

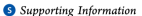


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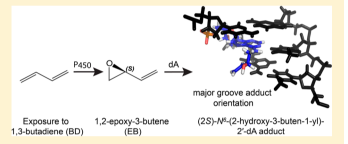
# Major Groove Orientation of the (2S)-N<sup>6</sup>-(2-Hydroxy-3-buten-1-yl)-2'deoxyadenosine DNA Adduct Induced by 1,2-Epoxy-3-butene

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ABSTRACT: 1,3-Butadiene (BD) is an environmental and occupational toxicant classified as a human carcinogen. It is oxidized by cytochrome P450 monooxygenases to 1,2-epoxy-3butene (EB), which alkylates DNA. BD exposures lead to large numbers of mutations at A:T base pairs even though alkylation of guanines is more prevalent, suggesting that one or more adenine adducts of BD play a role in BD-mediated genotoxicity. However, the etiology of BD-mediated genotoxicity at adenine remains poorly understood. EB alkylates the  $N^6$  exocyclic nitrogen of adenine to form  $N^6$ -(hydroxy-3-



buten-1-yl)-2'-dA ((2S)-N<sup>6</sup>-HB-dA) adducts (Tretyakova, N., Lin, Y., Sangaiah, R., Upton, P. B., and Swenberg, J. A. (1997) Carcinogenesis 18, 137–147). The structure of the (2S)-N<sup>6</sup>-HB-dA adduct has been determined in the 5'-d(C¹G²G³A⁴<u>C⁵Yʻ6</u>A²G<sup>8</sup>A<sup>9</sup>A¹0G¹¹¹)-3':5'-d(C¹²T¹³T¹⁴C¹⁵T¹G¹¹8T¹¹9 C²0C²¹G²²)-3' duplex [Y = (2S)-N<sup>6</sup>-HB-dA] containing codon 61 (underlined) of the human N-ras protooncogene, from NMR spectroscopy. The (2S)-N<sup>6</sup>-HB-dA adduct was positioned in the major groove, such that the butadiene moiety was oriented in the 3' direction. At the  $C_a$  carbon, the methylene protons of the modified nucleobase Y<sup>6</sup> faced the 5' direction, which placed the  $C_{\beta}$  carbon in the 3' direction. The  $C_{\beta}$  hydroxyl group faced toward the solvent, as did carbons  $C_{\gamma}$  and  $C_{\delta}$ . The  $C_{\beta}$  hydroxyl group did not form hydrogen bonds with either  $T^{16}$ O<sup>4</sup> or T<sup>17</sup> O<sup>4</sup>. The (2S)-N<sup>6</sup>-HB-dA nucleoside maintained the anti conformation about the glycosyl bond, and the modified base retained Watson–Crick base pairing with the complementary base ( $T^{17}$ ). The adduct perturbed stacking interactions at base pairs  $C^5:G^{18}$ ,  $Y^6:T^{17}$ , and  $A^7:T^{16}$  such that the  $Y^6$  base did not stack with its S' neighbor  $C^5$ , but it did with its 3' neighbor  $A^7$ . The complementary thymine  $T^{17}$  stacked well with both S' and  $S^{18}$  neighbors  $S^{18}$  and  $S^{18}$ . The presence of the ( $S^{18}$ )- $S^{18}$ -HB-dA resulted in a 5  $^{\circ}$ C reduction in the  $T_{\rm m}$  of the duplex, which is attributed to less favorable stacking interactions and adduct accommodation in the major groove.

# ■ INTRODUCTION

1,3-Butadiene (BD) is a genotoxic chemical strongly carcinogenic in laboratory mice $^{1-3}$  and to a lesser extent in rats.  $^4$  BD has been classified by the United States Environmental Protection Agency as "carcinogenic to humans by inhalation,"<sup>5</sup> and it has been also characterized as a known human carcinogen by the National Toxicology Program.<sup>6</sup> The International Agency for Cancer Research (IARC) lists BD as "carcinogenic to humans (Group 1)."<sup>7,8</sup> Accordingly, there has been interest in identifying human biomarkers of exposure to BD.<sup>9,10</sup> These exposures arise occupationally during the manufacture of styrene-butadiene rubber 11,12 and also environmentally since BD is found in automobile emissions 13 and in cigarette smoke. <sup>14</sup> Chronic exposures to BD may induce genotoxic effects <sup>15–17</sup> and have been associated with increased cancer risk. <sup>11,18–26</sup>

