Conformational Insights into the Lesion and Sequence Effects for Arylamine-Induced Translesion DNA Synthesis: $^{19}$F NMR, Surface Plasmon Resonance, and Primer Kinetic Studies

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**ABSTRACT:** Adduct-induced DNA damage can affect transcription efficiency and DNA replication and repair. We previously investigated the effects of the 3′-next flanking base (G*CT vs G*CA; G*, FABP, N-(2′-deoxyguanosin-8-yl)-4′-fluoro-4-aminobiphenyl; FAF, N-(2′-deoxyguanosin-8-yl)-7-fluoro-2-aminofluorene) on the conformation of arylamine-DNA lesions in relation to E. coli nucleotide excision repair (Jain, V., Hilton, B., Lin, B., Patnaik, S., Liang, F., Darian, E., Zou, Y., Mackerell, A. D., Jr., and Cho, B. P. (2013) *Nucleic Acids Res.*, 41, 869–880). Here, we report the differential effects of the same pair of sequences on DNA replication in vitro by the polymerases exofree Klenow fragment (KF-exo−) and Dpo4. We obtained dynamic $^{19}$F NMR spectra for two 19-mer modified templates during primer elongation: G*CA [d(5′-CTTACCATCG*CAACCTTC-3′)] and G*CT [d(5′-CTTACCATCG*CTACCCTTC-3′)]. We found that lesion stacking is favored in the G*CT sequence compared to the G*CA counterpart. Surface plasmon resonance binding results showed consistently weaker affinities for the modified DNA with the binding strength in the order of FABP > FAF and G*CA > G*CT. Primer extension was stalled at (n) and near (n + 1) the lesion site, and the extent of blockage and the extension rates across the lesion were influenced by not only the DNA sequences but also the nature of the adduct for the modiﬁed DNA with the binding strength in the order of FABP > FAF and G*CA > G*CT. In vitro, AF is processed by the high-fidelity polymerases but slows down replication, whereas the bulky N-acetylated AAF blocks the replication process and needs bypass polymerases for TLS. Thus, AF induces point mutations, and AAF produces point and frameshift mutations. In the DNA duplex, the N-deacetylated AF- and AFB-modiﬁed dG adducts tend to exist in equilibrium of two prototype conformers: B-type, in which the carcinogen occupies the major groove of a double helical DNA or a wedge (W) conformer, in which the hydrophobic structure is well placed in the narrow minor groove area. The S/B conformomeric equilibrium is dependent on the size, coplanarity, and topology of the arylamine carcinogen. For example, the AF/FAF adduct exists in an S/B mixture owing to the methylene linkage between the two aromatic rings that restricts the flexibility and makes it planar.
with efficient stacking ability (Figure 1A). Similarly, the more planar bulky amine adducts such as 1-aminopyrene and isoquinoline adopt predominantly S- and/or W-type conformers. In contrast, the flexible one-ring aniline and two-ring ABP adducts primarily adopt major groove B conformation. As shown in Figure 1A and B, ABP differs from AF in that it lacks a methylene bridge so that the biphenyl moiety is twisted. In addition, the nucleotide sequences surrounding the lesion also dictate the lesion-induced conformational heterogeneity, thus modulating repair and replication outcomes. Using the fluorinated analogues of AF, AAF, and ABP (e.g., FAF, FAAF, and FABP; Figure 1A), we have shown that these arylamine lesions undergo conformation-specific nucleotide excision repair (NER), i.e., the more thermodynamically unstable S-conformational lesions are repaired more efficiently than B-conformational lesions.

The mutagenicity of an adduct is also affected by its location within a DNA template and the neighboring bases. We recently studied the active site conformation of FAF in the presence of DNA polymerase β using 19F NMR, nucleotide insertion assays, and surface plasmon resonance (SPR). FAF in a single nucleotide gap adopted both S and B conformations, and heterogeneity was retained upon binding to the polymerase; however, it was altered by the incoming dNTP. One of the most striking examples of sequence effects involves the NarI sequence (5′-G1G2CG3C-3′), in which AF and AAF adducts have been shown to yield higher frequencies of frameshift mutations when they are associated with the third guanine (G3) of the sequence in E. coli. Interestingly, mutational frequencies are affected by the nature of the base at the 3′-next flanking position (N): the presence of dC opposite the lesion site results in a high rate of mutation compared to that of thymidine. In addition, structural studies have indicated that AF adducts adopt the S conformation when N = C but display heterogeneity when C is replaced by T.

Recently, we identified an unusual 3′-flanking T effect on a random 11-mer duplex sequence (5′-CCATCGTACC-3′), where the 3′-next flanking base was changed from T to A (5′-CCATCGTACCC-3′), the B conformation was adopted exclusively. Similar sequence effects also have been observed for FAF- and FAAF-modified G*N11-mer sequences. FAF...
modified G*CA and G*CT sequences adopted the S conformation in 66% and 90% of populations, respectively. Furthermore, E. coli repair studies have revealed that the bulky N-acetylated FAAF was repaired 3- to 4-fold more efficiently than the N-deacetylated FABP and FAF analogues, and all three adducts were repaired more efficiently in the G*CA sequence compared to that in the G*CT sequence context.

In the present study, we performed systematic dynamic 19F NMR, SPR, and primer elongation kinetic studies for the TLS of the G*CT and G*CA sequences containing FAF and FABP adducts (Figure 1). The results showed that the bulky lesions on the G*CT duplex exhibited greater populations of the stacked S conformation compared to the G*CA counterparts and that the S conformation decreased the binding affinity of complementary strands. Moreover, full length primer extension experiments were performed to investigate the similar sequence effects on in vitro translesion synthesis by two different polymerases: high fidelity replicative polymerase exofree Klenow fragment (Kf-exo−) and Dpo4. We found that the full-length primer extensions across the modified templates were significantly faster in the G*CA sequence, which commonly adopted the B conformation. Significant stalling occurred due to both lesions. The anti-B conformation of FABP showed more stalling at the prelesion (n − 1) site, whereas the S conformation of FAF showed more stalling at the lesion (n) site. The relative nucleotide insertion rates were significantly reduced immediately upstream of the lesion (n + 1 and n + 3). Together, the results of this study shed light on the roles of lesion-induced conformational heterogeneity in modulating the efficiency of TLS.

■ MATERIALS AND METHODS

Caution: 4-Aminobiphenyl and 2-aminofluorene derivatives are mutagens and suspected human carcinogens; therefore, they must be handled with caution.

Crude oligodeoxynucleotides (ODN, 1–10 μmol scale) in desalted form were purchased from Eurofins MWG Operon (Huntsville, AL). All HPLC solvents were purchased from Fisher Inc. (Pittsburgh, PA) and used as received. Kf-exo− and Dpo4 were received as gifts from Dr. Catherine Joyce (Yale University, CT) and Dr. F. Peter Guengerich (Vanderbilt University, TN), respectively.

