Quantifying the energetic contributions of desolvation and $\pi$-electron density during translesion DNA synthesis

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ABSTRACT

This report examines the molecular mechanism by which high-fidelity DNA polymerases select nucleotides during the replication of an abasic site, a non-instructional DNA lesion. This was accomplished by synthesizing several unique 5-substituted indolyl 2'-deoxyribose triphosphates and defining their kinetic parameters for incorporation opposite an abasic site to interrogate the contributions of $\pi$-electron density and solvation energies. In general, the $K_{d,\text{app}}$ values for hydrophobic non-natural nucleotides are $\sim$10-fold lower than those measured for isosteric hydrophilic analogs. In addition, $k_{\text{pol}}$ values for nucleotides that contain $\pi$-electron densities are slower than isosteric analogs possessing higher degrees of $\pi$-electron density. The differences in kinetic parameters were used to quantify the energetic contributions of desolvation and $\pi$-electron density on nucleotide binding and polymerization rate constant. We demonstrate that analogs lacking hydrogen-bonding capabilities act as chain terminators of translesion DNA replication while analogs with hydrogen bonding functional groups are extended when paired opposite an abasic site. Collectively, the data indicate that the efficiency of nucleotide incorporation opposite an abasic site is controlled by energies associated with nucleobase desolvation and $\pi$-electron stacking interactions whereas elongation beyond the lesion is achieved through a combination of base-stacking and hydrogen-bonding interactions.

INTRODUCTION

The central dogma of DNA replication is that the efficiency and fidelity of this process is dependent upon the formation of correct hydrogen-bonding interactions between an incoming nucleotide and its templating partner (1). The mutual recognition of adenine (A) by thymine (T) and of guanine (G) by cytosine (C) involves the ability of the DNA polymerase to consummate the formation of precise hydrogen-bonding interactions between each partner. The specific pairing of these hydrogen bonding groups provide high-fidelity DNA polymerases with enough flexibility to accurately recognize four distinct pairing partners (A:T, G:C, T:A and C:G), yet remain stringent enough to maintain exquisite genomic fidelity.

Under normal conditions, replicative DNA polymerases use hydrogen-bonding interactions to generate low error frequencies of approximately one mistake every $10^6$ opportunities (2–4). However, polymerization fidelity and efficiency can become altered when the chemical composition of the templating strand is modified by DNA damaging agents such as methylmethane sulfonate and nitrosurea (5). Since these agents alter hydrogen-bonding information, the risk of mutagenesis increases as the frequency of misincorporation events is enhanced. Mutagenesis, in some cases, can benefit an organism by generating genomic diversity required for evolution and/or adaptation (6). However, mutagenesis in humans is typically associated with disease development, most notably cancer (7,8). To better understand the underlying origins of disease development, it is important to define how genomic errors are generated by misreplicating damaged DNA.

One commonly formed DNA lesion is an abasic site that, due to the lack of coding information, is highly promutagenic (9). Although abasic sites are non-instructional, high-fidelity DNA polymerases such as the bacteriophage T4 DNA polymerase preferentially utilize dATP (10). This phenomenon is commonly referred to as the ‘A-rule’ of translesion DNA synthesis (11–18). While the mutagenic consequences of the ‘A-rule’ are well documented (19–21), the mechanistic basis for the preferential utilization of dATP has not been firmly established. Our investigations

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of the ‘A-rule’ have focused on characterizing the kinetic behavior of the library of non-natural nucleotides illustrated in Figure 1A during insertion opposite this DNA lesion (22–28). While these analogs mimic the core structure of dATP, the introduction of various functional groups to the 5-position of the indole ring provides a way to systematically modulate biophysical features such as shape, size, hydrophobicity, dipole moment and π-electron surface area. The kinetic parameters for their utilization were used to develop a structure–activity relationship (SAR) defining key biophysical features required for optimal incorporation opposite an abasic site (23). The parabolic nature of this plot reveals that the catalytic efficiency \(k_{\text{pol}}/K_{\text{d, app}}\) for nucleotide incorporation is dependent upon an optimal π-electron surface area that most closely mimics that of a natural Watson–Crick base pair.

This article further examines this model by quantifying the energetic contributions required for nucleotide binding and insertion of isosteric analogs of two unique non-natural nucleotides, 5-NITP (22) and 5-PhITP (27), that are themselves incorporated opposite an abasic site with high catalytic efficiencies \(>10^6\) M\(^{-1}\)s\(^{-1}\). The strategy employed here is to modulate the degree of π-electron density and hydrophobicity without compromising the overall shape, size or conformation of the parental nucleotide. Kinetic analyses indicate that the binding affinity of the incoming nucleotide, designated as \(K_{\text{d, app}}\), is linked with the overall shape and hydrophobicity of the incoming nucleotide whereas the maximal rate constant for incorporation or \(k_{\text{pol}}\) value depends heavily on the presence of π-electron density. These results are used to propose an enzymatic mechanism accounting for the roles of desolvation, steric fit and π-electron density during translesion DNA synthesis. In addition, elongation beyond the abasic site is dependent upon a combination of base-stacking properties and hydrogen-bonding potential of the nucleotide that is paired opposite the DNA lesion. This final aspect has important implications toward the rational design of novel nucleotides that can be used as chemical probes for DNA synthesis, for expanding the genetic code and/or as potential therapeutic agents.

**MATERIALS AND METHODS**

Reagents and general methods: anhydrous sodium methoxide, phosphoryl oxychloride, tetrakis triphenylphosphine palladium \([\text{Pd}(\text{PPh}_3)_4]\) and 1,8-bis(dimethylamino)naphthalene were purchased from ACROS. Sodium hydride (60% dispersion in mineral oil), tributylammonium pyrophosphate and dimethoxyethane (DME) were purchased from Sigma-Aldrich. Trimethylphosphate and tributylamine were purified via fractional distillation under reduced pressure. Acetonitrile was purified by refluxing with CaH\(_2\) for 2 h followed by simple distillation. DMF was purified by distillation with ninhydrin. All anhydrous solvents were dried over 4 Å molecular sieves and stored under argon. All other solvents and reagents were purchased at the highest purity. Water-sensitive reactions were performed in oven-dried glassware under argon. The intermediates and products were characterized by nuclear magnetic resonance (NMR), UV–vis absorbance and mass spectrometry. \(^1\)H, \(^31\)P and \(^19\)F NMR spectra were recorded in a Varian-400 FT NMR spectrometer. All chemical shifts are reported in ppm and the coupling constants are in Hertz (Hz). \(^1\)H NMR spectra were taken in deuterated chloroform (CDCl\(_3\)), or dimethylsulfoxide (DMSO-d6) using tetramethylsilane (TMS) as the external reference. \(^19\)F-NMR spectra were taken in CDCl\(_3\) or D\(_2\)O and trichlorofluoromethane was used as the external standard. \(^31\)P-NMR spectra were taken in D\(_2\)O with 50 mM Tris–HCl and 2 mM EDTA and phosphoric acid (85%) was used as the external standard. High-resolution electrospray ionization mass spectrometry (Hi-Res ESI-MS) was performed on an IonSpec HiRes ESI-FTICRMS at the University of Cincinnati Mass Spectrometry facility. The configuration of the nucleoside was confirmed by \(^1\)H NMR NOE difference spectroscopy on a Varian Inova 600 NMR instrument. Thin layer chromatography was carried out on a Whatman Silica Gel UV\(_{254}\) plates. Column

![Figure 1. (A) Library of non-natural nucleotides used to interrogate the mechanism accounting for the preferential utilization of dATP during the replication of an abasic site. For clarity, only the nucleobases of the 5-substituted indolyl nucleotides are shown. (B) SAR highlighting the importance of π-electron surface area of the incoming nucleotide for efficient incorporation opposite an abasic site. See text for details.](image-url)
chromatography was performed on a Fisher Scientific Silica Gel, sizes 32-63. HPLC analysis and purification were carried out using the JASCO Model PU-2089 instrument with the reverse phase C-18 Vydac column. UV absorption of the nucleosides used for their quantification was performed on a Cary 50 spectrophotometer.

