Substrate Rescue of DNA Polymerase β Containing a Catastrophic L22P Mutation

Thomas W. Kirby, Eugene F. DeRose, William A. Beard, David D. Shock, Samuel H. Wilson, and Robert E. London*

Laboratory of Structural Biology, National Institute of Environmental Health Sciences, Research Triangle Park, North Carolina 27709, United States

Supporting Information

ABSTRACT: DNA polymerase (pol) β is a multidomain enzyme with two enzymatic activities that plays a central role in the overlapping base excision repair and single-strand break repair pathways. The high frequency of pol β variants identified in tumor-derived tissues suggests a possible role in the progression of cancer, making the determination of the functional consequences of these variants of interest. Pol β containing a proline substitution for leucine 22 in the lyase domain (LD), identified in gastric tumors, has been reported to exhibit severe impairment of both lyase and polymerase activities. Nuclear magnetic resonance (NMR) spectroscopic evaluations of both pol β and the isolated LD containing the L22P mutation demonstrate destabilization sufficient to result in LD-selective unfolding with minimal structural perturbations to the polymerase domain. Unexpectedly, addition of single-stranded or hairpin DNA resulted in partial refolding of the mutated lyase domain, both in isolation and for the full-length enzyme. Further, formation of an abortive ternary complex using Ca²⁺ and a complementary dNTP indicates that the fraction of pol β(L22P) containing the folded LD undergoes conformational activation similar to that of the wild-type enzyme. Kinetic characterization of the polymerase activity of L22P pol β indicates that the L22P mutation compromises DNA binding, but nearly wild-type catalytic rates can be observed at elevated substrate concentrations. The organic osmolyte trimethylamine N-oxide (TMAO) is similarly able to induce folding and kinetic activation of both polymerase and lyase activities of the mutant. Kinetic data indicate synergy between the TMAO cosolvent and substrate binding. NMR data indicate that the effect of the DNA results primarily from interaction with the folded LD(L22P), while the effect of the TMAO results primarily from destabilization of the unfolded LD(L22P). These studies illustrate that substrate-induced catalytic activation of pol β provides an optimal enzyme conformation even in the presence of a strongly destabilizing point mutation. Accordingly, it remains to be determined whether this mutation alters the threshold of cellular repair activity needed for routine genome maintenance or whether the “inactive” variant interferes with DNA repair.

The genetic instability that leads to the development of cancer is driven by the accumulation of damaged DNA. Impairment of any of the DNA repair pathways represents one mechanism leading to such an accumulation.1,2 Approximately 30% of human tumors that have been sequenced show elevated numbers of tumor-associated mutations in pol β, an enzyme that is centrally important in the overlapping base excision repair (BER) and single-strand break repair pathways.3–5 A large number of tumor-associated mutations in pol β have been identified, and the functional effects of many of them have been characterized.6–12 Many of these mutations confer mutator phenotypes on the enzyme, and several of these cancer-associated mutations have catastrophic effects on enzymatic activity.13,14 The pol β(E295K) mutation alters a critical residue in the sensor–effector coupling pathway that constitutes the functional connection between correct base pair sensing and catalytic activation. Recent nuclear magnetic resonance (NMR) studies of pol β(E295K) demonstrate the failure of a correctly matched ternary complex to induce the conformationally activated state,13 a result consistent with the extremely low activity found for this mutant.10 In addition to impairing activity, some mutated forms of pol β can have additional consequences resulting from nonproductive binding of damaged DNA intermediates or the displacement of native, wild-type pol β from DNA repair complexes. Pol β is known to form a specific complex with the amino-terminal domain of the scaffold X-ray cross complementing group 1 protein (XRCC1) involving the carboxyl-terminal “N-subdomain” of the enzyme.15,16 Mutations that influence pol β structure or activity but leave the interface with XRCC1 unperturbed can interfere with the normal function of the repair complex. Thus, a binary complex formed between a truncated pol β, polβΔ208–236, and XRCC1 acts as a dominant-negative mutant, and transgenic mice expressing the truncated
form of pol β exhibit a significantly elevated incidence of tumor formation.27

Pol β containing the L22P mutation in the lyase domain (LD) has been identified in cells derived from a gastric carcinoma.6,9 The mutation was found to significantly impair both enzymatic activities; pol β(L22P) exhibits negligible S’-deoxyribose phosphate (DRP) lyase activity, and very low or no polymerase activity. Molecular dynamics simulations indicated that the L22P mutant is characterized by altered packing that results in considerable destabilization.8 In our study, we have utilized NMR spectroscopy to evaluate the structural impact of the L22P mutation. We also have evaluated the effects of the mutation on enzyme activity and substrate binding, as well as the structural effects of substrate interactions on the L22P-perturbed enzyme structure.