Preparation of FAF- and FABP-Modified ODNs. FABP modification of 11-mer ODNs (5′-CCATXGXTACC-3′, X = A, T, C, G; Figure 1C) and FAF/FABP modification of 19-mer ODNs (5′-CTTTACCACCG*NACCATTC-3′, G* = FABP or FAF; N = A or T; Figure 1D) were performed by the procedures described previously. Briefly, 5–10 mg of N-acetoxy-N-trifluoroacetyl-7-fluoro-4-aminobiphenyl or N-acetoxy-N-trifluoroacetyl-7-fluorofluorene dissolved in absolute ethanol was added to a sodium citrate buffer (pH 6.0) containing 200–250 ODs of unmodified ODN and placed in a 37 °C shaker overnight. The modified strands were purified by a reverse phase-HPLC system which consisted of a Hitachi EZChrom Elite HPLC unit with an L2450 diode array detector and a Phenomenex Luna C18 column (150 × 10 mm, 5.0 μm). We employed a gradient system involving 3–15% acetonitrile in water (pH 7.0) with a flow rate of 2.0 mL/min.

Dynamic 19F NMR. Approximately 15 ODs of a modified strand was annealed with an equimolar amount of a complementary strand to produce different duplexes (Figure 3).
1C and D) that were lyophilized. The samples were then dissolved in 300 μL of typical pH 7.0 NMR buffer containing 10% D2O/90% H2O with 100 mM NaCl, 10 mM sodium phosphate, and 100 μM EDTA, and filtered into a Shigemi tube through a 0.2 μm membrane filter. All 1H and 19F NMR results were recorded using a dedicated 5 mm 19F/1H dual probe on a Bruker DPX400 Avance spectrometer operating at 400.0 and 376.5 MHz, respectively, using acquisition parameters described previously.11,12,32,33 Imino proton spectra (Figure S1, Supporting Information) at 5 °C were obtained using a phase-sensitive jump-return sequence and referenced relative to that of DSS.19F NMR spectra were acquired in the 1H-decoupled mode and referenced relative to that of CFCl3 by assigning external C6F6 in C6D6 at −164.9 ppm. Dynamic 19F NMR spectra (Figures 2, 3, and 4) were measured between 5 and 60 °C with an increment of 5−10 °C. Temperatures were maintained by a Bruker-VT unit with the aid of controlled boiling of liquid N2 in the probe.

**Primer Extension Assays.** Steady-state kinetic experiments were performed as described previously.34,35 Briefly, the primers (29−33-mers) were S′-radiolabeled using [γ-32P]ATP and T4 polynucleotide kinase (T4 PNK) following the manufacturer’s protocol. The S′-32P-labeled primer (100 pmol) was annealed to either an unmodified or adducted template oligonucleotide (120 pmol) by heating to 95 °C for 5 min and then slowly cooled to room temperature in 3 h. The primer−template sequence (100 nM) was incubated with Kf-exo− (2.5 nM) and Dpo4 (50 nM FABP and 100 nM FAF) polymerase in the presence of all four dNTPs. Aliquots were withdrawn at regular intervals of time, and the reaction was quenched as mentioned above. The extended products were separated on a 20% denaturing polyacrylamide gel.

**Running Start Experiments.** A 25-mer primer was used with the FAF- or FABP-modified dG at position 31 of the 44-mer template (Figure 5). The extension was performed using both Kf-exo− (2.5 nM) and Dpo4 (50 nM FABP and 100 nM FAF) polymerase in the presence of all four dNTPs. Aliquots were withdrawn at regular intervals of time, and the reaction was quenched as mentioned above. The extended products were separated on a 20% denaturing polyacrylamide gel.

**Steady-State Kinetics Experiments.** Steady-state kinetic parameters for incorporation of the nucleotide opposite the unmodified and FAF- or FABP-modified templates were determined by following the procedures reported previously.34,35 The reactions were performed at 20 °C. For the unmodified sequence, reactions were performed in a shorter time period of 1 min for correct nucleotide incorporation and for up to 45 min in the case of incorrect nucleotide incorporation. The percentage of primer extended in kinetic assays was determined by taking the ratio of extended primer to the total amount of primer (unextended + extended primer). The kinetic parameters $k_{cat}$ and $K_m$ were determined as described earlier.34,35

![Figure 4](https://dx.doi.org/10.1021/bi5003212) Dynamic 19F NMR spectra of FAF-modified 19-mer (A) G*CT and (B) G*CA duplexes at different single/double strand junction positions, i.e., n − 1, n, n + 3, and full (see Figure 1D).
Surface Plasmon Resonance (SPR). SPR measurements were conducted using a Biacore T200 instrument (GE Healthcare) in order to probe the effect of lesion-induced conformational heterogeneity using different lengths of complementary strands from n−1 to n+5 full duplex in a polymerase-free condition (Figure 7A). A 14-mer biotinylated DNA strand (5′-biotin-CTATC\text{\underline{G}}\text{C}N\text{\underline{A}}C\text{C}A\text{C}A\text{C}T-3′, N = T or A) was used for the SPR work. The template strands have the same sequence contexts as those utilized in the NMR studies (Figure 1D) except for the variation in the underlined portions (CC to CT; ATT to ATC) to avoid complications in complementary strand elongation (n−1, S′-GATGGTXG-3′; n, S′-GATGGTXGC-3′; n + 1, S′-GATGGTXGC\text{G}-3′; n + 2, S′-GATGGTXGC\text{A}-3′; n + 3, S′-GATGGTXGC\text{G}-3′; and n + 5, S′-GATGGTXGC\text{A}-3′; X = T for the GCA sequence, and X = A for the GCT sequence; see Figure 7A). The biotinylated oligonucleotides were modified with FABP and FAF by using the usual biomimetic procedure, purified by HPLC and characterized using MALDI-TOF mass spectrometry (Supporting Information, Figures S2 and S3). Streptavidin (SA) was immobilized on a CM5 chip using the EDC-NHS coupling method using the manufacturer’s protocol. The unbound SA was removed from the chip by injecting five 1 min pulses of NaOH (50 mM), and we stabilized the surface with running buffer for 30 min. The unmodified and modified oligo strands were coated over the SA surface to about 450 resonance units (RU) through a manual command mode. The stability of duplexes was monitored by injecting different concentrations of complementary strands (analytes) in HBS-P+ buffer (10 mM Hepes at pH 7.4; 150 mM NaCl; and 0.05% surfactant P20) at 25 °C. Dissociation rate constant (k_d) for each duplex were determined by fitting the data in a 1:1 model using the k_d-alone fitting method available in Scrubber software, version 2.0 (Myszka and collaborators; BioLogic Software) (Supporting Information, Table S10). The concentration-independent parameter k_d was calculated by fitting the initial 100 s of the dissociation curves (Supporting Information, Figure S4). The goodness of the fit was determined from the residual standard deviation.

