Structure and Mechanism of DNA Polymerase β
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ABSTRACT: DNA polymerase (pol) β is a small eukaryotic DNA polymerase composed of two domains. Each domain contributes an enzymatic activity (DNA synthesis and deoxyribose phosphate lyase) during the repair of simple base lesions. These domains are termed the polymerase and lyase domains, respectively. Pol β has been an excellent model enzyme for studying the nucleotidyl transferase reaction and substrate discrimination at a molecular level. In this review, recent crystallographic studies of pol β in various liganded and conformational states during the insertion of right and wrong nucleotides as well as during the bypass of damaged DNA (apurinic sites and 8-oxoguanine) are described. Structures of these catalytic intermediates provide unexpected insights into mechanisms by which DNA polymerases enhance genome stability. These structures also provide an improved framework that permits computational studies to facilitate the interpretation of detailed kinetic analyses of this model enzyme.

DNA polymerases catalyze template-dependent DNA synthesis during genome replication and repair. These enzymes are responsible for preferentially binding and incorporating a nucleotide, from a pool of chemically and structurally similar molecules, that correctly base pairs with the appropriate templating base. DNA polymerase (pol) β has served as a model enzyme for studying this fundamental task, providing a detailed understanding of the events during substrate selection. Pol β is the smallest cellular DNA polymerase (335 residues, 39 kDa) and lacks a 3′ → 5′ proofreading exonuclease activity that enhances the accuracies of replicative DNA polymerases (e.g., pol ε and pol δ). On the basis of its primary sequence, pol β belongs to the X-family of DNA polymerases. This review will highlight recent advances and insights provided by the structural characterization of pol β in various liganded and conformational states. For earlier kinetic and structural descriptions, see previous in-depth reviews.

BIOLOGICAL ROLE

Endogenous and environmental agents continually modify genomic DNA, resulting in physical damage or modification that results in steady-state levels of 50000–200000 apurinic/apyrimidinic (AP) sites per eukaryotic cell. AP sites are generated through spontaneous depurination or lesion specific enzymatic hydrolysis of the N-glycosyl bond between the deoxyribose and base. The rate of spontaneous depurination has been estimated to be ~10⁴ depurinations per cell per day. The base excision repair (BER) pathway (Figure 1) is responsible for removing simple base lesions and AP sites in DNA. Pol β contributes two enzymatic activities, DNA synthesis and deoxyribose phosphate (dRP) lyase, during the repair of AP sites. AP sites represent potentially dangerous lesions because they can be mutagenic and cytotoxic. AP endonuclease 1 incises the AP site, resulting in 3′-hydroxyl and 5′-dRP termini. The dRP group is excised by the lyase activity of pol β, resulting in a 5′-phosphate and a one-nucleotide gap (i.e., a single templating base). Pol β fills this single-nucleotide gap, resulting in nicked DNA that will subsequently be ligated to restore DNA’s native structure.

As discussed in detail below and elsewhere, pol β and other members of the X-family of DNA polymerases have evolved to fill short DNA gaps during essential cellular transactions. In a mouse model system, the loss of pol β results in embryonic lethality; however, cultured embryonic mouse fibroblasts are viable. These cells are hypersensitive to genomic toxins because of the accumulation of cytotoxic repair intermediates. In addition to its enzymatic activities, pol β physically interacts with other key BER factors that hasten repair at AP sites.

The fundamental role that pol β plays in BER and high-fidelity gap-filling DNA synthesis implicates pol β and BER as tumor suppressors. Consistent with this idea is the observation that a high percentage of tumors have variants of pol β. These often have altered fidelity or catalytic activities and can induce cellular transformation. Many of these variants have amino acid changes that are distant from the polymerase or lyase active sites. It remains to be seen whether these alterations affect critical protein–protein interactions necessary for efficient BER or critical protein dynamic behavior that influences catalytic activity and/or fidelity (see below).

MINIMAL REACTION PATH FOR NUCLEOTIDE INSERTION

DNA polymerases are believed to follow the general pathway outlined in Scheme 1. The detailed kinetic and equilibrium constants are sensitive to the identity of the polymerase and
strains (e.g., base and sugar of the nucleoside triphosphate and DNA sequence context). In addition, the nature (kinetic and thermodynamic) of the conformational changes(s) before and after chemistry has been the subject of a long-standing debate as well as their impact on substrate discrimination (i.e., fidelity).14 Steady-state kinetic analyses indicate that pol β follows an ordered binding of substrates;15 after binding DNA (Scheme 1, step 1), DNA polymerases preferentially bind a DNA structure.

Figure 1. Base excision repair. Base excision repair involves the removal of a damaged nucleotide (red pointers) from DNA. It is replaced with an undamaged nucleotide (green pointers). The damaged base is removed by a damage specific DNA glycosylase (1) that hydrolyzes the N-glycosidic bond between the deoxyribose and damaged base. In this image, uracil would be removed by uracil DNA glycosylase. AP endonuclease 1 incises the sugar–phosphate backbone 5′ to the AP site (2). The lyase domain of pol β (3) removes the 5′-dRP group (red pointer), and the polymerase domain inserts a nucleotide in a template-dependent reaction (green pointer). DNA ligase (4) seals the nicked DNA, resulting in restoration of the original DNA structure.

Scheme 1. Minimal DNA Polymerase Reaction Pathway17

"After binding DNA (step 1), the nucleotide triphosphate binds, forming an initial ternary complex (circle, step 2). The polymerase–substrate complex undergoes rapid conformational adjustments that lead to a productive ternary substrate complex (square, step 3). Catalysis (step 4) leads to a postchemistry product ternary complex. Product release occurs concurrent with or following conformational changes (step 5) that facilitate PPᵢ release (step 6). Translocation of the nascent base pair upstream vacating the active site prepares the polymerase for the next insertion event (gray solid line). Alternatively, product DNAᵢ₊₁ may dissociate from the polymerase (step 7), terminating DNA synthesis. While two divalent metals (Mg²⁺) are required for catalysis, they are not explicitly shown in this scheme. See the text for the role of these metals.