# EXPERIMENTAL PROCEDURES

Proteins. [methyl-13C]Methionine-labeled pol β(L22P) and pol β(L22P)LD (residues 1–87) were prepared as described previously21 by growth of plasmid-containing Escherichia coli on a medium containing [methyl-13C]methionine (CIL, Cambridge, MA). The [U-2H], [Ile-δ-13C] samples were expressed in E. coli BL21(DE3) transforms grown in M9 deuterated (99% D2O) medium containing [U-13C]glycerol and 15NH4Cl as the sole carbon and nitrogen sources, respectively; 50 µg per liter of [3,3,4,4-2H4]-1-α-ketobutyric acid was added to the deuterated culture 30 min prior to protein induction by the addition of isopropyl β-D-thiogalactopyranoside. The L22P variants were generated using the QuickChange kit (Stratagene). The protein concentrations were determined using 280 nm extinction coefficients of 20088 M⁻¹ cm⁻¹ for full-length polymerases and 3591 M⁻¹ cm⁻¹ for the isolated lyase domains.

Oligonucleotides. Oligonucleotides for NMR (from Oligosetc or IDT) were dissolved in D2O to make an ~10 mM stock solution. The single-stranded DNA used for the NMR experiments has a 5′-CCG ACG GCG CAT CAG C-3′ sequence. The short hairpin DNA used for the NMR experiments has a 5′-CTG GCG AAG CCA G-3′ sequence. The double-hairpin, one-nucleotide gap DNA substrate used for the NMR experiments has a 5′-CCG ACG GCG CAT CAG C-3′ sequence (the templating T in the gap is underlined). The DNA concentrations were determined using their 260 nm extinction coefficients. The annealing reactions were conducted by incubating a solution of primer with downstream and template oligonucleotides (1:1.2:1.2 molar ratio) at 95 °C for 5 min and cooling the sample at a rate of 1 °C/min to 10 °C in a PCR thermocycler.

DNA Preparation. DNA substrates for single-nucleotide gap-filling DNA synthesis or DRP lyase activity measurements were prepared by annealing three purified oligonucleotides. Each oligonucleotide was suspended in 10 mM Tris-HCl (pH 7.4) and 1 mM EDTA, and the concentration was determined from their UV absorbance at 260 nm. The annealing reactions were conducted by incubating a solution of primer with downstream and template oligonucleotides (1:1.2:1.2 molar ratio) at 95 °C for 5 min and cooling the sample at a rate of 1 °C/min to 10 °C in a PCR thermocycler.

Single-Nucleotide Gap-Filling DNA Synthesis. Steady-state kinetic parameters for single-nucleotide gap-filling reactions at 37 °C were determined by initial velocity measurements as described previously.28 Unless noted otherwise, enzyme activities were determined using a standard reaction mixture containing 50 mM Tris-HCl (pH 7.4, 37 °C), 100 mM KCl, 10 mM MgCl2, 1 mM DTT, 100 µg/mL bovine serum albumin, 10% glycerol, and 500 nM single-nucleotide gapped DNA. For reactions with TMAO, glycerol was omitted. Enzyme concentrations and reaction time intervals were chosen so that substrate depletion or product inhibition did not influence initial velocity measurements. Reactions were stopped with EDTA and samples mixed with an equal volume of formamide dye. The substrates and products were separated on 16% denaturing (8 M urea) polyacrylamide gels. Because a 6-carboxyfluorescein 5′-labeled primer was used in these assays, the products were quantified using the GE Typhoon.
phosphorimager in fluorescence mode. Steady-state kinetic parameters were determined by fitting the rate data to the Michaelis equation. When the observed rates could not be saturated because of poor substrate binding, the data were fit to an alternate form of the Michaelis equation to extract the apparent catalytic efficiency ($k_{cat}/K_M$ best-fit initial slope).