**Method A: synthesis of 5-substituted indolyl 2'-deoxynucleosides**

To a solution of 5-substituted indole nucleobase (1 eq) dissolved in dry acetonitrile (50 ml) was added NaH (60% dispersion in mineral oil) (5 eq). The reaction mixture was stirred at room temperature for an hour under argon prior to the addition of the Hoffer’s chlorosugar (1.5 eq) prepared using established protocols (29,30). The resulting mixture was further stirred at room temperature under argon for 16 h and immediately rota-evaporated to dryness. The remaining solid residue was again stirred for 16 h at room temperature under argon prior to the addition of anhydrous sodium methoxide until the pH of the reaction mixture reaches 12. The reaction mixture was stirred at room temperature for an hour under argon before being evaporated to dryness under reduced pressure. The crude product was purified by silica flash column chromatography.

**Synthesis of 1-(2-deoxy-β-D-erythropentafuranosyl)-5-methylcarboxylate-1H-indole 1.** This compound was prepared using method A with 5-methylcarboxylate indole (300 mg, 1.7 mmol, 1 eq) as the starting material. The crude product was purified by silica flash column chromatography (10–25% methanol in chloroform). The isolated product was a light brown foam and the yield was 330 mg. 1H NMR (DMSO, 400 MHz, 298 K, TMS) δ: 2.23–2.28 (m, 1H, 2'-H), 2.47–2.54 (m, 1H, 2'-H), 3.48–3.56 (m, 2H, 5'-H), 3.83–3.85 (m, 1-H, 4'-H), 3.85 (s, 3H, –OCH3), 4.35–4.36 (m, 1H, 3'-H), 4.93 (t, J = 5.93 Hz, 1H, 5'-OH), 5.32 (d, J = 4.1 Hz, 1H, 3'-OH), 6.43 (Japp, J = 6.8 Hz, 1H, 1'-H), 6.67 (d, J = 3.13 Hz, 1H, Ar), 7.69 (d, J = 8.9 Hz, 1H, Ar), 7.73 (d, J = 3.5 Hz, 1H, Ar), 7.77 (dd, J = 8.7 Hz, 1.6 Hz, 1H, Ar), 8.25 (d, J = 1.1 Hz, 1H, Ar). UV (MeOH) λmax (nm) 240 (ε = 104 591 cm⁻¹ M⁻¹). HiRes ESI-MS (+): calculated mass spectrum formula C13H17NO5 for M+Na: 314.1004; experimental mass spectrum: 314.1017.

**Synthesis of 1-(2-deoxy-β-D-erythropentafuranosyl)-5-carboxylate-1H-indole 2.** This compound was prepared starting from 1 (204 mg, 0.7 mmol) dissolved in methanol (10 ml). To this reaction mixture was added NaN3 (220 mg, 5.5 mmol) solution dissolved in 1:1 H2O:MeOH solvent (10 ml). The reaction mixture was stirred vigorously at room temperature and the reaction was monitored by TLC using 3:1 Hexane:EtOAc as the solvent system. Conversion to the carboxylic acid derivative was visualized via bromocresol green TLC staining (31). After 72 h, 0.5 N HCl was added to the reaction mixture until pH <6, followed by extraction with 20 ml of ethyl acetate (3×). The organic layers were combined and dried with Na2SO4 and evaporated to dryness under reduced pressure. The resulting product was a brownish crystalline solid and the yield was 200 mg. 1H NMR (DMSO, 400 MHz, 298 K, TMS) δ: 2.18–2.24 (m, 1H, 2'-H), 2.45–2.48 (m, 1H, 2'-H), 3.44–3.51 (m, 2H, 5'-H), 3.78–3.81 (m, 1-H, 4'-H), 4.30–4.33 (m, 1H, 3'-H), 4.88 (br s, 1H, 5'-OH), 5.73 (br s, 1H, 3'-OH), 6.37–6.40 (Japp, J = 7.1 Hz, 1H, 1'-H), 6.62 (d, J = 3.8 Hz, 1H, Ar), 7.62 (d, J = 8.8 Hz, 1H, Ar), 7.67 (d, J = 3.5 Hz, 1H, Ar), 7.72 (dd, J = 8.7 Hz, 1.6 Hz, 1H, Ar) 8.18 (d, J = 1.4 Hz, 1H, Ar), 12.5 (br s, 1H, –COOH). UV (MeOH) λmax (nm) 235 (ε = 36 074 cm⁻¹ M⁻¹). HiRes ESI-MS (+): calculated mass spectrum formula C14H15NO5 for M+H: 278.1028; experimental mass spectrum: 278.1027.

**Synthesis of 5-perfluorophenylindole 5.** To the mixture of 1-bromo-2,3,4,5,6-pentafluorobenzene (1.25 g, 5 mmol), Pd(PPh3)4 (0.17 g, 0.15 mmol) in DME (20 ml) was added 5-boronic acid indole (0.89 g, 5.5 mmol) followed by sodium bicarbonate (1.26 g, 15 mmol) in water (15 ml). The reaction mixture was refluxed for 3 h and cooled to room temperature to quench the reaction. DME was immediately evaporated under reduced pressure followed by the addition of ether (50 ml) to the two-layer mixture for extraction. The aqueous phase was extracted with ether (3×) and the organic phases were pooled, dried over MgSO4, filtered and evaporated to dryness. The crude product was purified by flash column chromatography (30–50% ethyl acetate in hexane). 5-perfluorophenylindole was a yellow solid (1.06 g, 74% yield). 1H NMR (CDCl3, 400 MHz, 298 K, TMS) δ: 6.62–6.63 (m, 1H, 3-H), 7.22 (dd, J = 4.0, 1.2 Hz, 1H, 7-H), 7.29 (Japp, J = 2.8 Hz, 1H, 2-H), 7.50 (d, J = 8.4 Hz, 1H, 6-H), 7.71 (s, 1H, 4-H), 8.30 (br s, 1H, 1-H). 19F-NMR (CDCl3, 376 MHz, 293 K, CCl3F) δ: −163.4—−163.2 (m, 2F), −157.6 (t, 1F, J = 21.1 Hz), −143.8 (dd, 2F, J = 31.7, 7.9 Hz). HiRes ESI-MS (+): calculated mass spectrum formula C14H14F5N for M+H: 284.0949; experimental mass spectrum: 284.0949.