Albertini, Kirman, and co-workers have reviewed BD metabolism and genotoxicity. 27-29 It is of interest that BD exposures lead to large numbers of mutations at A:T base pairs, 30-34 even though alkylation of guanines by EB is more prevalent.<sup>35</sup> The number of A:T base pair substitutions has been reported to equal or exceed the number of mutations at G:C base pairs,<sup>34</sup> which implies that one or more adeninespecific lesions contribute significantly to butadiene-induced genotoxicity. <sup>32,34,36</sup> However, the etiology of adenine-specific mutations remains incompletely understood. 32,37,38 Electrophilic 1,2-epoxy-3-butenes (EB) are formed prevalently when BD is oxidized by cytochrome P450 enzymes (Scheme 1),<sup>39–41</sup> and alkylation products formed from the reactions of EB with adenine have been characterized. In each instance, a pair of regioisomeric 1-hydroxy-3-buten-2-yl and 2-hydroxy-3-buten-1yl and alkylation products is produced as a result of epoxide ring opening at either the internal or the terminal carbon atoms

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Scheme 1. Cytochrome P<sub>450</sub>-Mediated Oxidation of BD to EB, DEB, and EBD, Where EH Is Epoxide Hydrolase

of EB, respectively. Among the EB-dA products that have been identified in calf thymus DNA are N-1-(1-hydroxy-3-buten-2yl)-adenine, N-1-(2-hydroxy-3-buten-1-yl)-adenine, N-3-(1-hydroxy-3-buten-2-yl)-adenine, and N-3-(2-hydroxy-3-buten-1yl)-adenine. The N1-dA and N3-dA adducts occur at lower levels than do N7-dG adducts,<sup>39</sup> but they may be important for the ability of EB to induce mutations at A:T base pairs.<sup>30–3</sup> The N1-dA adducts are likely to be precursors of the regioisomeric  $N^6$ -(1-hydroxy-3-buten-2-yl)- 2'-dA adducts<sup>42</sup> and  $N^6$ -(2-hydroxy-3-buten-1-yl)-dA ( $N^6$ -HB-dA) adducts (Chart 1) through Dimroth rearrangement. 39,43-45 N<sup>6</sup>-HB-dA adducts have been detected in calf thymus DNA treated with EB in vitro. 42,46,47 N<sup>6</sup>-HB-dA adducts have also been detected in tissues of rodents exposed to BD by inhalation. 46,48 A second P450-catalyzed oxidation of EB leads to the more genotoxic diepoxybutane (DEB),<sup>7,12,30-32</sup> a bis-electrophile that forms DNA–DNA cross-links<sup>49–54</sup> and DNA–protein conjugates.<sup>50</sup> Thus, proximate electrophiles arising from BD metabolism include not only EB, and also DEB, and 1,2-dihydroxy-3,4epoxybutane (EBD).<sup>55-57</sup> Additionally, EBD is metabolized by cytochrome P450 to hydroxymethylvinylketones (HMVK). 58,59

The preparation of site-specific BD alkylation products in synthetic oligodeoxynucleotides provides a basis by which to probe the chemistry and biology of BD-derived electrophiles in DNA. The potential roles of both regio- and stereochemistry in modulating processing of these adducts are of interest, e.g., in light of the regio- and stereospecific processing of adducts arising from diol epoxides of various polycyclic aromatic hydrocarbons.<sup>60</sup> Harris and co-workers developed a post-oligomerization synthetic approach<sup>61,62</sup> in which oligodeoxynucleotides modified site-specifically with 6-chloropurine were reacted with regio- and stereospecific amino alcohol surrogates of specific BD-derived adducts.<sup>63</sup> Following this approach, oligodeoxynucleotides containing site-specific  $N^6$ -(2-hydroxy-3buten-1-yl)-dA (N<sup>6</sup>-HB-dA) adducts have been prepared by Quirk-Dorr et al.<sup>64</sup> The ras61 duplex 5'-d(CGGA<u>CAA</u>GAAG)-3':5'-d(CTTCTTGTCCG)-3' contains codons 60, 61 (underlined), and 62 of the human N-ras protooncogene. Feng and Stone 65 employed a restrained molecular dynamics and

Chart 1. (A) Structure of the (2S)-N<sup>6</sup>-HB-dA Adduct and (B) Sequences and Numbering of the Unmodified and Modified Duplexes<sup>a</sup>

 $^{\prime\prime}$  In A, H $_{\alpha}$  is the pro-R, and H $_{\alpha'}$  is the pro-S proton. In B, at the Y $^{6}$  position adenine has been replaced with the (2S)-N $^{6}$ -HB-dA adduct to form the modified duplex.