$$k_{obs} = [(k_{cat}/K_M)S]/(1 + S/K_M)$$  \hspace{1cm} (1)

dRP Lyase Activity. To generate a 5’-dRP group, the annealed DNA substrate with a 5’-uracil on the downstream oligonucleotide was treated with 50 nM uracil DNA glycosylase for 30 min at 37 °C. The DNA substrate was stored on ice until it was used. Prior to being used, the DNA substrate was incubated at 37 °C for 5 min before initiation of the reaction with enzyme. The reaction mixture included 50 mM HEPES-KOH (pH 7.5), 20 mM KCl, 0.5 mM EDTA, 1 mM DTT, 10% glycerol, and 100 nM DNA. For reactions with TMAO, glycerol was omitted. Reactions were quenched with freshly prepared cold 200 mM NaBH₄ (final concentration) and mixtures put on ice for at least 30 min. After the mixtures had been briefly heated (5 min at 95 °C), the DNA was resolved on 15% denaturing (8 M urea) polyacrylamide gels. Because a 6-carboxyfluorescein 3’-labeled downstream oligonucleotide was used in these assays, the products were visualized using the GE Typhoon phosphorimager in fluorescence mode.

## RESULTS

It has been reported that pol β(L22P) exhibits no lyase activity and negligible polymerase activity, so it was anticipated that the L22P substitution would significantly perturb the NMR resonances of the pol β lyase domain. A comparison of the $^1H$-$^{15}N$ HSQC spectra obtained for pol β(L22P) with that of pol β reveals that the amide resonances of the catalytic domain show negligible or small chemical shift perturbations, while the amide resonances of the lyase domain are largely absent (Figure 1A,B). The spectra also contain additional intense resonances consistent with substantial, domain-specific unfolding. In harmony with these observations, the $^1H$-$^{15}N$ HSQC spectrum obtained for the isolated mutant pol β lyase domain, LD(L22P), is generally consistent with expectations for a random coil (Figure 1C); the $^1H$ shift range of the amide protons is mostly localized to the region between ~8 and 8.5 ppm. Some resonance broadening is observed, suggesting a dynamic state that may include regions containing some degree of secondary structure. In contrast with these results, the spectrum of the wild-type LD exhibits dispersion typical of a folded domain (Figure 1C).

Previous NMR studies of pol β containing $^{13}C$-labeled methyl groups have proven to be particularly useful for the analysis of substrate binding and conformational activation. The region of the $^1H$-$^{13}C$ HSQC spectrum containing the isoleucine $\delta$-methyl resonances of binary complexes of pol β or the L22P variant with a double-hairpin gapped DNA substrate is shown in Figure 2A. The $\delta^{13}CH_3$-Ile resonances in the LD annotated with blue numbers are essentially missing for the L22P mutant, consistent with the conclusions regarding the loss of structure described above. Alternatively, the $\delta^{13}CH_3$-Ile resonances in the polymerase domain are unaffected (black annotations) or, in some cases, show small but significant shift perturbations (magenta annotations). Importantly, the latter resonances overlap quite well with resonances from uncomplexed wild-type pol β, indicating that there is very little binding of the double-hairpin substrate by the mutant enzyme (Figure 2B). Thus, the apparent indirect chemical shift perturbations of the isoleucine 106, 119, 138, 257, 260, and 277 resonances resulting from the L22P mutation can all be explained by the loss of DNA binding and the accompanying loss of the DNA-induced conformational perturbations (Figure 2C). Interestingly, the $\delta^{13}CH_3$ resonance for Ile97, corresponding to a residue on $\alpha$-helix F, has also disappeared and is annotated in blue (Figure 2A). Although this helix is typically assigned to the DNA-binding subdomain, it interacts closely with helices A and E in the LD,
so it is not surprising that the catastrophic loss of structure of the LD should significantly affect a residue on α-helix F. Examination of the well-resolved amide resonances of the variant enzyme indicates disappearance for residues 99 and 100, but not residue 104, suggesting that from a structural standpoint, the integrity and positioning of α-helix F are strongly dependent on the presence of a folded LD.