Structural Intermediate:

Structural biology offers a unique opportunity to visualize well-populated intermediate states during catalytic cycling at a molecular level. This approach requires that intermediate states be trapped using modified enzymes, inert cofactors, or substrate analogues. Alternatively, reactions can be initiated in the crystal and stopped at defined moments by rapid freezing. In each case, the structures should be interpreted in the context of a kinetic–thermodynamic model for ligand binding and catalysis. This is especially challenging because structural models often represent static snapshots of dynamic events. Computational studies that bridge the observed intermediate states can be very useful in correlating structural and kinetic observations.16 Crystallographic structures of most of the intermediates depicted in Scheme 1 have been determined at high resolution and provide molecular insights by which an induced-fit model bestows substrate specificity (Table 1). We next describe salient features of available crystallographic structures of the intermediates outlined in Scheme 1.

Apoenzyme (I). Controlled proteolytic or chemical cleavage of pol β first demonstrated that it is folded into discrete domains.17 The structure of the unliganded apoenzyme was first reported in 1994.18 It confirmed that the enzyme is folded into two domains: an amino-terminal 8 kDa domain and the 31 kDa polymerase domain (Figure 2). Consistent with sedimentation velocity measurements19 and small-angle X-ray scattering studies of the apoenzyme,20 the global structure displays an extended conformation (Figure 2A). It is now recognized that the dRP lyase activity resides in the amino-terminal domain.21,22 Like other DNA polymerases, the...
polymerase domain has a modular organization with three functionally distinct subdomains. The catalytic subdomain coordinates two divalent metal cations (Mg$^{2+}$) that facilitate DNA synthesis. The other two subdomains are spatially situated on opposite sides of the catalytic subdomain. While the catalytic subdomains of X- and C-family DNA polymerases share structural homology, those from other families (e.g., members of the A- and B-families) exhibit a similar but unique fold. Structures of A-, B-, and Y-family polymerases have likened these enzymes to a right hand with fingers subdomains, respectively, of right-handed DNA polymerases (Figure 2).

### Table 1. Crystallographic Structures of Pol β Catalytic Intermediates

| intermediate (conformation)$^a$ | ligand | PDB entry | refs |
|----------------------------------|--------|-----------|------|
| I (O) | none | 1BPD, 3UXN | 18, 78 |
| II (O) | DNA | 1BPX, 3ISB | 30, 66 |
| III (O) | DNA, dNTP | 4FSN, 4FSO, 4FS$^b$ | 34 |
| IV (C) | DNA, dNTP | 2BPF, 1BPE, 2FMS, 1HUO, 4KLL, 4KLE, 4KL, 3CL$^b$, 3C2M$^b$, 4KLQ$^b$, 4KLQ$^b$, 4KL5$^b$ | 30, 36, 37, 39, 40, 46, 65 |
| V (C) | DNA, PP$^b$ | 1HUZ, 4KLQ, 4KLH, 4KLI, 4KLI, 4KLI$^b$ | 39, 46 |
| VI (O) | DNA, PP$^b$ | 4KLL, 4KLM, 4KLO, 4KLT$^b$ | – |
| VII (O) | DNA, PP$^b$ | 1BPZ, 1TV9, 1TV8, 1KLU$^b$ | 30, 39, 79 |

$^a$The global conformation is specified as open (O) or closed (C).

$^b$Refers to a structure with an active site mismatch.

The catalytic subdomain includes the C-subdomain (catalytic), the D-subdomain (DNA binding), and the N-subdomain (nascent base pair binding) that are equivalent to the palm, thumb, and fingers subdomains, respectively, of right-handed DNA polymerases (Figure 2).

#### Binary DNA Substrate Complex (II).

In contrast to replicative DNA polymerases that are targeted to the growing 3′-terminus at the double-stranded—single-stranded DNA junction, pol β is targeted to the 5′-margin in gapped DNA. In contrast to the extended protein conformation of the apoenzyme, the binding to single-nucleotide gapped DNA results in a doughnutlike protein conformation (Figure 2B). The lyase domain strongly binds the 5′-phosphate or 5′-dRP intermediate in gapped DNA. The product of the dRP lyase reaction generates a 5′-phosphate; transient-state kinetic characterization of the dRP lyase reaction indicates that it is significantly more rapid than single-nucleotide insertion, so that DNA synthesis would occur using a “clean” gap (i.e., the 3′- and 5′-margins in the gap are appropriate for their respective enzymatic activities; DNA synthesis and subsequent ligation).

In the binary one-nucleotide gapped DNA structure, the 5′-phosphate in the gap is hydrogen bonded to Lys35 and Lys68 (Figure 3A). Importantly, pol β binds tightly to the 5′-phosphate only when there is single-stranded DNA adjacent to the 5′-phosphate. This can be nongapped single-stranded DNA because a 3′-terminus is not required for optimal binding. Accordingly, pol β is expected to bind to the 5′-phosphate in a DNA gap of any size. The observation that it will processively (i.e., insert several nucleotides before dissociating from the DNA substrate) fill short gaps (fewer than six nucleotides) suggests that the lyase domain tethering the polymerase domain to the downstream position in gapped DNA. When the primer terminus (i.e., 3′-OH) is within six nucleotides of the 5′-phosphate on the downstream DNA strand, the proximity of the polymerase domain to the primer terminus would hasten processive DNA synthesis.