### RESULTS

**Experimental DNA Sequences.** The chemical structures of ABP, AF, and AAF and their corresponding fluorine analogues, FABP, FAF, and FAAF, are shown in Figure 1A. In addition, the 11-mer G[\text{FABP}]CT duplex sequence used in our previous repair studies and the flanking sequence variations used are shown in Figure 1C.\textsuperscript{30} These modified DNA strands were annealed with complementary strands to form duplexes for the \textsuperscript{19}F NMR measurements. The CG*CN (N = A or T) series designed for the TLS experiments are shown in Figure 1D. The original 11-mer oligonucleotides were extended to 19-mers to improve thermal stability for the n−1 and n duplexes. As shown in Figure 1D, four 19-mer G*CN strands (G* = FABP or FAF; N = A or T) were each annealed with complementary strands of variable lengths (n−1, n, n + 3, and n + 9; n is the lesion site) to create four discrete model TLS systems.

**Flanking Base Effects.** The rationale behind this study is to investigate the uniqueness of 5′- or 3′-flanking C in the -CG*CT-sequence context in promoting lesion-induced conformational heterogeneity. As such, the 3′- and 5′-flanking C of CG[\text{FABP}]\text{C}N (N = A or T) was systematically switched to A, T, or G.
Conformational Heterogeneity at Simulated TLS Single/Double Strand Junctions. The goal of this study was to determine whether the S/B-conformational heterogeneity in the G*CA and G*CT duplexes also presents at the single/ double strand junctions formed during the primer elongation in TLS process. Therefore, we conducted 19F NMR experiments in buffer-only solutions in the absence of a polymerase on four discrete 19-mer G*CA duplexes (G* = FABP or FAF; N = A or T) (Figure 1D). These four model TLS systems were prepared by annealing the 19-mer modified strand with complementary strands of variable lengths (n − 1, n, n + 3, and n + 9; n is the lesion site).

FABP. Figure 3A−B shows the dynamic 19F NMR spectra (5−50 °C) for TLS of the FABP-modified G*CT* and G*CA* sequences (Figure 1D) at various elongation positions with a temperature range between 5−50 °C. Signals were assigned according to published procedures based on relative chemical shifts, dynamic NMR signal patterns, and H/D isotope effects.13,35 19F shielding is a hallmark of the van der Waals interactions and the ring current effects caused by the carboxyl group moiety within the stacked and bulge duplexes (S-type conformation). We have studied a number of 19F NMR spectra of arylamine-modified duplexes (including FABP and FAF) in various sequence settings and in all cases found that the fluorine of the S-type conformer resonates upfield relative to that of the external binding B-type conformer.12,35 The same trend has been observed with various fully paired and deletion duplexes modified with the bulky N-acetylated FAAF, which specifically revealed a mixture of B, S, and W conformations in the −115.0 to −115.5 ppm, −115.5 to −117.0 ppm, and −116.5 to −118.0 ppm ranges, respectively.21 In contrast, the N-deacetylated FABP and FAF adducts adopted an interchangeable mixture of the B- and S-conformers. Accordingly, the signals in Figures 3 and 4 were assigned as the B (downfield)- and S (upfield)-type conformers, respectively.

The fully paired 19-mer G*CT* and G*CA* duplexes (Figure 1D), which represent the end points of TLS, displayed comparable conformational differences (Figure 3A−B) similar to those of their respective 11-mer duplexes (Figure 2A). Populations of G*CT* sequences in the S conformation decreased significantly from the 11-mer (60%) to the 19-mer (33%) duplexes. The 19F signal patterns for the n + 3 and full duplex sequences for both G*CA* and G*CT* did not change much. The 19F NMR signals at the n and n − 1 sites were designated as B-like (B*) and S-like (S*), respectively, based on their chemical shift similarities to the duplexes. The B (B*)- and S (S*)-proportions were maintained as elongation progressed from n − 1 to full in the G*CT* TLS series. Interestingly, we observed a significantly larger population of the S conformation at the n − 1 and n positions as compared to that in the n + 3 or n + 6 position in the G*CA* series (Figure 3B). The conformer population ratios at different positions for each modified strand are summarized in Table 1.

| duplex | sequence context | FABP | FAF |
|--------|-----------------|------|-----|
| n − 1  | G*CA (n)        | B* (82%) | S* (18%) |
|        | G*CT (n)        | B* (70%) | S* (30%) |
| n      | G*CA (n)        | B* (76%) | S* (24%) |
|        | G*CT (n)        | B* (64%) | S* (36%) |
| n + 3  | G*CA (n)        | B (100%) | B (42%) |
|        | G*CT (n)        | B (65%)  | S (35%) |
| full   | G*CA (n)        | B (95%)  | B* (5%)  |
|        | G*CT (n)        | B (67%)  | S (33%)  |

FAF. The dynamic 19F NMR spectra for the FAF-modified G*CT* and G*CA* TLS series (n = 1 to full) are shown in Figure 4A−B. In both sequences, FAF exhibited B (B*)- and S (S*)-conformation heterogeneity, and G*CT* had approximately 16% more S-conformers than G*CA* in the full duplex. Like the FABP adduct, template strand elongation from the 11-mer to the 19-mer duplex reduced the G*CT* S-conformer population from 90% to 80%. At n + 3, the S-conformer population differed by approximately 23% between the two sequences, primarily due to the increase in the B-conformer population in the G*CA* template (Figure 4B). Moving from the n + 3 to the n and the n − 1 positions, FAF exhibited two 19F signals. The S conformation gap was further increased to 34% at the lesion site (n), mostly due to the increase in the B-conformer population of G*CA* (Figure 4B and Table 1). However, the n − 1 series displayed very similar 19F characteristics and a 3:7 ratio of B- and S-conformations. The conformational results of the FAF-induced TLS system are summarized in Table 1.

Primer Extension Experiments. The purpose of primer extension experiments was to determine the impact of lesion-induced conformational heterogeneity on the polymerase efficiency during TLS. Therefore, experiments were performed using 44-mer G*CA* (G* = FABP or FAF; N = A or T) templates. As shown in Figure 5A, 32P-labeled 25-mer primers were annealed to the templates, and primer extension was carried out in the presence of all four dNTPs and polymerases (KF-exo− or Dpo4). Running start experiments were specifically carried out to determine the overall impact on the rate of full length primer extension across the lesion and the major blockage sites for polymerase.
Running Start Experiments on FABP Adducts with Kf-exo Polymerase. The products of Kf-exo-mediated primer extension across FABP at different time intervals at room temperature are shown in Figure 5B (left). Kf-exo was able to extend the primer to a full-length 44-mer across G*C and G*CA in 2–5 min. However, two major stalls in extension were observed: one at the nucleotide before the lesion site (n−1, 30-mer) and the other at the lesion site (n). The blockage at n was stronger than that at n−1. The full-length product was formed from this blocked primer, although the primers at n−1 persisted even after incubation for 60 min. Additionally, both sequences exhibited unusual blockage at the site five bases upstream of the lesion (red arrows). Although the sequences G*CA and G*CT both showed similar blockage patterns, the durations of the blockages were different. In the initial few minutes, G*CT displayed a significant increase in blockage at the lesion site (green arrow), which decreased after the full-length product was produced. A similar pattern was observed for G*CA, but TLS occurred in a shorter time (see expanded red rectangular insets in Figure 5B). The prolonged blockage at the lesion site (n) was converted to a full-length product with G*CA displaying a faster rate of extension than G*CT.