 $Y = (2S)-N^6-HB-dA$ 

simulated annealing approach to refine the structure of this duplex and concluded that it maintained a B-like DNA helix.

In the present work, we have generated DNA strands containing site- and stereospecific  $N^6$ -HB-dA adducts of BD. The structure of the (2S)- $N^6$ -HB-dA adduct<sup>64</sup> has been determined in the ras61 5′-d(C¹G²G³A⁴Cౖ⁵Yౖ⁶Aౖ²G³A⁴A°A¹0G¹¹)-3′:5′-d(C¹²T¹³T¹⁴C¹⁵T¹6T¹³G¹¹8T¹9 C²0C²¹G²²)-3′ duplex [Y = (2S)- $N^6$ -HB-dA] (Chart 1). The structure reveals that the (2S)- $N^6$ -HB-dA adduct is positioned in the major groove such that the butadiene moiety is oriented in the 3′ direction. The modified base Y<sup>6</sup> maintains Watson—Crick base pairing with the complementary base T¹7. The (2S)- $N^6$ -HB-dA adduct perturbs stacking interactions at base pairs C⁵:G¹³8, Y⁶:T¹²7, and A³:T¹⁶ resulting in a 5 °C reduction in the  $T_m$  of the duplex.

# MATERIALS AND METHODS

Syntheses and Characterization of Oligodeoxynucleotides. Unmodified oligodeoxynucleotides 5'-d(CGGACAAGAAG)-3' and 5'-d(CTTCTTGTCCG)-3' were synthesized by the Midland Reagent Company (Midland, TX) and purified by anion-exchange HPLC. The 5'-O-(4,4'-dimethoxytrityl)-3'-O-(2-cyanoethyl)-N,N-diisopropyl phosphoramidite of 6-chloropurine-2'-deoxyribonucleoside was purchased from ChemGenes Co. (Wilmington, MA). The 2'deoxyribonucleoside-3'-phosphoramidites and all other reagents necessary for automated DNA synthesis were purchased from Glen Research (Sterling, VA). Oligodeoxynucleotides were synthesized by solid phase methods using an ABI 394 DNA synthesizer (Life Technologies, Carlsbad, CA). All solvents and chemical reagents were obtained from commercial sources and used without further purification. The oligodeoxynucleotide 5'-d(CGGACYAGAAG)-3' containing the (2S)-N<sup>6</sup>-HB-dA adduct was synthesized by coupling 6-chloropurine containing DNA with 1-aminobut-3-en-2-ol.<sup>64</sup> Briefly, an 11-mer oligodeoxynucleotide containing a 6-chloropurine at position Y<sup>6</sup> (210 nmol) was coupled with 1-aminobut-3-en-2-ol