**Synthesis of 1-(2-deoxy-β-D-erythropentafuranosyl)-5-perfluorophenyl-1H-indole 6.** This compound was prepared as described in method A using 5 (300 mg, 1.7 mmol, 1 eq) as the starting material. The crude product was purified by silica flash column chromatography using hexane:ethyl acetate (1:2) as eluent. The isolated product was a light brown solid and the yield was 445 mg. 1H NMR (DMSO, 400 MHz, 298 K, TMS) δ: 2.15–2.20 (m, 1H, 2'-H), 2.42–2.48 (m, 1H, 2'-H), 3.41–3.49 (m, 2H, 5'-H), 3.74–3.77 (m, 1-H, 4'-H), 4.29–4.33 (m, 1H, 3'-H), 4.83 (br s, 1H, 5'-OH), 5.23 (br s, 1H, 3'-OH), 6.35 (Japp, J = 6.47 Hz, 1H, 1'-H), 6.52–6.53 (d, J = 3.28 Hz, 1H, 3'-H), 7.15 (d, J = 8.80 Hz, 1H, Ar), 7.58–7.62 (m, 3H, 1H, Ar). 19F-NMR (CDCl3, 376 MHz, 293 K, CCl3F) δ: −163.2—−163.1 (m, 2F), −157.3 (t, 1F, J = 21.1 Hz), −143.8 (dd, 2F, J = 32.7, 8.7 Hz). UV (MeOH) λmax (nm) 248 (ε = 18 437 cm⁻¹ M⁻¹). HiRes ESI-MS (+): calculated mass spectrum formula C14H14F5NO5 for M+H: 400.0972; experimental mass spectrum: 400.0977.
Synthesis of 1-(2-deoxy-β-p-erythropentafuranosyl)-5-(2,3,5,6-tetrafluoro-4-methoxyphenyl)-1H-indole 7. This compound was prepared based on a slightly modified protocol (32) starting from 6 (300 mg, 0.8 mmol) dissolved in ether (5 ml). To this reaction mixture was added methanol (45 ml) and sodium methoxide until the pH of the solution is >12. The reaction mixture was stirred for 36 h at room temperature under argon and the product formation was monitored by 19F-NMR. This process afforded 70% conversion of the starting material to the product. The solution is then treated with 0.5 M DMF solution of tributylammonium hydrogen carbonate (31). Reverse-phase HPLC was also used to determine the progression of the deprotection process. Triphosphorylation was initiated by forming the phosphorodichloridated intermediate by the dropwise addition of POCl3 in the reaction mixture containing the 2′-mono-3′,5′-bis-O-(acrylamide)naphthalene (0.11 mmol) dissolved in 0.37 ml of dimethylphosphate pre-chilled at 0°C. This mixture was stirred for 3 h under argon and was then spontaneously treated with 0.5 M DMF solution of tributylammonium pyrophosphate (0.37 mmol) and tributylamine (0.37 mmol) and further stirred for 15 min at room temperature. Tributylaminumbicarbonate (1 M TEAB, pH 7.5) was added to quench the reaction and stirred at room temperature for 2 h to yield the nucleotide. The course of the reaction was monitored by TLC using the solvent system of 1-propanol:ammonium hydroxide:water (6:3:1) as the mobile phase A and 50% ACN in 0.1 M TEAB as mobile phase B. The desirable nucleotides were lyophilized to dryness and characterized by mass spectrometry and NMR. The concentration of the triphosphate was determined using the extinction coefficient at the UV λmax for the corresponding nucleoside (34).

1-(2-Deoxy-β-p-erythropentafuranosyl)-5-methylcarboxylate-1H-indole triphosphate 8. Using 6 as the starting material, this compound was prepared and purified as described in method B. 19F-NMR (D2O, 376 MHz, 293 K, CCl3F) δ: −164.31−164.16 (m, 2F); −158.15 (t, J = 21.6 Hz, 1F), −145.02 (dd, dd, J = 24.4 Hz, 8.6 Hz), −145.22 (dd, 2F, J = 23.7, 10.2 Hz). UV (MeOH) λmax (nm) 250 (ε = 30 342 cm−1 M−1). HiRes ESI-MS (+): calculated mass spectrum formula C20H17F4NO4 for M+Na: 434.0992; experimental mass spectrum: 434.0982 for M-H: 530.0024; experimental mass spectrum: 530.0036.

1-(2-Deoxy-β-p-erythropentafuranosyl)-5-carboxylate-1H-indole triphosphate 4. Compound 3 was converted to 4 by reacting with NaOH in water (2:1) overnight at room temperature in the presence of 15-crown-5 ether. The course of the reaction was monitored by TLC using 1-propanol:ammonium hydroxide:water (6:3:1) as the solvent system for elution and the conversion of the methyl carboxylate to the carboxylate moiety was visualized via bromocresol green staining which detects carboxylic acid (31). Reverse-phase HPLC was also used to determine the progression of the deprotection process. 5-MeCITP was eluted at 46% B (13 min retention time) using a C-18 analytical column from Vydac; buffer A: 0.1 M TEAB; buffer B: 50% ACN in 0.1 M TEAB, linear gradient of 20–50% within 15 min at a flow rate of 1 ml min−1 and monitored at a wavelength of 235 nm. Using the same conditions, 5-CITP was eluted at 27% B (3.5 min retention time). 31P-NMR (D2O, 162 MHz, H2PO4) δ: −4.9 (d, γ-P), −9.9 (d, α-P), −20.8 (d, β-P). HiRes ESI-MS (−): calculated mass spectrum formula C14H16NO14P3Na for M-H: 537.96868; experimental mass spectrum: 537.96875.

Method B: synthesis of non-natural deoxynucleoside triphosphates

The triphosphates were prepared as previously described (24,25,28,33) starting from the corresponding nucleosides. Triphosphorylation was initiated by forming the 5′-mono-phosphorodichloridated intermediate by the dropwise addition of POCl3 in the reaction mixture containing the 2′-deoxyribonucleoside (0.07 mmol) and 1,8-bis(dimethylamino)naphthalene (0.11 mmol) dissolved in 0.37 ml of trimethylphosphate pre-chilled at 0°C. This mixture was stirred for 3 h under argon and was then spontaneously treated with 0.5 M DMF solution of tributylammonium pyrophosphate (0.37 mmol) and tributylamine (0.37 mmol) and further stirred for 15 min at room temperature. Tributylaminumbicarbonate (1 M TEAB, pH 7.5) was added to quench the reaction and stirred at room temperature for 2 h to yield the nucleotide. The course of the reaction was monitored by TLC using the solvent system of 1-propanol:ammonium hydroxide:water (6:3:1). The crude product was concentrated by rotavaporation under reduced pressure prior to purification by preparative reverse phase HPLC using 0.1 M TEAB as mobile phase A and 50% ACN in 0.1 M TEAB as mobile phase B. The desirable nucleotides were lyophilized to dryness and characterized by mass spectrometry and NMR. The concentration of the triphosphate was determined using the extinction coefficient at the UV λmax for the corresponding nucleoside (34).