Running Start Experiments of FAF Adducts. The results of the running start experiments for the FAF adducts are shown in Figure 5B (right). Here, the primers were blocked at n and, to a lesser extent, at n−1 and n+1. Regardless, Kf-exo was able to extend more efficiently across the FAF adduct compared to that across FABP. In addition, a blunt-end addition was also observed. A similar sequence effect was observed with the FAF adduct, and G*CA showed a faster rate of extension compared to that of G*CT (see expanded red rectangular insets, Figure 5B). As shown in Figure 5C (right), primer extension was stalled at two sites, n−1 and n, in the Dpo4-mediated primer extension. Unlike with Kf-exo, there was no blockage at n+1; however, a slightly greater rate of extension across G*CA compared to that across G*CT was observed.

Steady-State Kinetics. Steady-state kinetic experiments in the presence of Kf-exo were performed in order to investigate the roles of lesion-induced conformational heterogeneity in nucleotide insertion kinetics. As shown in Figure 6A, lesions were positioned one nucleotide downstream of the template base (n−1) and compared to Kf-exo, the extent of full-length extension was significantly less, even at a high enzyme concentration (100 nM, data not shown). In addition, a stronger blockage occurred at n−1 with Dpo4, compared with that using Kf-exo. As with Kf-exo, the G*CT sequence exhibited an extended blockage at n (green arrow), resulting in slower extension of the primer to full length (rectangular insets, Figure 5C).
Table 2. Summary of the Insertion Efficiency against the FABP- and FAF-Modified Template in the Presence of KmexO at Different Insertion Sites in the G*CA and G*CT Sequence Contexta

| insertion site | sequence context | insertion efficiency (f_ins)b | FABP | FAF |
|---------------|------------------|-----------------------------|------|-----|
| n − 1         | G*CA             | dGTPw                        | 2.2 × 10⁻¹ | 4.0 × 10⁻¹ |
|               | G*CT             | dGTPw                        | 2.2 × 10⁻¹ | 3.1 × 10⁻¹ |
| n             | G*CA             | dCTP                         | 1.1 × 10⁻¹ | 4.6 × 10⁻² |
|               | G*CT             | dCTP                         | 3.0 × 10⁻⁴ | 1.5 × 10⁻⁴ |
| n + 1         | G*CA             | dG*+dC d²                  | 9.6 × 10⁻⁶ | 4.8 × 10⁻⁵ |
|               | G*CT             | dG*+dC d²                  | 9.1 × 10⁻⁶ | 5.1 × 10⁻⁵ |
| n + 3         | G*CA             | dG*+dC d²                  | 4.9 × 10⁻⁴ | 1.5 × 10⁻⁴ |
|               | G*CT             | dG*+dC d²                  | 2.0 × 10⁻² | 1.3 × 10⁻² |

aDetailed kinetic analysis with error limits are provided in the Supporting Information (Tables S1 to S8). bThe relative insertion efficiency f_ins = \( \frac{k_{cat}/K_m\text{modified}}{k_{cat}/K_m\text{unmodified}} \) − incoming nucleotide. cTemplate − primer terminus. "Incoming sequence, it was 1.5 × 10⁻², indicating that the dCTP insertion was preferred over the dATP insertion by 31-fold in G*CA and 8-fold in G*CT (Table 2).

For the FABP adducts, the f_ins values of dGTP opposite C at the n + 1 position were 9.6 × 10⁻⁶ and 9.1 × 10⁻⁶ in the G*CA and G*CT sequences, respectively. However, when the incorrect dA was present opposite the lesion, nucleotide insertion was blocked for both sequences and the efficiency could not be determined (ND, Table 2). For the FAF adducts (Figure 6B), the efficiencies of the correct nucleotide dGTP at n + 1 were similar in both the G*CA (4.8 × 10⁻³) and G*CT (5.1 × 10⁻³) sequences. The f_ins values for dGTP at n + 1 were reduced by 6- and 242-fold in the G*CA and G*CT sequences, respectively, when the incorrect dA was paired at the lesion site. However, the effect of a lesion at n + 3 only minimally affected the efficiency. The f_ins values for the FABP adducts were 4.9 × 10⁻¹ (G*CA) and 8.0 × 10⁻² (G*CT), and the f_ins values for the FAF adducts were 1.5 × 10⁻¹ (G*CA) and 6.8 × 10⁻² (G*CT) for the correct insertions (Table 2). For dA opposite the lesion site, the f_ins values at n + 3 between the G*CA and G*CT sequences differed by 10- and 3-fold in the FABP and FAF adducts, respectively.

**SPR Binding Affinity Measurements for Simulated TLS Single/Double Strand Junction Duplexes.** The objective of SPR experiments was to measure real-time association between

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Figure 7. (A) Primer−template duplex sequences for SPR measurements. SPR binding responses of complementary strands opposite unmodified and modified DNA for (B) GCA and (C) GCT sequence contexts. Strands concentrations are 50 nM (n − 1), 100 nM (n), 150 nM (n + 1), 200 nM (n + 2), and 250 nM (n + 3 and n + 5 full length complementary strands).
template and complementary strands in the absence of a polymerase. Binding data were obtained by injecting different lengths (8-mer n−1, 9-mer n, 10-mer n+1, 11-mer n+2, 12-mer n+3, and 14-mer n+5) of complementary strands to lesions containing 14-mer template strands (GCT vs GCA) coated on a streptavidin chip. For each elongation, it was necessary to employ different concentrations of complementary strands to achieve steady-state associations (Figure 7: 50 nM (n−1), 100 nM (n), 150 nM (n+1), 200 nM (n+2), and 250 nM (n+3 and n+5). Following each binding phase, the template-containing chip was washed with buffer as described in Materials and Methods. Figure 7B shows the representative sensograms as a function of different lengths of complementary strands. As a result, the usual Kd binding (k_d/k_a) kinetics of duplex formation for each elongation was not obtained. Since all of the experimental conditions included are identical for each elongation, direct qualitative comparison of RU values for each elongation is presented. Similarly, dissociation rates (k_a) are concentration-independent and thus could be used to estimate binding strengths for different lengths of complementary strands. Supporting Information, Figure S4 shows curve data analysis using Scrubber (BioLogic Software). Supporting Information, Table S10 lists the respective dissociation rate constants. The results are also summarized graphically in Figure 8 as plots of log k_d vs various elongation points for GCA and GCT sequences.