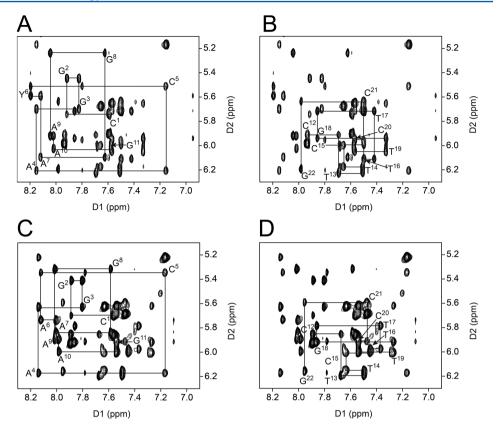


Figure 1. NOESY spectra for the modified and unmodified duplexes, showing sequential NOEs between the base aromatic and anomeric protons. (A) The (2S)- $N^6$ -HB-dA modified duplex, showing nucleotides  $C^1$  to  $G^{11}$ . The resonance at 6.95 ppm (D1) belongs to the  $Y^6$  H2 proton. (B) The (2S)- $N^6$ -HB-dA modified duplex, showing nucleotides  $C^{12}$  to  $G^{22}$ . (C) The unmodified duplex, showing nucleotides  $C^1$  to  $G^{11}$ . The resonance at 7.10 ppm (D1) belongs to the  $A^6$  H2 proton. (D) The unmodified duplex, showing nucleotides  $C^{12}$  to  $G^{22}$ . Spectra were obtained at 800 and 600 MHz for modified and unmodified duplexes, respectively, both with a mixing time of 250 ms at 15 °C.

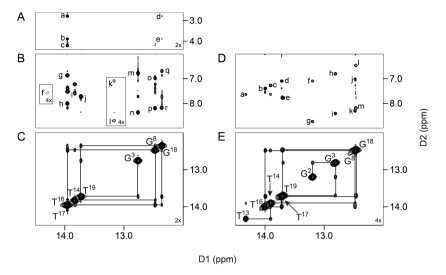
(7.44 mg), in the presence of DIPEA (210  $\mu$ L) in DMSO (700  $\mu$ L) for 16 h at 60 °C. The oligodeoxynucleotides were purified using semipreparative reverse-phase HPLC (YMC, Kyoto, Japan, Phenyl-Hexyl,  $5 \mu m$ , 250 mm  $\times 10.0$  mm) equilibrated with 0.1 M ammonium formate (pH 7.0) using an acetonitrile gradient. The oligodeoxynucleotides were desalted using Sephadex G-25 and lyophilized. The adducted oligodeoxynucleotides were characterized by capillary HPLC-ESI-MS. Sequence and site-specificity were confirmed by MALDI-TOF-MS of partial exonuclease digests. The unmodified oligodeoxynucleotides were characterized by capillary HPLC-ESI-MS. The concentrations of single-stranded oligodeoxynucleotides were determined by UV absorbance at 260 nm using extinction coefficients of 118,300 L M<sup>-1</sup>cm<sup>-1</sup> for 5'-d(CGGACYAGAAG)-3' and 5'd(CGGACAAGAAG)-3' and 90,800 L M<sup>-1</sup>cm<sup>-1</sup> for the complementary strand 5'-d(CTTCTTGTCCG)-3'.<sup>66</sup> Equimolar quantities of the complementary strands were combined and annealed by heating to 80  $^{\circ}\text{C}$  for 15 min and then slowly cooled to room temperature to form a duplex.

**DNA Melting Studies.** Absorption vs temperature profiles for each duplex were measured using a Varian Cary 100 Bio spectrophotometer (Varian Associates, Palo Alto, CA). The concentrations of the duplexes were 1.24  $\mu$ M. Samples were prepared in a solution of 10 mM NaH<sub>2</sub>PO<sub>4</sub> and 50  $\mu$ M Na<sub>2</sub>EDTA containing 0.1 M NaCl (pH 7.0). The temperature was increased from 5 to 90 °C for each duplex at a rate of 1.0 °C/min. The UV absorbance was monitored at 260 nm. The  $T_{\rm m}$  values were determined by taking the first derivatives of the melting curves and shape analyses.

NMR Spectroscopy. The DNA duplex containing the  $Y^6$  adduct was prepared at 0.51 mM concentration in 0.1 M NaCl and 50  $\mu$ M Na<sub>2</sub>EDTA in the presence of 10 mM NaH<sub>2</sub>PO<sub>4</sub> (pH 7.0). To observe nonexchangeable protons, the duplex was exchanged with D<sub>2</sub>O and dissolved in 99.96% D<sub>2</sub>O. To observe exchangeable protons, the