1-(2-Deoxy-β-p-erythropentafuranosyl)-5-perfluorophenyl-1H-indole triphosphate 8. Using 6 as the starting material, this compound was prepared and purified as described in method B. 19F-NMR (D2O, 376 MHz, 293 K, CCl3F) δ: −164.31−164.16 (m, 2F); −158.15 (t, J = 21.6 Hz, 1F), −145.02 (dd, dd, J = 24.4 Hz, 8.6 Hz, 2F), −145.22 (dd, 2F, J = 23.7, 10.2 Hz). 31P-NMR (D2O, 162 MHz, H2PO4) δ: −5.8 (d, γ-P), −10.2 (d, α-P), −21.4 (d, β-P). HiRes ESI-MS (−): calculated mass spectrum formula C14H16NO14P3Na for M-H: 637.9811; experimental mass spectrum: 637.9809.

Enzymatic assays

Materials. Magnesium acetate and Trizma base were from Sigma. [γ-32P]-ATP was purchased from Perkin Elmer Life and Analytical Sciences (Boston, MA, USA). Boric acid, urea, acrylamide and bis-acrylamide were from National Diagnostics (Rochester, NY, USA). Oligonucleotides, including those containing a tetrahydrofuran moiety mimicking an abasic site, were synthesized by Operon Technologies (Almeda, CA, USA). All other materials were obtained from commercial sources and were of the highest quality available. The exonuclease-deficient mutant of gp43 (Asp-219 to Ala mutation) was prepared based on a slightly modified protocol (32) starting from 6 (300 mg, 0.8 mmol) dissolved in ether (5 ml). To this reaction mixture was added methanol (45 ml) and sodium methoxide until the pH of the solution is >12. The reaction mixture was stirred for 36 h at room temperature under argon and the product formation was monitored by 19F-NMR. This process afforded 70% conversion of the starting material to the product. The solution is then treated with 0.5 M DMF solution of tributylammonium pyrophosphate (0.37 mmol) and tributylamine (0.37 mmol) and further stirred for 15 min at room temperature. Tributylaminumbicarbonate (1 M TEAB, pH 7.5) was added to quench the reaction and stirred at room temperature for 2 h to yield the nucleotide. The course of the reaction was monitored by TLC using the solvent system of 1-propanol:ammonium hydroxide:water (6:3:1). The crude product was concentrated by rotavaporation under reduced pressure prior to purification by preparative reverse phase HPLC using 0.1 M TEAB as mobile phase A and 50% ACN in 0.1 M TEAB as mobile phase B. The desirable nucleotides were lyophilized to dryness and characterized by mass spectrometry and NMR. The concentration of the triphosphate was determined using the extinction coefficient at the UV λmax for the corresponding nucleoside (34).
purified and quantified as previously described (35, 36).

Enzyme assays. The assay buffer used in all kinetic studies with the exonuclease-deficient bacteriophage T4 DNA polymerase consisted of 25 mM Tris-OAc (pH 7.5), 150 mM KOAc and 10 mM 2-mercaptoethanol. All enzymatic assays were performed at 25°C and polymerization reactions were monitored by analysis of the product in 20% sequencing gels as previously described (37). Gel images were obtained with a PhosphorImager instrument using the Optiquant software supplied by the manufacturer. Product formation was quantified by calculating the ratio of extended (14-mer) and non-extended (13-mer) 32P-radiolabeled primer which was corrected for substrate in the absence of polymerase (zero point). The total product was derived by multiplying the corrected ratios with the total concentration of primer/template used in each assay. All concentrations are listed as final solution concentrations.

Nucleotide incorporation assays. A rapid quench instrument (KinTek Corporation, Clarence, PA) was employed to monitor the time course in nucleotide incorporation by gp43 exo− opposite a single- and double abasic sites. $k_{pol}$ and $K_{d, app}$ values were determined using single-turnover reaction conditions in which excess gp43 exo− (500 nM) was preincubated with DNA (250 nM) in an assay buffer and mixed with variable concentrations of the nucleotide analogue (5–500 μM) and 10 mM Mg2+. The reactions were quenched with 350 mM EDTA at variable times (0.001–10 s) and analyzed as previously described. The time course in product formation was fitted using the following equation:

$$y = Ae^{-kt_{obs}} + C$$

(1)

where $y$ is the amount of product, $A$ is the amplitude, $k_{obs}$ is the rate constant, $t$ is time and $C$ is a defined constant. $K_{d, app}$ and $k_{pol}$ were determined by fits of the data points to the Michaelis–Menten equation:

$$k_{obs} = \frac{k_{pol}[dXTP]}{K_{d, app} + [dXTP]}$$

(2)

where $k_{obs}$ is the apparent first-order rate constant, $k_{pol}$ is the maximal polymerization rate constant, $K_{d, app}$ is the apparent dissociation constant for dXTP and [dXTP] is the concentration of nucleotide substrate.

Extension enzymatic assays. Rates of elongating beyond the incorporated nucleotide were measured under single-turnover conditions. In these experiments, gp43 exo− (500 nM) was incubated with 250 nM DNA in assay buffer containing 10 mM Mg2+ and mixed with 150 μM nucleotide. After four half-lives, an aliquot of the reaction was quenched with 350 mM EDTA to validate nucleotide insertion opposite the lesion. At this time, dGTP (500 μM) was then added. Aliquots were quenched with 350 mM EDTA at variable time points and analyzed as described above.

RESULTS

Synthesis of isosteric non-natural nucleotides

Two non-natural nucleotides designated as 5-CITP and 5-MeCITP (Figure 2A) were synthesized to further interrogate the role of hydrophobicity, π-electron interactions and steric constraints during translesion DNA synthesis. In this case, 5-MeCITP and 5-NITP are similar with respect to hydrophobicity and π-electron density yet are significantly different with respect to overall shape and size. In contrast, 5-CITP is nearly identical to 5-NITP with respect to shape, size and π-electron surface area yet differ with respect to hydrophobicity and solvation energies. These differences provide an excellent way to probe their roles during the misreplication of an abasic site. Both non-natural nucleotides were synthesized as outlined in Scheme 1. 5-MeCITP was prepared using the protocol established by Girgis et al. (34). Briefly, formation of the N-glycosidic bond between the 5-methylcarboxylate indole and Hoffer’s chlorosugar was achieved by employing sodium hydride as the base to form the sodium salt of the indole nucleobase in situ which then reacts with the electrophilic center of the anomic carbon (C-1′) of the deoxyribose sugar. Deprotection of the 3′- and 5′-hydroxyl moieties of the sugar was accomplished by employing the Zemplén conditions to yield mainly the β-anomeric isomer as confirmed by nuclear-Overhauser effect (NOE) difference experiments (NOEs of H-2, H-7, H-2′α and H-2′β upon irradiation of H-1′) as well as the presence of an apparent triplet ($t_{app}$) peak for the H-1′ resonance with a coupling constant of ~7.0 Hz which are characteristics for the β-anomer of N-nucleosides (38, 39). The one-pot synthesis procedure of Smith et al. (33) was used to convert the nucleoside to the corresponding triphosphate by first reacting the substituted indolyl deoxyribonucleoside with POCl3 to form the 5′-monophosphorodichloridated nucleoside intermediate which was then treated with pyrophosphosphate to form the nucleoside 5′-triphosphate. 5-CITP was prepared from 5-MeCITP in which the methyl ester protecting group was cleaved using sodium hydroxide in the presence of 15-crown-5 ether to chelate the sodium ions that act as a Lewis acid and jeopardize the integrity of the triphosphate group. In fact, attempts to cleave the methyl ester protecting group in the absence of 15-crown-5 ether produced low amounts of the desired product (data not shown). The conversion of 5-MeCITP to 5-CITP was monitored via TLC stained with bromocresol green dye (31) and analytical HPLC as further described in the Experimental Section.