The α-helical 8 kDa lyase domain includes a structural motif that binds a monovalent metal interacting with the DNA backbone downstream of the single-nucleotide gap (Figure 3B). This helix–hairpin–helix (HhH) motif (residues 55–79) includes Lys72 that serves as the primary amine that forms a Schiff base intermediate during excision of the 5′-dRP moiety in BER. In addition, the D-subdomain interacts with the DNA sugar—phosphate backbone of the duplex DNA upstream of the gap utilizing a second HhH motif (Figure 3B, residues 92–118). This HhH motif also interacts with the primer strand phosphate backbone through a monovalent metal ion. Thus, the two HhH motifs are observed to make DNA backbone interactions with each end of the incised DNA strand. In the structure of the binary gapped DNA complex, the DNA is bent ~90° as it enters the polymerase active site. The sharp bend occurs at the 5′-phosphodiester bond of the templating base. The function of the HhH motifs appears to be a sequence nonspecific phosphate backbone binding motif that stabilizes the pronounced bend observed in the gapped DNA structure. The abrupt bend in the DNA also exposes the terminal base pairs of each DNA duplex that is situated in the gap. His34 of the lyase domain interacts with the first base pair of the downstream duplex, whereas the N-subdomain contributes interactions with the nascent base pair in the closed ternary complex (see below). The altered path of the template strand as it enters the polymerase active site is a general feature observed in most structures of substrate complexes of DNA polymerases.

#### Open Ternary Substrate Complex (III).

Pol β and members of the A- and B-families of DNA polymerases exhibit
a rapid repositioning of the N-subdomain (fingers) upon nucleotide binding to close around the nascent base pair (Figure 4A). In the presence of an incoming dNTP, the closed complex of the wild-type enzyme is stable (i.e., $K_s \gg 1$).33 By weakening interactions of the N-subdomain believed to stabilize the closed complex, an intermediate complex with an incoming nucleotide bound to the open conformation was determined.34 Substituting lysine for arginine at residue 283 (Figure 4B) of $\alpha$-helix N results in a mutant polymerase with moderately reduced catalytic efficiency.35 The structure reveals that the coding template base facilitates binding of the incoming correct dNTP to the open form of pol $\beta$ through Watson–Crick hydrogen bonds (Figure 5A). However, the nascent base pair is severely buckled, because the sugar/
with an inert dUTP analogue (dUMPNPP). Importantly, the primer mechanism for nucleotidyl transfer. This closed structure was trapped complex (PDB entry 2FMS) are consistent with a two-metal are also indicated (T6 and O3α). The templating (coding) nucleotide and primer terminus (D190, D192, and D256) that coordinate active site metals are also indicated. The templating guanine. Asp276 (D276) hydrogen bonds with O3′ of the primer terminus (3′-endo; however, binding of the catalytic magnesium alters the sugar pucker of the primer terminus (3′-endo), thereby repositioning O3′ for in-line attack on the α-phosphate of the incoming nucleotide. These conformational adjustments facilitate binding of the catalytic magnesium necessary to activate the primer 3′-OH and position the α-phosphate of the incoming dNTP. Because the closed ternary substrate complex can be determined without the catalytic magnesium, it is believed that this is the last component that binds to complete the precatalytic complex. This is also consistent with the reported binding affinity for the catalytic metal.43

Recently, ternary complex crystallographic structures of intermediate complexes undergoing nucleotidyl transfer have been captured with natural substrates and metals.39,44 This is achieved by generating crystals of precatalytic substrate ternary complex (PDB entry 2FMS) are consistent with a two-metal mechanism for nucleotidyl transfer. This closed structure was trapped with an inert dUTP analogue (dUMPNPP).37 Importantly, the primer terminus O3′ coordinates the catalytic Mg2+, labeled Mg(C). The catalytic Mg2+ also coordinates all three active site aspartates (purple dashed lines). In this structure, O3′ of the primer terminus is 3.4 Å from the α-phosphate of dUMPNPP. A nucleotide binding metal, Mg(N), coordinates nonbridging oxygens on all three phosphates. The protein coordination of the triphosphate also differs from that observed in the absence of metals. Arg183 now only coordinates the β-phosphate, and Arg149 has lost direct contact with Pγ.

Figure 5. Nucleotide binding to pol β. (A) A ternary substrate complex with the correct incoming nucleotide was trapped in the open conformation using a mutant of pol β that destabilizes the closed conformation.34 In this structure (PDB entry 4FSN), protein side chains coordinate the extended anionic triphosphate moiety of the incoming nonhydrolyzable dCTP analogue, dCMP(CF3)PP. Although the nascent base pair (yellow carbon atoms) is severely buckled, the incoming cytosine base hydrogen bonds (green lines) with the templating guanine. Asp276 (D276) hydrogen bonds with O3′ of the incoming nucleotide, while Arg183 (R183) coordinates nonbridging oxygens on the α-phosphate (Pα) and β-phosphate (Pβ) of the incoming nucleotide. Arg149 (R149) and Gly189 (D189) coordinate the γ-phosphate (Pγ) of the incoming nucleotide. Active site aspartates (D190, D192, and D256) that coordinate active site metals are also indicated. The templating (coding) nucleotide and primer terminus are also indicated (T6 and O3′, respectively). (B) The active site structure and metal coordination of the precatalytic ternary substrate complex (PDB entry 2FMS) are consistent with a two-metal mechanism for nucleotidyl transfer. This closed structure was trapped with an inert dUTP analogue (dUMPNPP).37 Importantly, the primer terminus O3′ coordinates the catalytic Mg2+, labeled Mg(C). The catalytic Mg2+ also coordinates all three active site aspartates (purple dashed lines). In this structure, O3′ of the primer terminus is 3.4 Å from the α-phosphate of dUMPNPP. A nucleotide binding metal, Mg(N), coordinates nonbridging oxygens on all three phosphates. The protein coordination of the triphosphate also differs from that observed in the absence of metals. Arg183 now only coordinates the β-phosphate, and Arg149 has lost its direct contact with Pγ.