**FAFBP.** SPR data shows generally greater binding activities (RU) (Figure 7B) and k_d (Figure 8) values as the complementary strand length increased from n−1 to a full duplex in both sequences, with increasing duplex stability. Overall, we observed little difference between the unmodified GCA and GCT sequences. The effect of a lesion modification was evident when modified duplexes, which showed a substantial reduction in RU responses for the n to n+3 sites (Figure 7B). Interestingly, however, the FABP modification caused an increase of RU values at the pre lesion n−1 site compared to that in the unmodified controls. These results indicate a lesion-induced stabilization which is in agreement with the slower dissociation at the pre lesion site (Supporting Information, Table S10 and Figure 8). The effect of a lesion modification was evident with faster dissociations (k_a) than the unmodified one at the n to n+5 site (Figure 8 and Supporting Information, Figure S10). For example, the binding strength for G[FAFBP]CT at n+1 was decreased 3.3 (0.0443/0.0113)- and 37 (0.0443/0.0012)-fold relative to the G*CA and the unmodified GCT sequences, respectively (Supporting Information, Table S10). The sequence effect was found to be consistently around 3−4-fold different between the G*CT and G*CA sequences, respectively: n+2 (3.8-fold, 0.0307/0.00802), n+3 (3.7-fold, 0.00142/0.00378), and n+5 (3.6-fold, 0.00571/0.00156).

**FAF.** A similar lesion and sequence effects were observed for the FAF adducts. The RU intensities of all the elongations except for n−1 were generally suppressed compared to those of the unmodified controls (Figure 7). For example, the binding strength for G[FAF]CT at n+1 was decreased 2.9 (0.0326/0.0102)- and 27.1 (0.0326/0.0012)-fold relative to the G*CA and the unmodified GCT sequences, respectively (Supporting Information, Table S10). As in the case of the FABP adducts, the sequence effect was maintained at 2−3-fold differences between the G*CT and G*CA sequences.

## DISCUSSION

We studied a range of structural, biochemical, and biophysical properties of a pair of arylamine-DNA lesions, one (ABP) derived from the very important human bladder carcinogen 4-amino-biphenyl that is present in tobacco smoke and a second (AF) stemming from the model carcinogen 2-aminofluorene. These two lesions, bound to the C8-position of dG, differ only in that the biphenyl moiety lacks a methylene bridge that restrains the aminofluorene ring to planarity so that the biphenyl is twisted. Specifically, we characterized lesion-induced TLS involving two unique G*CA and G*CT sequences by dynamic 19F NMR, primer extension studies with two representative polymerases, Kf-exo− and Dpo4, with detailed steady state kinetic parameters, and surface plasmon resonance measurements.

### 3′-Next Flanking Base Effect on Lesion-Induced S/B Heterogeneity during Simulated TLS

The fully paired 19-mer G[FAF]CT duplex exhibited a 67:33 ratio of B- and S-conformers (Figure 3A and Table 1), consistent with the 60:40 B:S ratio that was previously observed for the 11-mer counterparts. A similar conformer ratio was intact at the n−1 and n sites. In contrast, the G[FAFBP]CT duplex adopted the B conformation predominantly (95% B:5% S) (see n+6, Figure 3B). The FAF-modified 19-mer duplex (Figure 4) exhibited S/B heterogeneity comparable to that of the previously studied 11-mer:38 90% S vs 80% S for G*CT and 66% S vs 64% S for G*CA. Similar conformational heterogeneity persisted even at the n−1 and n positions.

In general, the larger S-conformer population for the FAF adducts compared to the FABP adducts may be attributed to the presence of single carbon atom methylene bridging in FAF. This linkage restricts the twisting between the two aromatic rings that is possible in FABP, thus enhancing the stacking surface (Figure 1A,B). This simple topological difference is fascinating in its impact on the structural and biochemical properties of the lesion. Similar conformational behavior is observed with other planar bulky amine adducts such as 1-aminopyrene (AP) and the food-borne heterocyclic amine isoquinoine (IQ), which favor S and/or wedge (W)-type conformers. On the contrary, the AF analogue imidazopyrindine (PhIP) occupies the minor groove largely due to the freely rotatable phenyl at C6 position. These results clearly signify the...
role of a single linkage in dictating the conformations adopted by FAF.

A large S-conformational gap was noted in G*CA vs G*CT for FABP (30% vs 18%) at n − 1, while no such difference (67% vs 68%) was observed for the FAF adduct (Figures 3–4 and Table 1). At the lesion site n, FAF in the G*CT sequence exhibited a 34% higher S-conformer population than G*CA (Table 1). Complementary strand elongation from n to n + 3 increased the S-conformer population in both G*CA (42–58%) and G*CT (76–81%) for the FAF adduct. For fully paired duplexes, the S-conformer population of G*CA increased to 64%, whereas the conformation of G*CT was largely unaffected (80%). Taken together, these results indicated that lengthening the duplex sequence from 11 to 19 nucleotides had no major effects on the S/B heterogeneity of FABP or FAF adducts in the G*CA or G*CT sequences. The unusual 3′-next flanking base effects observed in the fully paired duplex were largely maintained at various stages of complementary strand elongation, including the preinsertion (n − 1) and lesion (n) sites.

We performed SPR experiments to investigate the impact of lesion heterogeneity on complementary strand binding (Figure 7). SPR is a chip-based and label-free procedure that allows real-time monitoring of various replication and repair interactions involving DNA lesions.27-43 SPR has also been used to study DNA assembly, DNA hybridization,44 and triplex formation.46 We obtained simple sensorgrams as a function of different complementary strands for each elongation, and consequently, we were unable to conduct the usual global binding analysis using $K_d$ affinity values (see Results). Instead, we utilize RU intensities (Figure 7) and $k_d$ dissociation rates (Supporting Information, Table S10) to assess lesion and sequence effects. In general, the binding strength of complementary strands toward the unmodified DNA sequence was higher than the modified sequences. In contrast, the effect of FABP or FAF modification was sensed primarily around the lesion (n − 1 to n + 2), but then the effect was reduced at n + 3 and full (n + 5) duplexes. It is worth noting that the RU for G*CT were consistently less than those for G*CA throughout the elongation. There was a trend in lesion-induced differences in terms of RU values in the G*CA/ G*CT sequence context, i.e., G*CT was a DNA destabilizer compared to G*CA. Figure 8 shows graphical plots of log $k_d$ dissociation rates vs various elongation points for the G*CA and G*CT TLSs. It is clear that the lesion effect is seen throughout the simulated elongation process (n to n + S) with greater impact for the G*CT compared to that for G*CA. As shown in Figure 8, however, the nature of the lesion (FABP vs FAF) had no discernible effect on $k_d$ values. Although no polymersases were involved, these SPR results are in line with the NMR and gel assay results as the greater S-conformation of the G*CT sequence posed a major hindrance toward the binding affinity. In other words, the lesion effect was propagated primarily upstream of the template strand and gradually diminished once the adduct came out of the pocket of the polymerase active site.35,34,47

