y108W-K26CCPM, (C) Y108W-S207CCPM, (D) Y108W-N200CCPM, (E) Y108W-K329C, and (F) Y108W-S307CCPM.
In the 32P-based kinetic assay, the loss of product formation was measured upon mixing with dNTP for a fixed time interval after a preincubated solution of Dpo4 and radiolabeled DNA had been mixed with an excess of unlabeled trap DNA for varying time intervals. This difference in rates obtained from the two techniques lends further support to a model in which the conformational change and DNA binding represent distinct steps. Notably, from these results, the possibility of a more complex binding process with additional steps in which Dpo4 adopts conformations intermediate between those observed in the apo and DNA-bound crystal structures cannot be ruled out.

To account for the results from fluorescence assays in this study, along with those from previous kinetic and structural studies, an expanded three-step model for DNA binding has been added to the previously proposed catalytic mechanism of Dpo4 (Figure 6C). In this model, it is proposed that, in the absence of DNA, the LF domain of Dpo4 is in equilibrium between two conformations (step 1, Figure 6C), only one of which (E) allows for DNA binding, and that after an initial rapid binding to DNA (step 2, Figure 6C), the binary complex is also in equilibrium between a nonproductive state and a productive state (step 3, Figure 6C), only one of which (E-DNA) is active for subsequent dNTP binding and incorporation. The inclusion of step 3 in the proposed mechanism is consistent with the binary crystal structure in which the terminal base pair occupies the dNTP binding site (E-DNA*).

In previous stopped-flow FRET studies, a very rapid step occurring prior to the P1 phase was attributed to a translocation of the DNA relative to Dpo4, corresponding to a shift in equilibrium from the nonproductive state to the productive state. In a recent single-molecule FRET investigation, Dpo4 was observed to alternate between these two conformations in the binary complex, with the binding of an incoming dNTP stabilizing the post-translocated (E-DNA) productive state, lending further support to the inclusion of step 3 in the proposed model (Figure 6C). This rapid conformational rearrangement was also proposed for the Y-family DNA polymerases from Sulfolobus acidocaldarius (Dbh)-based stopped-flow fluorescence measurements with a DNA substrate containing a 2-aminopurine fluorescent base analogue.

The observed rate of 0.02 s−1 in 32P-based kinetic assays corresponds to the loss of only the catalytically competent E-DNA complex (k−1) rather than much faster steps (k−1 and k−3), as the assay relies on observation of a decreasing amount of nucleotide incorporation product following the increased incubation time with the trap DNA. In our stopped-flow assays, the observed increase in fluorescence upon mixing Dpo4 with DNA (Figure 3A) would result from a shift in the equilibrium from the Eapo state to the E state [k1 (Figure 6C)] followed by steps 2 and 3 that do not lead to a change in fluorescence. Similarly, the decrease in fluorescence upon mixing the Trp-containing or labeled Dpo4 mutant-DNA complex with unlabeled wt Dpo4 (Figure 3C,D) results from the shift in equilibrium back toward the Eapo state [k−1 (Figure 6C)] following a more rapid dissociation of the E-DNA* complex in step 2. Notably, without step 3 in the proposed model in Figure 6C (i.e., the model in Figure 6B), one would expect the rate measured in panels C and D of Figure 3 to be much slower as observed in the previously described 32P-based kinetic trap assay, as k−3 (step 3, Figure 6C) would likely be lower at 20 °C than the value of 0.02 s−1 measured at 37 °C.

Clearly, the binding of Dpo4 to DNA is a complex process, and while the FRET data described here provide new insight into an early conformational change in the binding pathway, further investigation is necessary to fully elucidate the binding mechanism. To this end, additional stopped-flow as well as single-molecule fluorescence studies to further probe the complex mechanism of binding of DNA to Dpo4 are currently underway.

The conformational change in the orientation of the LF domain upon DNA binding may be a phenomenon common to all Y-family DNA polymerases. As with Dpo4, crystal structures of human DNA polymerases κ (hPolκ) also show a change of up to 50 Å in the position of the LF domain relative to the polymerase core between the apo and DNA-bound states. Furthermore, yeast DNA polymerase η (yPolη) also undergoes a structural change upon DNA binding, although it is much more modest, consisting of an 8° rotation of the LF domain. Interestingly, the LF domain of hPolκ was
observed in two different conformations in the apo state, both of which are distinct from the conformation observed in the DNA-bound state. While only one conformation of the LF domain has been observed for Dpo4 in the apo state, the crystal structure of Dpo4 in complex with the sliding clamp replication factor shows that the LF domain adopts a conformation different from that observed in the apo or DNA-bound form. This suggests that a certain degree of conformational flexibility exists in the position of the LF domain even in the absence of a DNA substrate for these Y-family DNA polymerases, which is consistent with the inclusion of step 1 in the proposed model (Figure 6C). However, further structural and kinetic investigations are necessary to determine whether the kinetic mechanism proposed here will apply to other Y-family DNA polymerases.

In structural studies with both Dpo4 and the bacterial Y-family enzyme *Escherichia coli* DNA polymerase IV, it was suggested each of the DNA polymerases primarily binds to a sliding clamp through its interactions with the LF domain in an inactive conformation that cannot bind to DNA, and thus, a conformational change would be required to allow for DNA binding by the Y-family DNA polymerase. To this end, it is possible that the conformational change in the LF domain prior to DNA binding (step 1, Figure 6C) may serve as a point of regulation during polymerase switching to prevent the low-fidelity Y-family DNA polymerases from accessing the replication fork prior to stalling of the replicative polymerases at DNA lesion sites.