Analogos of 5-PhITP were synthesized in which the π-electron density around the phenyl substituent group of 5-PhITP was reduced by substituting highly electron-negative fluorines for all of the hydrogen atoms. As illustrated in Figure 2B, this substitution reduces the
attached to the deoxyribose sugar using the protocol established by Girgis et al. (34) as previously described above. The one-pot synthesis procedure of Smith et al. (33) was used to obtain the desired product, 5-F5-PhITP, which was verified by nuclear magnetic resonance spectroscopy and high-resolution mass spectrometry analyses (Supplementary Data S2A and S2B).

5-F4OMe-PhITP, a slightly larger structural analog of 5-F5-PhITP (Figure 2B), was synthesized to further interrogate the influence of nucleobase size on the kinetic parameters of nucleotide incorporation. 5-F4OMe-PhIdR, 7, was obtained using a slightly modified procedure (32) starting from 5-F5-PhIdR, 6. In this procedure, 6 was dissolved in 10% ether in anhydrous methanol followed by the addition of sodium methoxide until pH >12 and vigorously stirred for 36 h. 19F-NMR and high-resolution mass spectrometry indicated that the fluorine atom at the para position of the perfluorophenyl ring was substituted by the methoxy moiety under these conditions. Phosphorylation of 7 was performed as previously described (33) and characterized by NMR and mass spectrometry after purification (Supplementary Data S2C and S2D).

**Nucleobase hydrophobicity and π-electron density influences translesion DNA synthesis**

The kinetic parameters for the utilization of 5-CITP opposite an abasic site were measured using a defined DNA substrate illustrated in Figure 2C. Time courses in product formation were generated using single-turnover conditions in which 500 nM of the exonuclease-deficient bacteriophage T4 DNA (gp43 exo-) was incubated with 250 nM DNA containing an abasic site (designated as 13/20SP-mer) and mixed with variable concentrations of non-natural nucleotide (5–1000 μM) and 10 mM Mg2+. Representative time courses provided in Figure 2D were fit to an equation describing a single exponential process (Equation 1) to define $k_{app}$, the rate constant in nucleotide incorporation.

$$k_{obs} = k_{pol}/K_d + 1$$

The plot of $k_{obs}$ versus 5-CITP concentration is hyperbolic (Figure 2E) and a fit of the data to the Michaelis–Menten equation (Equation 2) yields a $k_{pol}$ value of 67 ± 5 s$^{-1}$, a $K_d$ of 172 ± 31 μM and a $k_{pol}$/$K_d$ of 3.9 × 10$^5$ M$^{-1}$s$^{-1}$ for 5-CITP. The fast $k_{pol}$ of ~70 s$^{-1}$ is consistent with a role for π-electron density in modulating rate of the conformational change that precedes phosphoryl transfer. However, the $K_d$ of 170 μM for 5-CITP is ~10-fold higher than the hydrophobic analog, 5-NITP. This difference suggests that the binding of a non-natural nucleotide is governed largely by its hydrophobic nature (Table 2).

The kinetic parameters for 5-MeCITP were measured to further validate this mechanism. As reported in Table 1, the $K_d$ of 13 μM for 5-MeCITP is ~13-fold lower than that for 5-CITP and the increase in binding affinity is directly linked with the increase in hydrophobicity caused by the addition of the methyl group (Table 2). Moreover, 5-MeCITP displays a fast $k_{pol}$ value of 79 s$^{-1}$ that is nearly identical to that of 5-CITP. The fast $k_{pol}$ value with both analogs is consistent with the presence of π-electron density generated by the carbonyl groups.

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**Figure 2.** Structures of (A) 5-NTP, 5-CITP and 5-MeCITP; (B) 5-PhITP, 5-F5-PhITP and 5-F4OMe-PhITP. For clarity, only the nucleobases of the isosteric 5-substituted indolyl nucleotide analogues are shown; (C) Nucleotide sequence of the DNA substrate used in kinetic studies. (D) Dependency of the apparent burst rate constant on the concentration of 5-CITP as measured under single-turnover conditions. Assays were performed using 500 nM gp43 exo-; 250 nM 13/20SP-mer, 10 mM Mg(OAc)$_2$ and 5-CITP in variable concentrations of 10 (open circle), 25 (closed circle), 50 (open square), 100 (closed square), 200 (open diamond) and 500 μM (closed diamond). (E) Observed rate constants for incorporation (closed circle) were plotted against 5-CITP concentration. A fit of the data to the Michaelis–Menten equation yields a $k_{pol}$ of 67 ± 5 s$^{-1}$ and a $K_d$ of 172 ± 31 μM.

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overall π-electron density without significantly perturbing the overall size of the phenyl ring (Supplementary Data S1). Scheme 2 outlines the synthesis of the fluorophenyl-substituted indolyl nucleotides. The 5-perfluorophenylindolindole nucleobase 5 was first prepared using the Suzuki–Miyaura coupling method (40) and...
Collectively, these results are consistent with a model in which nucleotide binding is directly linked with its hydrophobic nature while the rate constant for incorporation is influenced by the amount of π-electron density present in the incoming nucleotide.

The role of π-electron density during translesion DNA synthesis was further investigated by measuring the kinetic parameters for the π-electron-deficient analogs, 5-F5-PhITP and 5-F4OMe-PhITP, opposite an abasic site (Table 2). Representative time courses for the incorporation of 5-F5-PhITP and the plot of $k_{\text{obs}}$ versus 5-F5-PhITP concentration are provided as Supplementary Data S3A and S3B. A fit of the data to the Michaelis–Menten equation yields a $k_{\text{pol}}$ of 15 ± 1 s$^{-1}$, a $K_{\text{d, app}}$ value of 2.3 ± 0.7 mM and a $k_{\text{pol}}/K_{\text{d, app}}$ of $6.5 \times 10^6$ M$^{-1}$ s$^{-1}$ for this π-electron-deficient analog (Table 1). It is surprising that the overall catalytic efficiency of 5-F5-PhITP is ∼2-fold higher than 5-PhITP since introducing fluorine groups should have reduced the amount of π-electron density within the aromatic phenyl ring and thus ‘lowered’ the efficiency of incorporation. However, close inspection of the kinetic parameters indicates that reducing π-electron stacking potential indeed lowers the polymerization rate constant by 3-fold (15 s$^{-1}$ versus 53 s$^{-1}$ using 5-F5-PhITP and 5-PhITP, respectively). As such, the higher catalytic efficiency for the fluorinated analog results from a 6-fold reduction in the $K_{\text{d, app}}$ value (compare a $K_{\text{d, app}}$ of 2.3 μM for 5-F5-PhITP with 14 μM for 5-PhITP). This effect coincides with the slightly larger size of 5-F5-PhITP compared with 5-PhITP and could reflect better shape complementarity with the DNA lesion. An alternative explanation is that the fluorinated nucleobase is more hydrophobic as it has a lower solvation energy (41).