triphosphate moieties interact with protein side chains that have not moved to their closed positions. Surprisingly, the negative charge on the incoming nucleotide triphosphate was neutralized through protein interactions (Figure 5A). Although free nucleotides are usually associated with a magnesium ion, the nucleotide–metal complex exists in several coordination states and diastereoisomers. The ability to trap a metal-free complex suggests that the polymerase has a strong influence on metal coordination and triphosphate reorganization. It remains to be determined how the polymerase directs and/or deters (e.g., for an incorrect nucleotide) the metal coordination state of the incoming nucleotide.

Closed Ternary Substrate Complex (IV). Precatalytic closed crystallographic structures have been trapped by employing inert analogues of catalytic participants; the dideoxy-terminated primer (i.e., absence of primer O3′).36 an incoming nucleotide where the bridging oxygen between the α- and β-phosphates has been replaced with an imido or methylene group;37,38 or substituting calcium for magnesium.39 All three approaches have been successfully utilized to trap closed ternary complexes of pol β. The structures show that the N-subdomain has closed around the nascent base pair (Figures 4 and 5). This global structural transition induces many subtle conformational adjustments that lead to conformational activation. One key aspect of this activation is that Asp192 is released from its salt bridge interaction with Arg258, permitting it to coordinate both the catalytic and nucleotide binding metals (Figure 4B). The importance of this conformational activation is illustrated by the observation that mutagenesis of Arg283 and Glu295, which are >10 Å from the active site, dramatically decreases activity and the extent of conformational activation.35,40–42
internal chemical equilibrium). In addition, there are basic side chains in the N-subdomain that stabilize the incoming dNTP in the closed conformation.

Closed Ternary Product Complex (VI). After chemistry, the ternary product complex remains in the closed conformation. However, there are subtle active site changes concomitant with chemistry. The resulting phosphodiester bond alters the coordination state of the catalytic Mg$^{2+}$, hastening its dissociation and its replacement with Na$^+$. In contrast, the nucleotide-associated Mg$^{2+}$ remains in the active site coordinating aspartates and PP$_i$ (Figure 6B). After longer periods of time, the PP$_i$ coordination is altered as one phosphate, the Py remnant, is stripped from the nucleotide metal by a competing water molecule, indicating that metal solvation may play a role in PP$_i$ dissociation and/or subdomain opening. Surprisingly, the product complex after correct nucleotide insertion (i.e., nicked DNA) remains in a closed conformation with bound PP$_i$ even with extended incubations. Because catalytic cycling in solution occurs much more rapidly, other factors such as crystal packing or excessive divalent metals are influencing the open or closed state of the enzyme.

The ternary product complex can also be formed by crystallizing pol $\beta$ with nicked DNA (annealed oligonucleotides) to generate an open binary complex. Addition of PP$_i$ and Mg$^{2+}$ produced a closed ternary product complex similar to that described above in which the nucleotide and product metal sites are occupied with Mg$^{2+}$ and the catalytic metal site is occupied with Na$^+$. In this case, a water molecule (S) replaces the remnant of the $\gamma$-phosphate for nucleotide metal coordination. Hydrogen bonds are displayed as green lines.

![Figure 6. Structures of the closed product complex of pol $\beta$. (A) As the nucleotidyl transfer reaction proceeds in the crystal, an additional divalent metal is observed in the closed product complex. After a 40 s reaction in the crystal (PDB entry 4KLG), the active site structure reveals a new metal [Mg(P)] that bridges the two products, i.e., coordinates nonbridging oxygens on the phosphates of the incorporated dCMP and the remnant $\beta$-phosphate of PP$_i$ (yellow carbons). Water molecules (red spheres) complete the octahedral coordination (purple dashed lines). A sodium ion (purple sphere, Na$^+$) replaces the catalytic magnesium, but the nucleotide-associated Mg$^{2+}$ still coordinates nonbridging oxygens on the phosphates of the products (dCMP and PP$_i$). (B) After an extended reaction in the crystal (45 min, PDB entry 4KLL), the polymerase remains closed, but the PP$_i$ appears to be preparing to dissociate. In this case, a water molecule (S) replaces the remnant of the $\gamma$-phosphate for nucleotide metal coordination.](Image 61x529 to 564x749)

DNA POLYMERASE NUCLEOTIDYL TRANSFERASE

A model for the nucleotidyl transferase enzymatic mechanism is supported by structures of ternary substrate complexes. The chemical mechanism proceeds by an in-line nucleophile attack of the Mg$^{2+}$-activated primer O$_3^-$ anion on the $\alpha$-phosphate of the incoming nucleotide, leading to a pentacoordinated bipyramidal $\alpha$-phosphate transition state. The transition state is resolved by release of PP$_i$ from the opposite side of the attacking nucleophile, resulting in stereochemical inversion about the $\alpha$-phosphorus of the newly incorporated nucleotide. The pol $\beta$ active site includes...
three conserved acidic residues that bind two divalent magnesium ions. One Mg\(^{2+}\) (nucleotide binding metal) coordinates two aspartate residues (Asp190 and Asp192) and the triphosphate moiety of the incoming nucleotide, thereby facilitating nucleotide binding. The other Mg\(^{2+}\) (catalytic) coordinates all three active site aspartates (Asp190, Asp192, and Asp256) and O3' of the primer terminus (Figures 4A and SB). It lowers the pK\(_{a}\) of the primer terminus 3'-OH, hastening attack on P\(_{\alpha}\) of the incoming nucleotide, and serves as a general base on O3' activation.48 Thus, the pK\(_{a}\) values of both the donor (primer O3') and the acceptor group (OD2 of Asp256) are modulated by the catalytic Mg\(^{2+}\).