Differences in DNA dissociation rates as measured by fluorescence versus 32P-based kinetic assays have also been reported for A-family and B-family DNA polymerases as well, suggesting that these DNA polymerase families may utilize a multistep DNA binding mechanism similar to that of Dpo4. For example, crystal structures of the large fragment of *Thermus aquaticus* DNA polymerase I (Klenaq1) show that, in the E-DNA binary complex, the template base of the DNA substrate stacks with a conserved Tyr residue, and thus, a conformational rearrangement is required for dNTP binding. Subsequently, stopped-flow FRET investigations with Klenaq1 and the Klenow fragment of *E. coli* DNA polymerase I have revealed evidence of a pre-equilibrium step prior to nucleotide binding similar to step 3 in the proposed Dpo4 mechanism (Figure 6C). However, because other DNA polymerase families do not possess a domain analogous to the LF domain, it is unclear whether a transition analogous to step 1 in the proposed mechanism (Figure 6C) might be a general feature for DNA polymerases beyond those in the Y-family.

The interdomain FRET changes observed upon mixing a preformed Dpo4-DNA binary complex with dNTPs (Figures 4 and 5) are consistent with the previously proposed nucleotide incorporation mechanism involving pre- and postcatalytic conformational change steps (steps 4–10, Figure 6C). The biphasic traces observed with the Trp donor in the LF domain and the CPM acceptor in the Palm domain here (Figure 4) or in the Finger domain in a previous study suggested that the LF domain moved away from the polymerase core before catalysis [P1 (Figure 6C)] and returned toward it after catalysis [P2 (Figure 6C)]. Similar results were obtained using mutant Y108W-S307C-CpM (Figure 5F), which also allowed us to monitor the change in distance between the Palm and LF domains. These observations are consistent with previous studies that showed that the LF domain also moves in a similar fashion relative to the DNA substrate and is further supported by computational predictions.

Despite the similarity in the results obtained for the motion of LF residue Y274W relative to the Finger or Palm domain residues, the motion of the LF domain cannot be considered as a simple whole domain translational movement outward from the polymerase core and DNA substrate prior to catalysis, as revealed by mutant Y108W-K329C (Figure 5E). Mutant Y108W-K329C exhibited FRET changes in the opposite direction of those of all other FRET pairs between the LF and Palm domains. Inspection of the LF’s crystal structure revealed that residues K329C and S307C are located on the loops on either side of α-helix M and that Y274W is located on α-helix L, near S307C (Figure 1). A possible interpretation of the results of this study is that the LF domain rotates or bends about an axis perpendicular to helices M and L, where both residues S307C and Y274W are located on the one side of the axis and K329C is positioned on the other side of the axis. The possibility of a rotational axis is supported by molecular simulation, which suggests that the LF domain pivots around the DNA major groove by approximately 12°, as measured by the rotation of α helices L and M. In previous FRET studies, similar rotational components were observed for the motion of the Finger and Palm domains relative to DNA during DNA lesion bypass.

While most mutants exhibited two clear FRET phases, a clear P2 phase was not apparent for mutants Y274W-S96C-CpM (black trace, Figure 4B) and Y108W-N200C-CpM (black trace, Figure 5D). It is possible that a change in distance between the FRET pairs in these mutants occurs during P2, but that the amplitude of this change is too small to be detected given the inherent noise in the stopped-flow data. Conversely, the possibility that additional conformational change steps may be occurring during incorporation of the nucleotide by Dpo4 that complicate the data analysis for these particular FRET pairs cannot be ruled out. The observation of a second fluorescence phase for mutant Y108W-S307C-CpM even with the dideoxy-terminated primer (red trace, Figure 5F) may result from an additional conformational change step that occurs at a rate similar to the steps that give rise to P3 for the other mutants but is independent of covalent nucleotide incorporation. If such a step also affects the motions of the FRET pairs in mutants Y274W-S96C-CpM and Y108W-N200C-CpM, it may obscure the P2 phase for these mutants. However, the results of this study are not sufficient to determine whether this additional step or some other mechanistic phenomenon is responsible for the observed fluorescence changes in these three mutants.

It is clear from these FRET studies that the conformational dynamics of Dpo4 are complicated, and while the use of multiple FRET pairs allows us to provide some insight into the structural details of these conformational changes, further solution-state structure studies are necessary to gain a clear picture of how each domain moves throughout the entire reaction pathway. Recent chemical shift assignments of the backbone nitrogens, α and β carbons, and amide protons of the polymerase core and the LF domain of Dpo4 will likely facilitate future investigations of this nature at atomic resolution.

This work provides fresh insight into the importance of domain motions of Dpo4 to both DNA binding and catalysis and creates a global, complex dynamic picture of domain motions during the catalytic cycle. It would be very interesting to determine whether Dpo4 employs different conformational
motions while incorporating correct or incorrect dNTP into undamaged DNA or encountering and bypassing a variety of lesions. Future research to answer these questions will provide us the basis for understanding the mechanism of the reactions that Y-family DNA polymerases were evolved to catalyze.

**ASSOCIATED CONTENT**

- Supporting Information
  Figures showing the CD spectra of wt Dpo4 and several representative mutants and stopped-flow control experiments. This material is available free of charge via the Internet at http://pubs.acs.org.

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**ABBREVIATIONS**

CPM, 7-(diethylamino)-3-(4′-maleimidophenyl)-4-methylcoumarin; Dpo4, *S. solfataricus* DNA polymerase IV; FRET, Förster resonance energy transfer; LF, Little Finger; DNAβ, DNA substrate 21/30-mer containing a 3′-deoxy-terminated 21-mer; DNAβ(H), normal DNA substrate 21/30-mer; wt, wild-type; PDB, Protein Data Bank.

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