These mechanisms were evaluated by monitoring the incorporation of 5-F4OMe-PhITP opposite the abasic site (Supplementary Data S4A and S4B). Kinetic analyses yield a $k_{\text{pol}}$ of 18 ± 1 s$^{-1}$, a $K_{\text{d, app}}$ of 4 ± 1 μM and a $k_{\text{pol}}/K_{\text{d, app}}$ of $4.5 \times 10^6$ M$^{-1}$ s$^{-1}$ for this π-electron-deficient analog (Table 1). The $k_{\text{pol}}$ value for 5-F4OMe-PhITP is 3-fold lower than 5-PhITP, again indicating that reductions in π-electron stacking potential slow the rate constant for this kinetic step. In addition, the $K_{\text{d, app}}$ of 4 mM for 5-F4OMe-PhITP is 4-fold lower than 5-PhITP, which coincides with the slightly larger size and lower solvation energy associated with the fluorinated nucleobase. While this suggests that shape complementarity is an important determinant in substrate binding, it should be noted that the $K_{\text{d, app}}$ of 4 μM for 5-F4OMe-PhITP is lower than that of 14 μM measured with 5-PhITP. This ∼4-fold increase in binding affinity coincides with a slightly larger size and lower solvation energy associated with the fluorinated nucleobase. While this suggests that shape complementarity is important in determining the binding site. However, it should be noted that the $K_{\text{d, app}}$ of 4 μM for 5-F4OMe-PhITP is ∼2-fold higher than that for 5-5-PhITP ($K_{\text{d, app}}$ of 2.3 μM). This reduction in binding affinity suggests that optimizing the size of the nucleobase to fit within the void of the abasic site has a minimal effect on binding affinity.

**Incorporation opposite other non-instructional DNA lesions**

The results presented above are consistent with a mechanism in which nucleotide binding is controlled by the
hydrophobic nature of the incoming nucleotide while the polymerization rate constant is influenced by π-electron density. To test whether this mechanism applies to other non-instructional DNA lesions, we measured the kinetic parameters for these non-natural nucleotides using DNA containing two adjacent abasic sites. As summarized in Table 1, the $K_{d, \text{app}}$ and $k_{\text{pol}}$ values for most non-natural nucleotides are similar regardless of whether the DNA contains a single abasic site or two adjacent DNA lesions. The best example for this behavior is 5-CITP as it displays identical $K_{d, \text{app}}$ values of 170 $\mu$M and identical $k_{\text{pol}}$ values of 70 s$^{-1}/C_0$ using either DNA substrate. However, there are some important exceptions that highlight nuances in the replication of non-instructional DNA lesions. For instance, the $K_{d, \text{app}}$ values for both fluorinated non-natural nucleotides are ~5-fold higher at the double abasic site compared with a single abasic site. This difference could reflect weaker binding interactions caused by the larger void of adjacent abasic sites compared with a single abasic site. In addition, differential effects are observed on the $k_{\text{pol}}$ value for both fluorinated nucleotides. The $k_{\text{pol}}$ for 5-F5-PhITP is slightly slower measured opposite a double abasic site compared with a single abasic site. Surprisingly, an opposite effect is observed with 5-F4OMe-PhITP as the $k_{\text{pol}}$ value of 30 s$^{-1}$ measured opposite the double abasic site is 2-fold faster than that of 15 s$^{-1}$ measured with a single-DNA lesion. In this case, the overall catalytic efficiency ($k_{\text{pol}}/K_{d, \text{app}}$) for 5-F4OMe-PhITP is nearly identical between the two types of DNA lesion. This occurs as larger $K_{d, \text{app}}$ values are offset by faster $k_{\text{pol}}$ values. Similar effects are observed with the natural nucleotide, dATP, as its binding affinity is weakened 5-fold while the rate constant for incorporation is increased 4-fold as a consequence of expanding the size of the ‘void’ in DNA.

More significant differences are observed during blunt-end extension, another form of non-instructional DNA lesion (42). As indicated in Table 1, only 5-MeCITP is utilized during blunt-end DNA synthesis, albeit with a low catalytic efficiency ($k_{\text{pol}}/K_{d, \text{app}}$) for 5-F4OMe-PhITP is nearly identical between the two types of DNA lesion.

Scheme 2. Synthetic scheme for the preparation of fluorinated analogs. Reagents and conditions were as follows: (a) Pd(PPh$_3$)$_4$, NaHCO$_3$ in H$_2$O, DME, reflux, 2 h; (b) NaH, anhydrous ACN, RT, 16 h; (c) NaOMe, MeOH, pH $\geq$12, RT, 16 h; (d) (i) POCl$_3$, Proton Sponge$^\circ$, trimethylphosphate, 0°C; (ii) Tributylammonium pyrophosphate, DMF, tributylamine, RT, 15 min and (iii) 1 M TEAB, RT, 2 h and (e) NaOMe, 1:10 (ether:MeOH), pH $\geq$12, RT, 36 h.
effectively during blunt-end extension compared with the fluorinated analogs, 5-F5-PhITP and 5-F4OMe-PhITP. The reduced capacity of the fluorinated analogs to act as efficient substrates is likely due to deficiencies in π-electron density. Although 5-CITP possesses π-electron density, it is not utilized during blunt-end DNA synthesis since its extremely hydrophilic nature adversely affects its binding affinity. Collectively, these data indicate that efficient nucleotide incorporation opposite any non-instructional DNA lesion depends upon a combination of low solvation energy coupled with significant π-electron density.

**Incorporation opposite templating nucleobases**

The incorporation of these analogs opposite a templating thymine were also measured (Table 1). Although all of the non-natural nucleotides reported here are inserted opposite T, their overall catalytic efficiencies are ~100-fold lower compared with the non-instructional abasic site. Their reduced efficiencies are caused by lower $k_{pol}$ values rather than through perturbations in their $K_{d, app}$ values and this result is similar to that obtained using the other non-natural nucleotides displayed in Figure 1A. Despite this difference, it is clear that insertion of a non-natural nucleotide opposite a templating nucleobase is also influenced by the combination of nucleobase hydrophobicity and π-electron density. For example, fluorinated analogs of 5-PhITP have higher catalytic efficiencies compared with 5-PhITP (Table 1). In both cases, the increase in catalytic efficiency for the fluorinated analogs is caused by significant decreases in their $K_{d, app}$ values that compensate for slower $k_{pol}$ values.