**Interaction of DNA with the Pol β LD.** The amino-terminal LD of pol β has a pI of 10.4 and a high affinity for both single- and double-stranded DNA. As shown previously, the methionine methyl resonances in pol β provide useful indicators of both ligand binding and conformational activation, with Met18 specifically sensing interactions with the DNA located immediately downstream of the single-nucleotide gap. As expected from the behavior of amide resonances discussed above, the Met18 methyl resonance of LD(L22P) is strongly shifted from its position in the wild-type domain, exhibiting an intense peak at δ(1H,13C) = (2.07,16.97), close to the expected position for a methionyl methyl resonance in a random coil peptide. Further addition of ssDNA leads to only minimal changes, indicating that the stabilization resulting from ssDNA binding is insufficient to fully compensate for the destabilization produced by the L22P mutation. Analogs results were obtained using hairpin DNA, similar to those used in our previous studies (Figure 3C). Although the intensity of the shifted Met18 resonance in the DNA–LD(L22P) complex is considerably lower than that of the resonance in the uncomplexed domain, the fraction of folded LD(L22P) is higher than a direct comparison of the intensities would indicate, because of the relaxation differences between the folded and unfolded conformations.

**Effects of TMAO on the Conformation of the LD.** TMAO is a naturally occurring organic osmolyte that has been shown to promote protein folding by destabilizing the unfolded state. Titration of a sample of [13CH₃-Met]LD with TMAO results in a loss of intensity of the random coil Met18 resonance, and a parallel increase in the intensity of a broad resonance near the folded position (Figure 3D). Interestingly, the TMAO titration does not produce a concentration-dependent shift perturbation, but rather a shift in the ratio of intensities of the two observed methionine methyl resonances. These results are consistent with a two-state conformational equilibrium of the LD(L22P), in which the primary effect of the TMAO is to alter the relative stability of the two component states. In comparison with the effects of the DNA ligands shown in panels B and C of Figure 3, the TMAO-induced Met18 resonance is broader and further shifted from its position in the wild-type LD, and the Met18 peak at the random coil position is much more effectively eliminated. Despite these changes suggesting that TMAO promotes folding of the LD(L22P) to a conformation similar to that of the wild-type domain, the 1H−15N HSQC spectrum of the [U-15N]-LD(L22P) shows considerably less dispersion and uniformity than the spectrum of the domain lacking the mutation (Figure S1 of the Supporting Information). In general, the effects observed are consistent with studies indicating that the primary effect of TMAO is destabilization of the random coil form of the domain as a result of unfavorable interactions between the TMAO and the peptide backbone.
Substrate-Induced Folding and Conformational Activation of Pol β(L22P). In response to the addition of its substrates, pol β undergoes a series of conformational changes that are conveniently monitored with the methionine methyl resonances. The $^1\text{H}$$-^{13}\text{C}$ spectrum of $[^{13}\text{CH}_3\text{-Met}]\text{pol β}(L22P)$ is very similar to that of the wild-type enzyme, except that the Met18 resonance typically observed at $\delta(^1\text{H},^{13}\text{C}) = (1.66,16.10)$ is replaced by an intense resonance at $\delta(^1\text{H},^{13}\text{C}) = (2.08,16.97)$, as seen for the isolated LD (Figure 4, black spectrum). Addition of a single-nucleotide gapped double-hairpin DNA substrate to $[^{13}\text{CH}_3\text{-Met}]\text{pol β}(L22P)$ produces a typical, small shift of the Met236 resonance. For Met18, two resonances are observed: a sharp and intense resonance near the random coil position and a broader resonance near the position expected for Met18 in the wild-type DNA complex (Figure 4, red spectrum). On the basis of comparisons of the methionine resonance intensities, we estimate that in the presence of the double-hairpin DNA, $\sim 20\%$ of the pol β(L22P) has adopted a folded state approximating that of the wild-type enzyme. Because the DNA-stabilized pol β(L22P) contains the mutation, the substrate-rescued conformation is expected to approximate rather than duplicate that of the wild-type enzyme.