Conformational Effects on Stalling during TLS in the Presence of Kf-exo and Dpo4. Kf-exo is a high-fidelity replicative polymerase lacking the 3′–5′ proofreading exonuclease activity used for TLS of various types of DNA damage, including bulky arylamine lesions.34 High fidelity polymerases generally are blocked by bulky lesions.5 The DNA polymerase Dpo4 from Sulfolobus solfataricus is a Y-family bypass polymerase, characterized by a loose active site that is accessible to solvent, thus known to promote TLS.48 The running start results (Figure S5) showed that primer extension was stalled at the lesion site (n) and near the lesion site (n − 1 and n + 1). The extent of blockage was clearly influenced by the nature of adduct structure, i.e., both FABP and FAF lesions blocked Kf-exo at n − 1 and n and additionally at n + 1 for the FAF adducts. Both lesions strongly blocked at the lesion site (n); however, they exhibited similar sequence effects, i.e., the G*CA sequence exhibited a greater extension efficiency compared to that of G*CT. With Dpo4, the lesions stalled primer extension only at n − 1 and n, but the major blockage site occurred at n − 1 for FABP, compared to n for FAF. The different stalling characteristics could be due to the spacious nature of the Dpo4 active site compared to that of Kf-exo. The high steric flexibility of Dpo4 has been shown to allow bulky and distorted DNA lesions to proceed through the active site differently.49–52 Surprisingly, however, we observed that Dpo4 exhibited a significantly slower rate of primer extension for FABP, even at higher enzyme concentrations (Figure 5C, left). Dpo4 also causes deletions with bulky adducts.

The G*CT and G*CA sequences displayed similar blocking patterns (Figure 5), consistent with their conformational profiles around the lesion sites (Table 1). However, full-length extension across the G*CA sequence was faster compared to that across the G*CT sequence of both lesions as well as in the presence of Kf-exo and Dpo4 (insets, Figure 5B and C). As mentioned above, the polymerase prefers the modified dG in the anti-glycosyl conformation for efficient replication across the lesion. The relatively slower extension across G*CT may be due to the higher percentages of the S-conformer at the lesion site.

Hsu et al.47 have reported the crystal structures of AF during accurate replication by BF, a high-fidelity DNA polymerase analogous to Kf-exo. They found that the AF adduct exhibited a syn conformation at the preinsertion site (n − 1) and underwent a transition to an anti-conformation at the insertion site (n), allowing it to base pair with an incoming dCTP. According to our NMR results (Figures 3–4; Table 1), the planar FAF at n − 1 exhibited a greater percentage of the syn-S-type conformation, while the nonplanar FAF favored the anti-B-type conformation (Table 1). It is plausible that the B-conformer of FABP stalls at n − 1, thus resulting in the accumulation of an extension product at the prelesion site. We have reported previously that FAF in the TG*A sequence adopted a higher percentage of the B-conformer and exhibited a stronger stalling effect at the n − 1 site, while the S-conformer of CG*A stalled primer extension at the lesion site (n).53 In addition, the B-conformer in the FAF-modified TG*A sequence context favored a misinsertion of dATP by 80-fold compared to the CG*A sequence due to the presence of 5′-T.

Lesion and Sequence Effects on Insertion Efficiencies during TLS in the Presence of Kf-exo. Both FABP and FAF lesions exhibited different relative insertion efficiencies ($f_{\text{ins}}$), depending on the primer positions (Table 2). The impact of the lesion was generally minimal at the prelesion (n − 1) and lesion (n) sites. The conformational characteristics of FABP and FAF and their $f_{\text{ins}}$ values for the matched (dG*:dC) series were comparable for the G*CA and G*CT sequences (Figure 6). Although both FABP and FAF at the lesion site (n) favored insertion of dCTP, the preference was more prominent for FABP than for FAF. This result may be due to the high S-conformer percentage of the FAF adduct (Table 1). Similar trends in extension efficiency patterns were observed for the dA-mismatched series, although extremely low $f_{\text{ins}}$ values were observed for FABP at the lesion site (ND, Table 2). The most dramatic lesion effects were observed at the n + 1 site. We noted a slightly greater lesion effect of FAF over FABP in both G*CA and G*CT sequences.
G*CT sequence contexts. These results are consistent with the blocking patterns observed in the running start and SPR binding results discussed above and could be attributed to the relatively higher S-conformer population of the FAF adducts compared to the FABP adducts (Table 1).

Figure 9 shows difference factor (DF) histograms to compare the relative insertion efficiency between two sequences (G*CA vs G*CT) at various insertion points in the matched (dG*:dC) series. DF is defined as the f_{ins} of G*CT over G*CA (f_{ins} G*CT/ f_{ins} G*CA), and thus, DF below one signifies a better insertion efficiency in the G*CA sequence compared to that of the G*CT at a particular insertion site, whereas a factor above one has opposite efficiencies. It is clear that the sequence effect was most dramatic at the lesion site (n), i.e., DF = 0.18 and 2.61 for the FABP and FAF adducts, respectively. We observed that the sequence effects were negligible (DF ≈ 1.0) at the pre lesion site (n − 1) (Figure 9 and Supporting Information, Table S9). These results seem to be in line with the NMR results showing that the nature of the 3′-next flanking base (G*CA vs G*CT) did not affect the conformational patterns at n − 1. In the FABP adducts at the lesion site, dATP insertion did not lead to any significant sequence effects (Table 2). In contrast, in the FAF adducts, the G*CT sequence favored dATP insertion by 11-fold compared to that of G*CA. This result is probably due to the relatively higher population of the S-conformer (76%) in the G*CT sequence compared to that of the G*CA (42%). Extension of the dG insertion at the n + 1 site was significantly reduced due to perturbations at the active site of the polymerase. As mentioned above, an extremely low insertion rate was noted for the FABP adducts in the dA-mismatched series. The insertion of dATP at n + 1 was observed for the FAF adducts, but with low efficiency. However, no significant sequence effects were observed at the n + 1 position in either the matched (Figure 9, DF = 0.95 and 1.06 for the FABP and FAF adducts, respectively) or the mismatched series. The FABP in the G*CA series at n + 3 adopted exclusively the B-conformation, yet exhibited a 6-fold greater insertion extension efficiency of dATP than G*CT for the matched series (Figure 9, DF = 0.16). In contrast, the planar FAF showed only a 16% difference in the B:S population ratios (Table 1), thus resulting in a minimal sequence effect (DF = 0.45) (Figure 8).