**Extension beyond templating DNA and non-instructional lesions**

We previously demonstrated that the bacteriophage T4 DNA polymerase extends beyond dATP or dGTP when paired opposite an abasic site (10,26). However, this high-fidelity DNA polymerase cannot extend beyond any of the 5-substituted indolyl nucleotides illustrated in Figure 1A. The lack of extension could be caused by alterations in the conformation of the primer template (43) or by a lack of hydrogen-bonding functional groups that serve as recognition elements for elongation (44). To differentiate between these models, we used the experimental protocol outlined in Figure 3A to quantify extension beyond the non-natural nucleotides developed in this study. These experiments employed single-turnover conditions in which DNA polymerase (500 nM) and DNA substrate (250 nM) were mixed with 150 μM non-natural nucleotide and 10 mM Mg$^{2+}$ to allow for incorporation opposite the abasic site. After four half-lives (time required to obtain 95% mispair formation), 500 μM dGTP was added to initiate elongation beyond the abasic site. Aliquots were quenched with EDTA at variable times and analyzed via denaturing gel electrophoresis to visualize product formation. Representative data provided in Figure 3B shows that the T4 DNA polymerase does not extend beyond hydrophobic nucleotides such as 5-NITP, 5-F5-PhITP, 5-F4OMe-PhITP and 5-PhITP since only 14-mer product is observed (Figure 4B, lanes 2, 5, 6 and 7, respectively). In contrast, the T4 DNA polymerase extends beyond 5-CITP (lane 3) and 5-MeCITP (lane 4) almost as efficiently as dATP (lane 8). In fact, quantifying the amount of primer extension reveals that the efficiency for extending beyond 5-CITP and 5-MeCITP is only 20% lower than that for dATP (Figure 3C). These data support a mechanism in which the presence of functional groups that can participate in hydrogen-bonding interactions is an important determinant for elongation.

**DISCUSSION**

Although hydrogen bonding interactions are widely associated with efficient and faithful DNA replication, other biophysical features such as π–π stacking, desolvation and geometrical constraints also contribute to this process (45). Unfortunately, it is difficult to accurately define the individual contributions of one feature
versus another since they are interrelated. This is best exemplified by examining the consequences of exposing nucleic acid to DNA damaging agents. For example, alkylation of the O6 position of guanine creates a miscoding DNA lesion, O6-methylguanine, that has altered hydrogen bonding potential compared with the unmodified nucleobase. However, alkylation also influences other biophysical features including nucleobase hydrophobicity, tautomeric form and shape/size of the nucleobase. These global changes make it difficult to unambiguously assign the energetic contributions of hydrogen bonding, \( \pi-\pi \) stacking interactions, desolvation energies and geometric constraints during correct and pro-mutagenic DNA synthesis.

To circumvent these problems, we studied the replication of an abasic site, a non-instructional DNA lesion that is devoid of a templating base and thus alleviates complications arising from hydrogen bonding interactions. In addition, we utilized a series of isosteric 5-substituted indolyl deoxynucleotides to probe the mechanism of nucleotide incorporation opposite this lesion. Since these nucleotide analogs contain modified functional groups attached to a common indole scaffold, the degree of \( \pi \)-electron density, desolvation energies, dipole moments and shape/size can be varied (Table 2) and quantified to evaluate their influence on the two most important kinetic steps associated with polymerization. These include nucleotide binding and the conformation change step preceding phosphoryl transfer. In this study, the use of single-turnover reaction conditions allows the first round of enzyme turnover to be accurately measured. This feature provides constraints on the derived kinetic parameters such that \( K_{d, \text{app}} \) values represent close approximations to true binding affinities for the non-natural nucleotides. In addition, we have demonstrated through elemental effects on nucleotide incorporation (10,46), pulse chase (47,48) and fluorescence analyses (49) that \( k_{\text{pol}} \) values reflect the conformational change step that precedes phosphoryl transfer. As a result, measurable differences in these kinetic parameters as a function of

| Nucleotide | Nucleobase structure | Surface area (Å\(^2\)) | Volume (Å\(^3\)) | Dipole moment (debye) | Solvation energy (kJ/mol) |
|-----------|----------------------|-------------------------|------------------|----------------------|--------------------------|
| dATP      |                      | 143.0                   | 124.5            | 2.68                 | -50.48                   |
| 5-NITP    |                      | 174.0                   | 154.5            | 9.80                 | -27.67                   |
| 5-CITP    |                      | 176.0                   | 158.5            | 16.1                 | -309.12                  |
| 5-MeCITP  |                      | 201.0                   | 181.3            | 5.20                 | -26.14                   |
| 5-PhITP   |                      | 223.2                   | 216.2            | 1.08                 | -10.15                   |
| 5-F5-PhITP|                      | 256.2                   | 240.5            | 4.09                 | -11.67                   |
| 5-F4OMe-PhITP |               | 278.3                   | 262.8            | 3.30                 | -13.26                   |

*Surface areas and volumes (used as an indicator of the relative size of the nucleobase), dipole moments and solvation energies (in the aqueous phase) for each nucleobase were calculated using Spartan '08 software. The semi-empirical method of calculation with AM1 was used for geometry optimization and the Density Functional Theory level of calculation with B3LYP 6-31G** basis sets were used for energy minimization.
non-natural nucleotide can be used to quantify the importance of various biophysical features.

The utility of this approach is apparent by comparing the kinetic parameters for utilizing 5-NITP and 5-CITP during translesion DNA synthesis. These particular nucleotides are essentially identical with respect to shape, size, \( \pi \)-electron density and dipole moment. However, their solvation energies are significantly different so that any variation in their catalytic efficiencies can be attributed to the energetic requirements for nucleobase desolvation. In fact, the differences in \( k_{\text{pol}}/K_{\text{d, app}} \) values of \( 7 \times 10^6 \text{M}^{-1}\text{s}^{-1} \) and \( 3.9 \times 10^5 \text{M}^{-1}\text{s}^{-1} \) for 5-NITP and 5-CITP, respectively, indicate that desolvation contributes \( \sim 1.71 \text{kcal/mol} \) (The overall change in Gibb’s free energies was calculated using the equation, \( \Delta \Delta G^\circ = RT \ln K \), where \( R = 1.9872 \text{cal mol}^{-1} \text{K}^{-1} \), \( T = 298 \text{K} \) and \( K \) is the ratio of respective kinetic parameters for the incorporation of these non-natural nucleotides opposite the lesion.) to the overall efficiency for nucleotide incorporation opposite the abasic site. The influence of desolvation

**Figure 3.** Measuring extension beyond an abasic site. (A) Experiments were performed under single-turnover conditions in which 500 nM bacteriophage T4 DNA polymerase was incubated with 250 nM 13/20SP-mer in assay buffer containing 10 mM Mg\(^2+\) and mixed with 150 \( \mu \text{M} \) dXTP for 5 min. To initiate extension beyond the lesion, 500 \( \mu \text{M} \) dGTP, the correct nucleotide for the next three insertion positions, was added and the reactions were quenched with 350 mM EDTA at 180 s. (B) Denaturing gel electrophoresis data for the incorporation and extension of nucleotides opposite an abasic site catalyzed by the bacteriophage T4 DNA polymerase. Only nucleotides containing hydrogen-bonding functional groups (dATP, 5-CITP and 5-MeCITP) showed extension beyond the abasic site. (C) Quantification of gel electrophoresis data provided in panel B. Blue bars represent percentage of 14-mer product (incorporation of nucleotide opposite abasic site) while red bars represent percentage of 14-mer product that is elongated after addition of dGTP.
toward nucleotide binding ($K_d, \text{ app value}$) and the conformational change step ($k_{pol}$ value) can be further broken down using an identical approach. The energetic contribution of desolvation is 1.34 kcal/mol for nucleotide binding, while that for the conformational change step is 0.37 kcal/mol. As such, this analysis indicates that nucleobase desolvation plays a more significant role during the binding of non-natural nucleotides compared with the conformational change step preceding phosphoryl transfer.