Next, an abortive ternary complex was formed by addition of complementary dATP and CaCl$_2$ to the binary pol β(L22P)·DNA complex (Figure 4, blue spectrum). This strategy represents an alternative to that used previously, in which gap-filling synthesis was blocked using either a deoxy-
terminated primer or a nonhydrolyzable dNTP analogue. The use of Ca^{2+} represents an attractive alternative because it promotes a ternary complex with the true enzyme substrates but fails to support catalysis. Although in the binary complex only the Met18 resonance exhibits two peaks, formation of the ternary complex results in two sets of resonances for each methionine methyl group, except for Met191. The methionine resonance shifts indicate that in the presence of these substrates, the fraction of the pol β-DNA-dATP ternary complex in which the LD has been conformationally rescued by the DNA substrate undergoes further conformational activation similar to that of the wild-type ternary complex. In particular, the abortive ternary complex is characterized by a downfield shift of the Met158 resonance in the $^{13}$C dimension, and an upfield shift of the Met282 resonance in the $^1$H dimension. As discussed previously, this $^1$H shift is consistent with the transition to a closed enzyme conformation and the increased proximity of the Met282 methyl group to Phe320.

Although in previous studies of the wild-type enzyme, the Met18 resonance was observed to respond to the DNA substrate but not to the formation of the ternary complex, the Met18 resonance of pol β(L22P) undergoes an additional shift change upon ternary complex formation (Figure 4). Apparently, the reduced stability of the LD in the mutant enzyme makes the domain conformation more susceptible to the perturbation produced by ternary complex formation.

One other characteristic worth noting is the observation of a shifted Met155 resonance in the ternary complex. Previous studies of Mg-containing ternary complexes did not result in an observable Met155 peak, which was too severely broadened to permit unequivocal observation. In contrast, the Ca^{2+}-containing ternary complex appears to provide greater stabilization of the closed enzyme conformation, so that the shifted resonance for Met155 can be observed. This observation is independent of the presence of the L22P mutation, and we have made similar observations in other studies of Ca^{2+}-containing pol β ternary complexes (unpublished results). For the Ca^{2+} ternary complex, Met191 is also subject to a small resonance shift. Overall, these studies demonstrate that the DNA substrate is able to induce a folded conformation of the mutated LD that approximates that of the wild-type enzyme. Formation of an abortive ternary complex is then able to induce the conformationally activated state in a fraction of the substrate-rescued enzyme. A larger fraction of the enzyme contains an apparently unfolded LD, binds DNA poorly, and fails to undergo conformational activation (Figure 4).

A comparison of the $^1$H–$^{13}$C HSQC spectra obtained for ternary complexes of the wild-type and L22P [13CH3-Met]pol β mutant is shown in Figure 5. In both cases, the abortive ternary complex exhibits the resonance character-

of the conformationally activated state. In contrast, the methionine methyl resonances in the pol β(L22P)-DNA-Ca^{2+}dATP complex are split into two components that we assign to species containing either the disordered or the folded LD. In the complex formed with wild-type pol β, the resonances arising from Met155, Met158, and Met282 are all shifted to positions that correspond to the closed enzyme structure, while in the complex formed with the L22P mutant, each of these residues gives rise to both a shifted resonance and an unshifted resonance. For these three pairs of resonances, the intensity ratio of the shifted to unshifted peaks provides an estimate of the fraction of active to inactive enzyme complex. Evaluation of peak intensities is consistent with ~20–40% folded enzyme complex.

**Figure 5.** Spectral comparison of abortive ternary complexes of wild-type and L22P pol β. Overlaid $^1$H–$^{13}$C HSQC spectra of 100 μM [methyl-$^{13}$C]methionine-labeled pol β with a one-nucleotide gap DNA substrate and dATP in the presence of CaCl2 (magenta) and 100 μM [methyl-$^{13}$C]methionine-labeled pol β L22P variant with a one-nucleotide gap DNA substrate and dATP in the presence of CaCl2 (blue). Samples were in 50 mM Tris-d$_1$ (pH 7.6), 150 mM KCl, 1 mM CDTA, 10 mM NaN$_3$, (pH 7.6), and 100% D$_2$O and run at 25°C.