In summary, we conducted systematic 19F NMR, SPR, and primer elongation kinetic studies for the TLS of the G*CT and G*CA sequences containing the bulky arylamine FAF and FABP adducts. The results are summarized in Figure 10. DNA adducts in the G*CT duplex exhibited greater populations of the S-conformer compared to that in the the G*CA duplex with the same adducts. We also found the 3′-next flanking T in the former promotes lesion stacking, thus supporting the greater S conformation. These lesions exhibited unique sequence-dependent conformational heterogeneities at various elongation positions, including the replication fork that differentially contributes to template—primer bindings and insertion efficiency during TLS. In addition, the SPR binding results revealed that the presence of adduct in the stacked conformation decreases the binding affinity of the complementary strands in the order of unmodified > FABP > FAF and G*CA > G*CT. Full-length primer extensions across the modified templates were significantly faster in the G*CA sequence, which commonly adopted the B-conformation. Primer extension was stalled at (n) and near (n − 1 and n + 1) the lesion site, and the extent of blockage and the extension rates across the lesion were influenced by the DNA sequences as well as the nature of the lesion (FAF vs FABP) and the polymerase employed (Kf-exo- vs Dpo4). For example, with Kf-exo the anti-B-conformic FABP showed stalling at both the pre lesion (n − 1) and lesion (n) sites, whereas the syn-S-conformic FAF showed stalling at all three sites (n − 1, n, and n + 1). Moreover, the relative nucleotide insertion rates (f_{ins}) were small at n − 1 but were significantly reduced immediately 5′ upstream of the lesion site (n + 1 and n + 3). Together, these results demonstrated the roles of lesion-induced conformational heterogeneity in modulating the efficiency of TLS.

Figure 10. Summary of present results in terms of lesion stacking, primer binding, lesion stalling, and nucleotide insertion rates.
ASSOCIATED CONTENT

Supporting Information

Imino proton and full dynamic 31P NMR spectra of FABP-modified 11-mer duplexes, MALDI-TOF spectra of biotinylated templates, and simulated SPR sensograms; kinetic parameters for dNTP insertion at different primer positions; and the dissociation constant values ($K_d$) obtained from SPR binding experiments. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

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ABBREVIATIONS

FAF, N-(2′-deoxyguanosin-8-yl)-7-fluoro-2-aminofluorene; FABP, N-(2′-deoxyguanosin-8-yl)-4′-fluoro-4-aminobiphenyl; K-fexo+, exonuclease deficient Klenow fragment; SPR, surface plasmon resonance; TLS, translesion synthesis.

REFERENCES

(1) Luch, A. (2005) Nature and nurture - lessons from chemical carcinogenesis. Nat. Rev. Cancer 5, 113–125.

(2) Friedberg, E. C., Walker, G. C., Siede, W., Wood, R. D., Schultz, R. A., and Ellenberger, T., Eds. (2006) DNA Repair and Mutagenesis, 2nd ed., ASM Press, Washington, DC.

(3) Hubscher, U., and Maga, G. (2011) DNA replication and repair bypass machin. Cell 146, 627–635.

(4) Cho, B. P., Beland, F. A., and Kadlubar, F. F. (1990) Handbook of Experimental Pharmacology, Spring-Verlag, Heidelberg, Germany.

(5) Hefflich, R. H., and Neff, R. E. (1994) Genetic toxicity of 2-acetylaminofluorene, 2-aminofluorene and some of their metabolites and model metabolites. Mutat. Res. 318, 113–114.

(6) Shibutani, S., Suzuki, N., and Grobman, A. P. (1998) Mutagenic specificity of (acetylamino)fluorene-derived DNA adducts in mammalian cells. Biochemistry 37, 12034–12041.

(7) Cho, B. P. (2004) Dynamic conformational heterogeneities of carcinogen-DNA adducts and their mutagenic relevance. J. Environ. Sci. Health, Part C 22, 57–90.

(8) Cho, B. P., Beland, F. A., and Marques, M. M. (1992) NMR structural studies of a 15-mer DNA sequence from a ras protooncogene, modified at the first base of codon 61 with the carcinogen 4-aminobiphenyl. Biochemistry 31, 9587–9602.

(9) Jain, N., Li, Y., Zhang, L., Meneni, S. R., and Cho, B. P. (2007) Probing the sequence effects on N-aril-induced −2 frameshift mutagenesis by dynamic 19F NMR, UV, and CD spectroscopy. Biochemistry 46, 13310–13321.

(10) Meneni, S. R., Shell, S. M., Gao, L., Jurecka, P., Lee, W., Sponer, J., Zou, Y., Chiarelli, M. P., and Cho, B. P. (2007) Spectroscopic and theoretical insights into sequence effects of aminofluorene-induced conformational heterogeneity and nucleotide excision repair. Biochemistry 46, 11263–11278.

(11) O’Handley, S. F., Sanford, D. G., Xu, R., Lester, C. C., Hingerty, B. E., Broyde, S., and Krugh, T. R. (1993) Structural characterization of an N-acetyl-2-aminofluorene (AAF) modified DNA oligomer by NMR, energy minimization, and molecular dynamics. Biochemistry 32, 2481–2497.

(12) Zhou, L., Rajabzadeh, M., Traficante, D. D., and Cho, B. P. (1997) Conformational heterogeneity of alyriline-modified DNA: 19F NMR evidence. J. Am. Chem. Soc. 119, 5384–5389.

(13) Mao, B., Vyas, R. R., Hingerty, B. E., Broyde, S., Basu, A. K., and Patel, D. J. (1996) Solution conformation of the N-(deoxyguanosin-8-yl)-1-aminoapryne ([API[dl]] adduct opposite dc in a DNA duplex. Biochemistry 35, 12659–12670.

(14) Wang, F., DeMuro, N. E., Elmasi, C. E., Stover, J. S., Rizzo, C. J., and Stone, M. P. (2006) Base-displaced intercalated structure of the food mutagen 2-amin-3-methylimidazo[4,5-f]quinoline in the recognition sequence of the Narl restriction enzyme, a hotspot for −2 bp deletions. J. Am. Chem. Soc. 128, 10085–10095.

(15) Wang, F., Elmasi, C. E., Stover, J. S., Rizzo, C. J., and Stone, M. P. (2007) DNA sequence modules the conformation of the food mutagen 2-amin-3-methylimidazo[4,5-f]quinoline in the recognition sequence of the Narl restriction enzyme. Biochemistry 46, 8498–8516.

(16) Cho, B. P., Beland, F. A., and Marques, M. M. (1994) NMR structural studies of a 15-mer DNA duplex from a ras protooncogene modified with the carcinogen 2-aminofluorene: conformational heterogeneity. Biochemistry 33, 1373–1384.

(17) Shapiro, R., Ellis, S., Hingerty, B. E., and Broyde, S. (1998) Effect of ring size on conformations of aromatic amine-DNA adducts: the aniline-C8 guanine adduct resides in the B-DNA major groove. Chem. Res. Toxicol. 11, 335–341.

(18) Meneni, S. R., D’Mello, R., Norigian, G., Baker, G., Gao, L., Chiarelli, M. P., and Cho, B. P. (2006) Sequence effects of aminofluorene-modified DNA duplexes: thermodynamic and circular dichroism properties. Nucleic Acids Res. 34, 755–763.