Site-directed mutagenesis of Asp256 coupled with crystallographic, activity–pH profile, and computational studies confirms that this residue plays a fundamental role in nucleotidyl transfer.49 In this study, quantum calculations revealed the transfer of charge into the catalytic metal and a decrease in the charge of Asp256(OD2) that accompanies the proton jump from the primer terminus (O3') to OD2. In spite of the loss of the proton, the charge on O3' remains almost constant, facilitating its approach toward P\(_{\alpha}\) of the incoming nucleotide. Interestingly, in the structure of a D256E mutant, a water molecule replaces OD2 of Asp256, and this water is seen to coordinate the catalytic Mg\(^{2+}\). However, in quantum calculations, this water does not substitute for Asp256(OD2) in the transfer of charge to the catalytic Mg\(^{2+}\). In this situation, the water molecule jumps to OE1 of Glu256, and the energy barrier for the transition state is much higher than that for the wild-type enzyme.50 The critical role of Asp256(OD2) appears to be facilitated by a stabilizing salt bridge interaction between the water molecule and the nearby Arg254. In the D256E mutant, Arg254 is repositioned and does not interact with Glu256.

## FIDELITY

### Correct Nucleotide Insertion Efficiency Controls Fidelity.

DNA polymerase fidelity, specificity, and discrimination are relative kinetic terms used to describe the probability of a polymerase producing a base substitution error (i.e., mismatch). The base substitution error frequency for DNA replication and repair polymerases is generally between 10\(^{-3}\) and 10\(^{-6}\).50 These frequencies represent one error per 1000 and 1 million nucleotides synthesized, respectively. These levels of discrimination are far greater than those predicted by free energy differences between matched and mismatched DNA termini (predicted error frequency of ~0.4; one error per three nucleotides synthesized), indicating that DNA polymerases enhance fidelity by a large factor.51 DNA polymerase specificity is commonly characterized by determining the misinsertion frequency.52 The misinsertion frequency is the insertion efficiency of an incorrect nucleotide divided by the sum of the insertion efficiencies for incorporation of incorrect and correct nucleotides at the same concentration of nucleotides. Quantitatively, fidelity is simply the reciprocal of the misinsertion frequency. In general, the specificity constants for incorrect nucleotides are much lower than for the correct nucleotide, so that fidelity is simply the ratio of specificity constants ([k\(_{\text{cat}}/K_{m}\)\(_{\text{cat}}\)]/([k\(_{\text{cat}}/K_{m}\)\(_{\text{inc}}\)]). Because relative misinsertion efficiency and fidelity are the same ratio of specificity constants for insertion of correct and incorrect nucleotides, they can be altered by a change in one specificity constant or both.

Importantly, low- and high-fidelity polymerases insert incorrect nucleotides with similar efficiencies.53 Consequently, fidelity is modulated by the efficiency of correct nucleotide insertion, and the molecular strategies that contribute to efficient DNA synthesis are dependent on the specific polymerase. Thus, an understanding of fidelity at the molecular level requires structural insight into the attributes that contribute to correct insertion efficiency rather than the molecular interactions that may occur between a low-fidelity polymerase during incorrect insertion. The lower efficiency and fidelity of DNA synthesis displayed by Y-family DNA polymerases is most likely reflected in the positioning of charged active site residues in the catalytic core.23 As noted previously,50 X- and Y-family DNA polymerases do not exhibit a conserved basic side chain interacting with a nonbridging oxygen of the α-phosphate. The lack of this interaction would be expected to diminish the rate of correct nucleotide insertion, thereby decreasing fidelity.

### Induced Fit.

The induced-fit hypothesis proposes that ligand-induced conformational changes align catalytic groups for optimal activity.54 Poor substrates would deter catalysis through the misalignment of the reactive atoms. As described above, crystallographic structures of polymerase binary DNA and ternary (+dNTP) complexes indicate that the N-subdomain closes around the nascent base pair.55 For 25 years, researchers characterizing the polymerase reaction have attempted to show that correct nucleotide insertion was limited by a nonchemical step. For most DNA polymerases, small elemental effects on the rate of nucleotide insertion of α-thio-substituted nucleotide analogues relative to the natural substrates were provided as evidence that chemistry was only partially rate-limiting. On the basis of model compounds, a significant decrease in rate upon sulfur substitution would suggest that chemistry is rate-limiting. However, there appears to be significant steric considerations, in addition to the electronnegativity of sulfur, that influence the measured rate.56 Following polymerase–DNA conformational changes with fluorescently labeled DNA and the effect of viscosity on both nucleotide insertion and fluorescent transients, the Tsai laboratory has provided compelling evidence that for pol β, chemistry is generally rate-limiting for nucleotide insertion.57–59 Likewise, Sucato et al.60,61 and Oertell et al.62 employing nucleotide analogues that alter the pK\(_{a}\) of the leaving group provide direct kinetic evidence that the rate-determining step during transient-state nucleotide insertion involves bond breaking in the transition state.