**Effect of the L22P Mutation on DNA Synthesis and dRP Lyase Activities.** Not unexpectedly, the L22P mutant exhibits very low activity when it is assayed at substrate concentrations that would saturate the wild-type enzyme. However, attempts to measure steady-state kinetic constants showed that the activity of the mutant enzyme increases approximately linearly with increasing DNA and dNTP concentrations, reaching a level similar to that of the saturated wild-type enzyme (Figure 6). This behavior is consistent with a DNA binding defect in pol β(L22P) rather than a nucleotide insertion deficiency. The loss of catalytic efficiency can be completely ascribed to the higher apparent $K_{M_{dNTP}}$, characterizing the mutant enzyme (Table 1). An increase in $K_{M_{dNTP}}$ is expected for a DNA polymerase that binds DNA more weakly (i.e., lower processivity because of an increase in its dissociation rate constant). Although it is difficult to saturate the L22P mutant with substrates, an apparent catalytic efficiency ($k_{cat}/K_{M_{dNTP}}$) can be easily quantified. In the presence of 1 μM DNA, the apparent efficiency is 0.69 × 10$^{-3}$ μM$^{-1}$ s$^{-1}$, more than 1000-fold lower than that of wild-type pol β [dG-dCTP (Table 1)].

The amino-terminal LD is responsible for targeting pol β to gapped DNA substrates bearing a 5′-phosphate or dRP group. During base excision repair, the LD removes the 5′-dRP group generating a 5′-phosphate required for DNA ligation after DNA gap-filling synthesis. The destabilizing effect of the L22P mutation on the structure of the LD would also be expected to strongly reduce the dRP lyase activity of pol β. A qualitative dRP activity assay indicates that while activity is reduced, it is not eliminated (Figure 7). This residual activity is consistent with DNA-induced folding of the mutated domain.

Because TMAO appears to stabilize the folded form of the lyase domain of the mutant enzyme, activity was measured in the presence of TMAO. In the presence of 1 M TMAO, steady-
state kinetic parameters for gap-filling DNA synthesis were easily quantified (Table 1). The apparent binding affinities for substrates are increased significantly, resulting in ~20-fold increases in catalytic efficiencies. Higher concentrations of TMAO do not increase activity. The fidelity [i.e., \( \frac{k_{\text{cat}}}{K_{M,dCTP}}/\frac{k_{\text{cat}}}{K_{M,dTTP}} \)] of the mutant enzyme is similar to that of the wild-type enzyme in the presence of TMAO, suggesting that the L22P substitution does not influence nucleotide discrimination. Interestingly, the fidelity of the wild-type enzyme is significantly reduced in the presence of TMAO (Table 1; −TMAO, 10175; +TMAO, 1430). With respect to dRP lyase activity, the activity of the isolated L22P LD is increased significantly in the presence of TMAO (Figure 7), whereas the activity of the wild-type LD is moderately decreased.

**DISCUSSION**

DNA pol β is composed of two domains that complement its biological function in base excision repair. The amino-terminal 8 kDa LD removes the 5′-deoxyribose phosphate generated after incision by apurinic/apyrimidinic endonuclease during the repair of simple base lesions. The LD recognizes the 5′-phosphate in DNA gaps, thereby targeting the polymerase for gap-filling DNA synthesis. The nucleotidyl transferase activity

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**Table 1. Steady-State Kinetic Parameters for Single-Nucleotide Gap-Filling DNA Synthesis**

| enzyme | incoming nucleotide | TMAO (1 M) | \( K_{M,dNTP} \) (μM) | \( k_{\text{cat}} \) (s\(^{-1}\)) | \( k_{\text{cat}}/K_{M,dNTP} \) (×10\(^{-3}\) μM\(^{-1}\) s\(^{-1}\)) | fidelity\(^b\) |
|--------|------------------|-----------|-----------------|-----------------|-----------------|------|
| WT     | dCTP             | −         | 1.18 (0.08)     | 0.96 (0.07)     | 814 (80)        | −    |
| WT     | dCTP             | +         | 0.26 (0.07)     | 0.26 (0.06)     | 1000 (355)      | −    |
| WT     | TTP              | −         | ND\(^d\)       | ND\(^d\)       | 0.08 (0.02)     | 10175 |
| WT     | TTP              | +         | 125 (25)        | 0.088 (0.002)   | 0.7 (0.1)       | 1430 |
| L22P   | dCTP             | −         | ND\(^d\)       | ND\(^d\)       | 0.69            | −    |
| L22P   | dCTP             | +         | 38 (2)          | 0.51 (0.03)     | 13 (1)          | −    |
| L22P   | TTP              | −         | NA\(^c\)       | NA\(^c\)       | NA\(^c\)       | NA\(^c\) |
| L22P   | TTP              | +         | ND\(^d\)       | ND\(^d\)       | 0.0042          | 3100 |

\(^a\)The templating base in the gap is guanine. When standard errors are given, the results represent the mean of at least two independent determinations. \(^b\)Fidelity = \[\frac{(k_{\text{cat}}/K_{M,dCTP})}{(k_{\text{cat}}/K_{M,dTTP})}\]. \(^c\)Wild-type enzyme. \(^d\)Not determined because of weak substrate binding. In this situation, the concentration dependence of the observed activities was fit to eq 1. \(^e\)No activity was observed.