Although desolvation may occur during the conformational change step, we envisioned that this kinetic step is more sensitive to the presence of $\pi$-electron density. This conclusion is based upon differences in the measured $k_{pol}$ values for 5-PhITP and the fluorinated analogs, 5-F5-PhITP and 5-F4OMe-PhITP. Since these analogs possess similar shapes, sizes and desolvation energies, the quantifiable disparities in their $k_{pol}$ values should primarily be attributed to energetic differences associated with $\pi$-electron density. Using the $k_{pol}$ values of 53 s$^{-1}$ versus 15 s$^{-1}$ for 5-PhITP and 5-F5-PhITP, respectively, a $\Delta \Delta G^0$ value of 0.75 kcal/mol is calculated for the conformational change step. This energetic difference may not be as large as expected for $\pi-\pi$ stacking interactions that must occur inside the interior of duplex DNA. However, reducing the $\pi$-electron density within the phenyl ring of 5-PhITP via substitution of hydrogen with electronegative fluorine (5-F5-PhITP) also introduces a partial positive charge at the center of the perfluorinated analog. As a result, favorable $\pi-\pi$ stacking interactions could be replaced by electrostatic quadrupolar interactions that are similar to $\pi$-cation interactions. Indeed, studies have shown that fluorinated aryl compounds have the ability to base stack with electron-rich aromatic rings through quadrupolar electrostatic interactions (50–52). This argument is consistent with the ~2-fold higher $k_{pol}$ value of 5-NITP (126 s$^{-1}$) compared with 5-CITP (67 s$^{-1}$). While both molecules possess similar $\pi$-electron densities, the positive charge on the nitrogen of 5-NITP may slightly enhance its base-stacking abilities (53). However, we note that the dipole moments and/or van der Waals contacts could also contribute to the mechanism of nucleotide incorporation opposite an abasic site. Further experimentation is being performed to address these features.

These data were used to propose the model outlined in Figure 4 that highlights the roles of nucleobase desolvation and $\pi-\pi$ stacking interactions during the replication of an abasic site. In this model, nucleobase desolvation primarily occurs during the initial binding

Figure 4. Simplified model illustrating the kinetic steps involved in desolvation and $\pi-\pi$ stacking interactions during the replication of an abasic site. See text for complete details. The model was generated using the Spartan '08, ChemBioOffice 2010 and Microsoft Office 2010 Powerpoint software packages.
undergoes a conformational change that aligns the
*P*-phosphate of the incoming nucleotide with the 3'-OH
of the primer for phosphoryl transfer. This step is
controlled predominantly by the *P*-electron density
present on the incoming nucleotide and allows for
proper *P*-*P* stacking interactions within an intrahelical
conformation to commence phosphoryl transfer. With
the bacteriophage T4 DNA polymerase, this kinetic step
is also influenced by base-stacking interactions with the
downstream templating nucleobase since polymerization
using blunt-end DNA or DNA containing a double
abasic site occurs with reduced *K*₂₀ values without a
negative effect on *K*~d, app~.

This model can also be applied to describe the roles
of desolvation, *P*-*P* stacking, steric fit and hydrogen-bonding
interactions during normal DNA synthesis. In the
presence of a templating nucleobase, it is clear that
desolvation must also occur during the initial binding
step as water molecules must be effectively stripped
away from functional groups that eventually participate
in hydrogen-bonding interactions. In this case, energetic
penalties caused by the removal of water molecules are
adequately offset by formation of favorable enthalpic
contributions by hydrogen bonding interactions between
correctly paired nucleobases. Subsequent formation of
proper *P*-*P* stacking interactions occur during the
conformational change step then allows for phosphoryl
transfer and primer elongation to occur. As such, there
are two major energetic differences between correct
versus translesion DNA synthesis. The first is with
respect to desolvation and enthalpic compensation
during the binding of a nucleotide. During correct DNA
synthesis, energetic penalties associated with nucleobase
desolvation are offset by the formation of correct and
complementary hydrogen bonding interactions. Enthalpic compensation via hydrogen-bonding inter-
actions does not occur at non-instructional DNA lesions
such as an abasic site and this deficiency adversely affects
the mechanism of nucleotide binding. The other major
difference is the involvement of nucleobase desolvation
during the conformational change step. During correct
DNA synthesis, there is no need for nucleobase desolvation
during this kinetic step as functional groups on the templating and incoming nucleobase are already
locked into proper hydrogen bonding interactions.

During the replication of a DNA lesion, however,
desolvation likely occurs during the conformational
change step and this energetic demand also reduces the
overall efficiency of nucleotide incorporation. A provoc-
tive implication is that the primary determinant for
efficient and faithful DNA synthesis is the ability of the
DNA polymerase to effectively perform nucleobase
desolvation.

Primer extension depends upon hydrogen-bonding
interactions and shape complementarity

While *P*-electron density and hydrophobicity are impor-
tant for efficient nucleotide incorporation opposite an
abasic site, these features are not required for primer
elongation. In fact, nucleotide extension depends heavily
upon the presence of functional groups that provide
hydrogen-bonding interactions. This conclusion is based
upon the observation that the bacteriophage T4 polymer-
ase does not extend beyond 5-PhThT or 5-NITP but can
extend beyond 5-CITP and 5-McCITP when paired
opposite an abasic site. Elongation beyond 5-CITP and
5-McCITP likely occurs since both analogs contain a
carbonyl group that can hydrogen bond with active site
amino acids necessary for elongation. This observation
is similar to results indicating that most high-fidelity DNA
polymerases are incapable of extending non-natural nu-
cleotides unless the formed base pair contains at least
one functional group that participate as a hydrogen
bond acceptor or donor (44).

The presence of hydrogen-bonding groups does not
always guarantee that a non-natural nucleotide will be
elongated. This is evident as the bacteriophage T4 DNA
polymerase cannot extend beyond 5-CITP or 5-McCITP
when they are paired opposite thymine (data not shown),
the predicted pairing partner for these non-natural nucleo-
tides. The lack of extension argues that the conformation
of the primer-template also plays an important role in
defining if a base pair will be efficiently extended or not.
This interpretation is consistent with work published by
Romesberg indicating that extension beyond a base-pair
containing a non-natural nucleotide depends upon the
proper geometry of the terminal base pair (43,54). Our
data extend these findings and indicate that efficient elong-
ation depends upon the presence of hydrogen bonding
groups that must be in a precise geometrical orientation.
If either condition is not met, primer elongation cannot
occur. This model may also explain why most mispairs
formed between natural nucleotide and DNA lesions are
poorly elongated by high-fidelity DNA polymerases
(55,56).

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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