A significant effort has been invested in determining the identity of the rate-determining step because it was assumed that a rate-limiting conformational change must limit correct nucleotide insertion for polymerases that utilize an induced-fit model. However, Post and Ray63 have pointed out that induced fit can alter enzyme specificity even when critical conformational changes are kinetically silent (i.e., fast), such as when the transition states for correct and incorrect nucleotide incorporation are unique. Additionally, they show that an induced-fit model reduces specificity. This reduced level of discrimination represents an acceptable compromise for an enzyme such as a DNA polymerase that must select a different and/or new substrate (DNA and dNTP) with each catalytic cycle.

More recently, Tsai and Johnson highlighted that catalytic efficiency (k\(_{\text{cat}}/K_{m}\)) of correct nucleotide insertion is independent of the chemical step (k\(_{\text{cat}}\)) even when it is the slowest step in the forward direction.64 Critically, efficiency is linked to the reversal of the polymerase conformational change (k\(_{\text{−}}\)). Thus, the concentration of complex IV may accumulate
for correct insertion, but not for incorrect insertion (i.e., the concentration of complex III is high because $k_{-3,\text{incorrect}} \gg k_{-3,\text{correct}}$). Consequently, the conformational change insulates the correct nucleotide from dissociating from the ternary substrate complex and commits it to the forward insertion reaction.

**Structural Characterization of a Base Substitution Error.** Crystallographic structures of pol $\beta$ with an incoming incorrect nucleotide indicate that the ternary complex is in a closed conformation (Figure 7A). Using a mutant enzyme that destabilizes the closed conformation of the polymerase is in the closed conformation. The coding template base (T6) is shifted upstream 3.2 Å, while the incoming nucleotide is positioned in the dNTP binding pocket. The $\alpha$, $\beta$, and $\gamma$-phosphates of the incoming nucleotide are denoted P$\alpha$, P$\beta$, and P$\gamma$, respectively. The primer terminus (P10) of the mismatched structure rotates to follow its templating base that has shifted upstream, as the coding templating nucleotide vacates its binding site. This displaces O3$'$ from the primer terminus (highlighted), thereby deterring incorrect nucleotide insertion. In contrast to insertion of the correct nucleotide in the crystal, misinsertion of an incorrect nucleotide results in an open binary complex in which PP$\gamma$ has dissociated (PDB entry 4KLU). The structure of the binary product complex following misinsertion indicates that the enzyme is in the open conformation (pink carbons). In this open conformation, the density for the misinserted nucleotide is poor, indicating that it can assume multiple conformations.

**Figure 7. Intermediate pol $\beta$ structures for insertion of the wrong nucleotide.** (A) Overlay of the ternary substrate complex structure with a correct incoming nucleotide (PDB entry 2FMS, light green carbons) with a precatalytic complex with an active site mismatch (PDB entry 3C2M, yellow carbons; dG-dAMP-CPP$\gamma$). The position of $\alpha$-helix N (ribbon) of the N-subdomain indicates that the polymerase is in the closed conformation. The coding template base (T6) is shifted upstream 3.2 Å, while the incoming nucleotide is positioned in the dNTP binding pocket. The $\alpha$, $\beta$, and $\gamma$-phosphates of the incoming nucleotide are denoted P$\alpha$, P$\beta$, and P$\gamma$, respectively. The primer terminus (P10) of the mismatched structure rotates to follow its templating base that has shifted upstream, as the coding templating nucleotide vacates its binding site. This displaces O3$'$ from the primer terminus (highlighted), thereby deterring incorrect nucleotide insertion. (B) In contrast to insertion of the correct nucleotide in the crystal, misinsertion of an incorrect nucleotide results in an open binary complex in which PP$\gamma$ has dissociated (PDB entry 4KLU). The structure of the binary product complex following misinsertion indicates that the enzyme is in the open conformation (pink carbons). In this open conformation, the density for the misinserted nucleotide is poor, indicating that it can assume multiple conformations.
to that observed for a true abasic site.66 Importantly, the shift in the template strand results in rotation of the primer terminus as it remains hydrogen bonded to its templating base. The rotation of the primer terminus displaces O3′ to a position that deters misinsertion (Figure 7B). Misinsertion would require that the primer terminus sugar realign through transient template strand slippage (i.e., partially subdomain opening) or melting of the primer terminus from the template strand.

In contrast to the stable closed conformation observed after correct insertion, crystallographic structures after nucleotide misinsertion demonstrate that the N-subdomain can open and release PPi and metals.39 The phosphate of the misinserted guanine is repositioned to relieve steric and electrostatic clashes. A structure of a ternary substrate complex with an incoming 8-oxodGTP in the syn conformation paired with adenine (PDB entry 3MBY, yellow carbons).77 In contrast to a staggered arrangement of bases described previously with active site mismatches, the 8-oxodGTP-dA mispair is planar because 8-oxodGTP assumes the syn conformation while the templating deoxyadenine (dA) remains in the anti conformation providing for good Watson–Crick geometry. The syn conformation is stabilized through Hoogsteen hydrogen bonding with the templating adenine (yellow dashed lines) and a hydrogen bond with Asn279 (not shown). The syn conformation of 8-oxodGTP positions O8 in the DNA minor groove in a position similar to that of O2 for a Watson–Crick base pair. The structure of the ternary substrate complex with dUMPnPPP paired with adenine is shown for reference (PDB entry 2FMS, gray carbons).37 Additionally, an intramolecular hydrogen bond between N2 and a nonbridging oxygen on Pro (pro-5′) of 8-oxodGTP could stabilize the syn conformation. (C) Structures of precatalytic ternary substrate complexes of pol β with an incoming CTP (wild-type enzyme, PDB entry 3RH4, light green carbons; Y271A mutant, PDB entry 3RH6, yellow carbons) were superimposed with the wild-type enzyme with an incoming dUMPnPPP (PDB entry 2FMS, gray carbons).37,73 The 2′-ribose oxygen is unusually close to the carbonyl oxygen of Tyr271 (2.54 Å, dashed line) but is well-accommodated in the closed complex of these structures. Replacing the tyrosine side chain with a methyl group (Y271A) provides additional freedom to subtly displace this carbonyl from O2′, resulting in a mutant enzyme that displays a decreased level of discrimination for ribonucleotides.75 (D) Tyr271 hydrogen bonds to the base of the minor groove edge of the primer terminus (dashed black line). Substitution of Tyr271 with alanine results in the loss of this hydrogen bond and displacement of the dideoxy-terminated primer into the major groove. This is illustrated by examining the position of N3 of the guanine base in the structures of wild-type and mutant enzymes with an incoming CTP. The carbonyl of Ala271 is also illustrated.