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**Figure 6.** Steady-state kinetic characterization of pol β(L22P). The left panel shows the DNA concentration dependence of the observed rate of insertion of dCMP opposite guanine in a single-nucleotide gapped DNA substrate. The concentration of dCTP was 1 mM. The right panel shows the dCTP concentration dependence of the observed rate of insertion of dCMP opposite guanine. The concentration of the single-nucleotide gapped DNA substrate was 1 μM. Because the observed rate increased in an approximately linear fashion with substrate concentration, the data were fit to a hyperbolic equation to extract the best-fit initial slope (i.e., apparent catalytic efficiency, gray line; see Experimental Procedures).

**Figure 7.** Influence of TMAO on the dRP lyase activity of the wild-type and L22P LD. A 5′-uracil-containing downstream oligonucleotide labeled at its 3′-end with 6-FAM (U, lane D) was treated with uracil DNA glycosylase as described in Experimental Procedures to create a substrate (S) for the dRP lyase reaction. The 5′-terminal dRP-containing oligonucleotide migrates farther than the U-containing strand. Removal of the dRP group results in a shorter product (P). The dRP lyase reaction was monitored for 5 and 10 min in the absence (−) or presence (+) of 2 M TMAO with 50 nM enzyme. The last lane included 500 nM wild-type (WT) enzyme, demonstrating complete conversion of substrate to product.
of pol β resides in the 31 kDa carboxyl-terminal polymerase domain. A helix–hairpin–helix structural motif is found in each domain and interacts with the DNA backbone in a non-sequence-dependent manner on opposite sides of gapped DNA. Genomic mutations that alter DNA repair systems are a common feature of cancer cells, and changes in the DNA repair enzyme pol β have been identified in several cancers.3−5,9,12 A leucine to proline change at residue 22 of pol β was isolated from a gastric carcinoma.2 The change is situated in the middle of α-helix A in the amino-terminal LD and forms part of a hydrophobic core. The catastrophic structural effect resulting from the L22P mutation was not totally unexpected. It is well-known that proline residues disfavor interior positions in α-helices;40 in the referenced study, the authors were not able to identify any examples of α-helices containing proline at an interior position. An analogous unfolding effect has been reported to result from the introduction of a T62P substitution in α-helix A of the model enzyme, staphylococcal nuclease.20 These profound perturbations highlight the delicate balance that exists between the folded and unfolded states of proteins. The ability of single- or double-stranded DNA to partially rescue the conformation of the unfolded lyase domain was, however, unexpected. Apparently, the binding energy is able to provide some but not all of the energy required to compensate for the destabilizing effect of the mutation. Crystallographic structures of pol β-DNA binary complexes indicate that the short downstream fragment of gapped DNA interacts with residues on α-helices B−D, but not directly with helix A. An even stronger effect might have been observed if the DNA interacted directly with residues on this helix.

In this series of solution NMR studies, the Met18 methyl resonance was found to provide a particularly useful readout for the structural status of the LD, both in isolation and in the full-length protein. From the perspective of Met18, both pol β(L22P) and the LD(L22P) behave as bistable systems, with folded and unfolded fractions that are dependent on the presence of single- or double-stranded DNA. Additionally, the folded/unfolded ratio can also be influenced by other environmental factors such as TMAO. In both cases, the exchange between the folded and unfolded states is apparently slow on the NMR time scale, consistent with an interconversion rate below ~10^6 s^−1. The NMR data presented here indicate that the conformational rescue by DNA is based more on its effective stabilization of the folded LD, while the effect of the TMAO results more from its destabilization of the unfolded LD, as has been observed for other proteins.18,19,35 Thus, the DNA complex with LD(L22P) appears to provide a closer approximation of the wild-type structure. The ability of the TMAO to enhance the activity of pol β(L22P) indicates that there is some cooperativity between the effects of the cosolvent and the substrate. A reasonable interpretation of this result is that the TMAO stabilizes a conformational ensemble that more closely approximates that of the wild-type enzyme, providing a more substrate-accommodating binding site than the structure with an unfolded LD.