(19) Patnaik, S., and Cho, B. P. (2010) Structures of 2-acetylaminofluorene modified DNA revisited: Insight into conformational heterogeneity. Chem. Res. Toxicol. 23, 1650–1652.

(20) Meneni, S., Shell, S. M., Zou, Y., and Cho, B. P. (2007) Conformation-specific recognition of carcinogen-DNA adduct in Escherichia coli nucleotide excision repair. Chem. Res. Toxicol. 20, 6–10.

(21) Geacintov, N. E., Cosman, M., Hingerty, B. E., Amin, S., and Patel, D. J. (1997) NMR solution structures of stereoisometric covaent polycyclic aromatic carcinogen-DNA adduct: principles, patterns, and diversity. Chem. Res. Toxicol. 10, 111–146.

(22) Liu, Y., Reeves, D., Kropachev, K., Cai, Y., Ding, S., Kolbanovskiy, M., Kolbanovskiy, A., Mello, R., Norigian, G., Baker, G., Zou, Y., Chiarelli, M. P., and Cho, B. P. (2006) Probing for DNA damage with beta-hairpins: similarities in incision efficiencies of bulky DNA adducts by prokaryotic and human nucleotide excision repair systems in vitro. DNA Repair 10, 684–696.

(23) Geacintov, N. E., Cosman, M., and Patel, D. J. (2014) Probing DNA damage with beta-hairpins: similarities in incision efficiencies of bulky DNA adducts by prokaryotic and human nucleotide excision repair systems in vitro. DNA Repair 10, 1126–1136.

(24) Kolbanovskiy, M., Kolbanovskiy, A., Bolton, J. L., Broyde, S., and Geacintov, N. E. (2014) Nucleotide excision repair of 2-aminofluorene and 2-amino(8-C8)-guanine adducts: molecular dynamics simulations elucidate how lesion structure and base sequence context impact repair efficiencies. Nucleic Acids Res. 40, 9675–9690.

(25) Jain, V., Hilton, B., Patnaik, S., Zou, Y., Chiarelli, M. P., and Cho, B. P. (2012) Conformational and thermodynamic properties modulate the nucleotide excision repair of 2-aminofluorene and 2-acetylamino-
fluorene dG adducts in the NarI sequence. *Nucleic Acids Res.* 40, 3939–3951.
(27) Vaidyanathan, V. G., Liang, F., Beard, W. A., Shock, D. D., Wilson, S. H., and Cho, B. P. (2013) Insights into the conformation of aminofluorene-deoxyguanine adduct in a DNA polymerase active site. *J. Biol. Chem.* 288, 23573–23585.
(28) Burnouf, D., Koehl, P., and Fuchs, R. P. (1989) Single adduct mutagenesis: strong effect of the position of a single acetylaminofluorene adduct within a mutation hot spot. *Proc. Natl. Acad. Sci. U.S.A.* 86, 4147–4151.
(29) Broschard, T. H., Koffel-Schwartz, N., and Fuchs, R. P. (1999) Sequence-dependent modulation of frameshift mutagenesis at NarI-derived mutation hot spots. *J. Mol. Biol.* 288, 191–199.
(30) Jain, V., Hilton, B., Lin, B., Patnaik, S., Liang, F., Darian, E., Zou, Y., Mackrell, A. D. Jr., and Cho, B. P. (2013) Unusual sequence effects on nucleotide excision repair of arylamine lesions: DNA bending/distortion as a primary recognition factor. *Nucleic Acids Res.* 41, 869–880.
(31) Liang, F., and Cho, B. P. (2010) Enthrally-entropy contribution to carcinogen-induced DNA conformational heterogeneity. *Biochemistry* 49, 259–266.
(32) Jain, N., Meneni, S., Jain, V., and Cho, B. P. (2009) Influence of flanking sequence context on the conformational flexibility of aminofluorene-modified dG adduct in dA mismatch DNA duplexes. *Nucleic Acids Res.* 37, 1628–1637.
(33) Meneni, S., Liang, F., and Cho, B. P. (2007) Examination of the long-range effects of aminofluorene-induced conformational heterogeneity and its relevance to the mechanism of translational DNA synthesis. *J. Mol. Biol.* 366, 1387–1400.
(34) Miller, H., and Grollman, A. P. (1997) Kinetics of DNA polymerase I (Klenow fragment exo-+) activity on damaged DNA templates: Effect of proximal and distal template damage on DNA synthesis. *Biochemistry* 36, 15336–15342.
(35) Vaidyanathan, V. G., and Cho, B. P. (2012) Sequence effects on lesion syntheses of an aminofluorene-DNA adduct: Conformational, thermodynamic, and primer extension kinetic studies. *Biochemistry* 51, 1983–1995.
(36) Cho, B. (2010) Structure-Function Characteristics of Aromatic Amines-DNA Adducts, in *The Chemical Biology of DNA Damage*, pp 217–238, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim, Germany.
(37) Shapiro, R. U. G., Zawadzka, H., Broyde, S., and Hingerty, B. E. (1986) Conformation of d(CpG) modified by the carcinogen 4-aminobiphenyl: a combined experimental and theoretical analysis. *Biochemistry* 25, 2198–2205.
(38) Brown, K., Hingerty, B. E., Guenther, E. A., Krishnan, V. V., Broyde, S., Turteltaub, K. W., and Cosman, M. (2001) Solution structure of the 2-amino-1-methyl-6-phenylimidazo[4,5-b]pyridine C8-deoxyguanosine adduct in duplex DNA. *Proc. Natl. Acad. Sci. U.S.A.* 98, 8507–8512.
(39) Vaidyanathan, V. G., Xu, L., and Cho, B. P. (2012) Binary and ternary binding affinities between exonuclease-deficient Klenow fragment (Kf-exo-) and various arylamine DNA lesions characterized by surface plasmon resonance. *Chem. Rev. Toxicol.* 25, 1568–1570.
(40) Wang, M., Mahrenholz, A., and Lee, S. H. (2000) XPA stabilizes the XPA-DNA complex through protein-protein interaction. *Biochemistry* 39, 6433–6439.
(41) You, J. S., Wang, M., and Lee, S. H. (2003) Biochemical analysis of the damage recognition process in nucleotide excision repair. *J. Biol. Chem.* 278, 7476–7485.
(42) Lebbink, J. H., Fish, A., Reumer, A., Natraj, G., Winterwerp, H. H., and Siima, T. K. (2010) Magnesium coordination controls the molecular switch function of DNA mismatch repair protein MutS. *J. Biol. Chem.* 285, 13131–13141.
(43) Sedletska, Y., Culard, F., Midoux, P., and Malinge, J. M. (2013) Interaction studies of mutS and mutL with DNA containing the major cisplatin lesion and its mismatched counterpart under equilibrium and nonequilibrium conditions. *Biopolymers* 99, 636–647.