Oxidative DNA Damage. Environmental and endogenous toxic chemicals lead to oxidative stress that threatens the integrity of genomic DNA. Cells maintain an intricate surveillance system for protecting themselves against adverse genotoxic stress. A major lesion found in DNA and dNTP pools exposed to reactive oxygen species is the promutagenic lesion 8-oxo-7,8-dihydro-2′-deoxyguanosine (8-oxoG). Thus, oxidative stress leads to production of 8-oxodGTP in the dNTP pool.
pool and 8-oxoG in DNA. At neutral pH, the major tautomeric form of 8-oxoG has a carbonyl group at C8 and is protonated at N7. Thus, guanine oxidation results in the alteration of the hydrogen bonding capacity of its Hoogsteen edge. Whereas the unmodified deoxyguanine glycosidic torsion-angle preference is anti, 8-oxoG favors a syn conformation that can form a Hoogsteen base pair with adenine. The altered glycosidic torsion-angle preference is due to steric repulsion between O8 and deoxyribosyl. Although the 8-oxoG(anti)-(anti) base pair does not exhibit Watson–Crick hydrogen bonding, this mispair is well-accommodated within duplex DNA.67

The mutagenic effects of 8-oxoG are mediated by the action of DNA polymerases, because the molecular interactions of 8-oxoG in the confines of the active site can influence its anti–syn conformation. Structural characterization of pol β ternary complex structures with DNA containing 8-oxoG indicates that the glycosidic torsion-angle preference is determined by its base pairing partner, being anti with a complementary cytosine and syn when base-paired with adenine.66–70 The structures reveal that the template binding pocket will permit 8-oxoG to assume an anti or syn conformation and encode incorporation of an incoming cytosine or adenine, respectively. However, the binding pocket for the incoming nucleotide does not have this flexibility, so that insertion of 8-oxodGTP opposite cytosine is strongly discouraged.70

A binary complex crystal structure was obtained with 8-oxoG in the template position of the single-nucleotide gapped DNA substrate.68,70 As expected, the enzyme is in the open conformation. In this binary complex structure with unpaired 8-oxoG, the 8-oxoG base could be modeled into the electron density in both the syn conformation and the anti conformation, indicating that the 8-oxoG is in conformational equilibrium in the absence of enzyme; once the enzyme is bound, the base is sterically restrained from altering its general glycosidic angle. In this case, the 8-oxoG base may be available for pairing with an incoming dATP or dCTP. Similarly, the structure of the phosphate backbone of the 5′-phosphate of the 8-oxoG nucleotide suggests two positions similar to that observed with alternate templating bases [G and 8-oxoG (see below)].

Structures of preinsertion complexes of 8-oxoG paired with an incoming dCTP or ddCTP in the confines of the pol β active site have been determined.68–70 The N-subdomain is in the closed conformation with the bases of the nascent base pair in anti conformations. Only a minor change in the phosphate backbone conformation of the templating 8-oxoG is required to relieve the steric clash of O8 with the sugar–phosphate backbone (Figure 8A). A ternary complex structure of pol β with 8-oxoG as the templating base and an incoming nonhydrolyzable nucleotide analogue, dAMP CPP, has also been determined.70 In this case, the templating base 8-oxoG is in a syn conformation and forms a Hoogsteen base pair with the incoming adenine. The syn conformation of 8-oxoG is stabilized by stacking with an adjacent lysine residue (Lys280).

In contrast to the template binding pocket, there is a severe constraint for the incoming nucleotide for insertion of 8-oxodGTP.70 This is expressed kinetically as moderate insertion efficiency for 8-oxodGTP opposite adenine and a severely reduced efficiency opposite cytosine. The structure of a ternary complex of 8-oxodGTP opposite adenine indicates that 8-oxodGTP assumes the syn conformation and forms a Hoogsteen base pair with the templating adenine (Figure 8B). In contrast to previously published structures of pol β with active site mismatches, this mispair is planar and exhibits Watson–Crick-like geometry, albeit with Hoogsteen hydrogen bonds. The syn conformation of 8-oxodGTP is stabilized through Hoogsteen hydrogen bonding with the templating adenine, a hydrogen bond with Asn279, and an intramolecular hydrogen bond between N2 and a nonbridging oxygen on the α-phosphate.

For most DNA polymerases, 8-oxodGTP is preferentially misinserted opposite adenine rather than cytosine.70 This is consistent with the lack of polymerase crystal structures with 8-oxodGTP paired with a templating cytosine. Modeling an incoming 8-oxodGTP in an anti conformation paired with cytosine indicates that steric repulsion between O8 and its deoxyribose phosphate would distort the active site. Although DNA polymerases can modulate the backbone position of the templating nucleotide, perturbing the position of the α-phosphate of the incoming 8-oxodGTP to accommodate an anti conformation would be expected to severely compromise its insertion.