In contrast with the analysis based on the methionine resonances, a 1H−15N HSQC spectrum of the [U-15N]LD-(L22P) indicates that even 3 M TMAO is unable to compensate for the effects of the L22P mutation (Figure S1 of the Supporting Information). The more dynamic structure of the LD(L22P) indicated by the 1H−15N HSQC spectrum is consistent with the broader Met18 methyl resonance of the folded structure observed in the presence of TMAO (Figure 3D). For this system, the observation of two resolved Met18 resonances provides a useful basis for determining the ratio of folded to unfolded states, while the amide HSQC spectrum does not lend itself to such a direct evaluation.

Two models are generally considered for ligand-assisted protein folding: (1) an induced-fit model in which binding of a ligand to a catalytically inactive conformation results in a catalytically active conformation41 and (2) a conformational selection model in which binding of a ligand to a minor population exhibiting a relatively high ligand affinity shifts the equilibrium to a folded state.52 The substrate-induced rescue of full-length pol β(L22P) is consistent with the induced-fit model. However, because the ligand-binding competent population is expected to be small and difficult to detect, some degree of conformational selection cannot be fully discounted. More recently, an extension of the conformational selection model that also incorporates an induced-fit component has been proposed.43 Although for DNA polymerases, induced fit generally has been discussed in terms of substrate specificity (i.e., right and wrong nucleotide selection),44 binding of a ligand to unfolded or partially unfolded proteins (e.g., intrinsically disordered proteins) alters their conformational distribution, and this type of ligand-induced folding is significant for many cellular interactions. Initial DNA gap binding by pol β is expected to occur through the LD that recognizes the 5′-phosphate in gapped DNA.53 In the absence of a folded LD, the level of binding of DNA to pol β(L22P) is reduced, thereby reducing the catalytic activities even in the presence of DNA. In addition, nucleotide binding is also affected. Generally, the binary DNA complex exists in an open conformation where the nucleotide binding pocket is exposed. Binding of a nucleotide results in a conformational change (repositioning of the N-subdomain of the polymerase domain) that provides the enzyme the opportunity to probe proper base pair geometry. Normally, the LD forms intimate contacts with the N-subdomain that are necessary for a stable ternary complex. In the absence of a properly folded LD, the catalytic efficiencies for nucleotide insertion and nucleotide binding are strongly compromised (Table 1). It previously has been suggested that the L22P pol β variant can produce a mutator phenotype,6 and this could be a direct consequence of the expression of a variant that exhibits low fidelity. It is already known that many of the cancer-associated variants of pol β that produce structural perturbations much less profound than those produced by the L22P mutation can significantly reduce polymerase fidelity. The very low activity of the variant and its limited ability to insert incorrect nucleotides suggest that poor fidelity is not the direct cause of a mutator phenotype in this case, as the fidelity of the variant in the presence of TMAO was similar to that of the wild-type enzyme. Alternatively, a reduced level of DNA repair, expected because of the loss of activity, and/or the possibility that the variant interferes with DNA repair (trans-dominant inhibition), e.g., by forming inactive complexes with XRCC1, may be expected to indirectly increase the level of mutations.

## ASSOCIATED CONTENT

### Supporting Information

Effects of 2 and 3 M TMAO on the 1H−15N HSQC spectra of the LD(L22P) (Figure S1). This material is available free of charge via the Internet at http://pubs.acs.org.
Corresponding Author
*Address: 111 TW Alexander Dr., P.O. Box 12233, MD MR-01, Research Triangle Park, NC 27709. E-mail: london@niehs.nih.gov. Phone: (919) 541-4879.

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Notes
The authors declare no competing financial interest.

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ABBREVIATIONS
5′-dRP, 5-deoxyribose phosphate; BER, base excision repair; DTT, dithiothreitol; LD, lyase domain; pol β, DNA polymerase β; TMAO, trimethylamine N-oxide; XRCC1, X-ray cross complementing group 1 protein.

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