Recent progress in understanding the structural basis of 8-oxoG mutagenesis by DNA polymerases has provided insight into how the architecture of the DNA polymerase active site is able to adapt to the Hoogsteen base pair. All DNA polymerases can accommodate a Hoogsteen base pair with an 8-oxoG(syn)-A(anti) mispair much more effectively than with a G-A mispair. Ultimately, discrimination will rely on the effect of the anti–syn equilibrium imposed on 8-oxoG by the DNA polymerase active site.

**Sugar Discrimination. **Ribonucleoside triphosphates differ from their deoxynucleotide counterparts by a single atom (oxygen) at C2′ of the sugar, and their cellular concentrations significantly exceed those of dNTPs. Thus, ribonucleotides would be inserted during DNA replication and repair at frequencies much higher than those observed for deoxynucleotides with the wrong base. The presence of a ribose 2′-hydroxyl group stabilizes the glycosyl bond but makes the DNA phosphodiester backbone more susceptible to hydrolysis. Spontaneous or enzyme-catalyzed DNA strand breaks initiate repair and cellular signaling events that would impact overall genome stability and cell survival.

In most instances, DNA polymerases discriminate against ribonucleotide insertion by binding them weakly and inserting them more slowly than their natural substrate.71 Crystallographic structures of substrate complexes of DNA polymerases from different families have indicated that a side chain could stERIC ally interfere with binding of a ribonucleoside triphosphate. This side chain has been termed a “steric gate”.72 In contrast, X-family DNA polymerases deter insertion of ribonucleotides using the protein backbone near the carboxyl terminus of α-helix M [i.e., Tyr271 of pol β (Figure 8C)].73 The backbone carbonyl of Tyr271 would be expected to clash with the hydroxyl group on C2′ of the incoming ribonucleotide. This would be expected to have important consequences because after binding a deoxynucleotide, α-helix M rotates so that Tyr271 forms a hydrogen bond with the minor groove edge of the primer terminus (Figure 8D). Therefore, ribonucleotide binding could alter interactions at the primer terminus transmitted through the altered interactions with α-helix M.

Wild-type pol β inserts ribonucleotides with an efficiency comparable to those of other polymerases that have been examined.71 With a decrease in the size of the side chain of residue 271 by alanine substitution (i.e., Y271A), the level of
pol β ribonucleotide discrimination decreases. Surprisingly, ternary substrate complex structures of wild-type and Y271A pol β with an incoming CTP indicated that the incoming ribonucleotide is well-accommodated in the nascent base pair binding pocket of pol β (Figure 8C). Comparing the crystallographic structures of the wild-type enzyme with bound dCTP or CTP and that of the Y271A mutant with CTP reveals that Tyr271 appears to play two significant roles in ribonucleotide discrimination. The backbone carbonyl at Tyr271 gets unfavorably close to 2′-OH of the ribose in the wild-type structure, and the structure of the mutant indicates that this carbonyl attempts to move farther from O2′ of the incoming ribonucleotide (Figure 8C). Quantum calculations suggest that the energetic cost for the proximity between the backbone carbonyl and ribose O2′ was only ∼2.2 kcal/mol, similar to the observed loss of binding affinity when the wild-type enzyme binds ribonucleotides. Thus, favorable interactions in one region of an enzyme can overcome smaller repulsive interactions in adjacent regions. Significantly, the alanine side chain also removes a hydrogen bond between the tyrosine hydroxyl group and the minor groove edge of the primer terminal base in the closed ternary substrate conformation (Figure 8D). In the wild-type enzyme, this hydrogen bond may alter the active site geometry, thereby deterring insertion of substrates with the wrong sugar. Therefore, discrimination of the ribonucleotide by pol β is attributed to the loss of a contact with the primer terminus and a steric clash between O2′ of the ribose ring and the adjacent polymerase backbone carbonyl.

The dNTP binding affinity for many DNA polymerases is in the range of concentrations found in cells. Although DNA polymerases bind ribonucleotides weakly, their elevated concentration in the cell would permit them to effectively compete for binding. Although ribonucleotides may not be incorporated, they would impact the rate of DNA polymerization. In contrast, because RNA polymerases bind dNTPs weakly and their cellular concentration is low, dNTPs would exert a weaker impact on RNA polymerization.

### FUTURE OUTLOOK

Structural, kinetic, and computational approaches have provided powerful tools for the development of molecular insights into DNA polymerase function. These experimental approaches indicate that dynamic events within the protein, substrates, and cofactors underlie molecular events that hasten correct and deter incorrect DNA synthesis. The observation that additional divalent metals may participate in catalysis, in addition to the two metals that have been traditionally proposed, provides additional motivation to probe chemistry in greater detail. Clearly, the highly charged active site provides an environment that could modulate chemistry through subtle molecular (charge and position) changes. Although the experimental focus has been on the forward DNA synthesis reaction, further characterization of the reverse reaction, pyrophosphorolysis, is warranted, especially because the reverse reaction can play an important role in nucleoside drug resistance. Additionally, solution studies indicate that divalent metals can influence the conformational equilibrium (open—closed) of the pol β DNA binary complex that would influence the distribution of active enzyme species. DNA polymerases often have accessory factors, and our understanding of the kinetic and structural effects these factors have on substrate binding and chemistry is lacking. Likewise, how substrates and/or products are optimally processed through a pathway with several enzymes (substrate channeling) provides important experimental opportunities that will certainly uncover cellular strategies for enhancing enzyme efficiency. The polymerase’s ability to replicate DNA of varying sequence with high fidelity represents an evolutionary achievement that perpetuates life and evolution. The elegant mechanisms that contribute to this elementary reaction are finally being uncovered. Even though structural biology has uncovered key molecular details during DNA synthesis, there are sure to be new details that have not been considered.

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