duplex was dissolved in 9:1 H<sub>2</sub>O/D<sub>2</sub>O. <sup>1</sup>H NMR spectra were recorded using 800 and 600 MHz spectrometers equipped with cryogenic probes (Bruker Biospin, Billerica, MA). Chemical shifts were referenced to the chemical shift of water at the corresponding temperature, with respect to 4,4-dimethyl-4-silapentane-1-sulfonic acid (DSS). Data were processed using TOPSPIN (Bruker Biospin Inc., Billerica, MA). NOESY $^{67,68}$  and DQF-COSY $^{69}$  spectra in D<sub>2</sub>O were collected at 15 °C at 800 MHz; NOESY experiments were conducted at a mixing time of 60 and 250 ms. These experiments were performed with a relaxation delay of 2.0 s. The NOESY spectrum in 9:1 H<sub>2</sub>O/ D<sub>2</sub>O was collected at 10 °C at 600 MHz for the modified and unmodified duplexes with a 250 ms mixing time. NMR experiments as a function of temperature in 9:1 H<sub>2</sub>O/D<sub>2</sub>O solution were collected at 5, 10, 15, 20, 25, and 30 °C at 600 MHz. These experiments were performed with a relaxation delay of 1.5 s. Water suppression was performed using the WATERGATE pulse sequence.7 peaks were assigned using the program SPARKY.<sup>7</sup>

NMR Distance Restraints. NOESY spectra cross-peak volumes were measured by volume integrations of the NOESY spectra using SPARKY. They were divided into five classes based on confidence in the integrations. The intrinsic integration error was assigned to be one-half the volume of the cross-peak of lowest intensity. The overlapping of cross-peaks, spectroscopic line broadening of cross-peaks, which was particularly an issue for integrations of cross-peaks involving exchangeable protons, and the potential for spin diffusion provided additional sources of integration errors. The class 1 cross-peak volumes were derived from well-resolved strong nonoverlapping cross-peaks and were assigned 10% error. The class 2 cross-peak volumes were derived from strong but slightly overlapped cross-peaks and were assigned 20% error. The class 3 cross-peak volumes were derived from strong but medially overlapped cross-peaks and were assigned 30% error. Classes 4 and 5 of the cross-peak volumes were derived from



**Figure 2.** NOESY spectra for the (2*S*)-*N*<sup>6</sup>-HB-dA modified (A−C) and unmodified (D−E) duplexes, showing NOEs between the base imino protons and the amino protons. (A) Interstrand NOEs between the Y<sup>6</sup> adduct and the T<sup>17</sup> base. The cross-peaks are assigned as a, Y<sup>6</sup> H<sub>α</sub>′ → T<sup>17</sup> N3H; b, Y<sup>6</sup> H<sub>α</sub> → T<sup>17</sup> N3H; c, Y<sup>6</sup> H<sub>β</sub> → T<sup>17</sup> N3H; d, Y<sup>6</sup> H<sub>α</sub>′ → G<sup>18</sup> N1H; and e, Y<sup>6</sup> H<sub>α</sub> → G<sup>18</sup> N1H. (B) Interstrand NOEs between complementary bases. The cross-peaks are assigned as f, A<sup>10</sup> H2 → T<sup>13</sup> N3H; g, Y<sup>6</sup> H2 → T<sup>17</sup> N3H; h, A<sup>7</sup> H2 → T<sup>16</sup> N3H; i, A<sup>9</sup> H2 → T<sup>14</sup> N3H; j, A<sup>4</sup> H2 → T<sup>19</sup> N3H; k, C<sup>21</sup> N<sup>4</sup>H1 → G<sup>2</sup> N1H; l, C<sup>21</sup> N<sup>4</sup>H2 → G<sup>2</sup> N1H; m, C<sup>20</sup> N<sup>4</sup>H1 → G<sup>3</sup> N1H; n, C<sup>20</sup> N<sup>4</sup>H2 → G<sup>3</sup> N1H; o, C<sup>15</sup> N<sup>4</sup>H1 → G<sup>8</sup> N1H; p, C<sup>15</sup> N<sup>4</sup>H2 → G<sup>8</sup> N1H; q, C<sup>5</sup> N<sup>4</sup>H1 → G<sup>18</sup> N1H; and r, C<sup>5</sup> N<sup>4</sup>H2 → G<sup>18</sup> N1H. (C) NOE connectivity for the imino protons for the base pairs G<sup>2</sup>:C<sup>21</sup>, G<sup>3</sup>:C<sup>20</sup>, A<sup>4</sup>:T<sup>19</sup>, C<sup>5</sup>:G<sup>18</sup>, Y<sup>6</sup>:T<sup>17</sup>, A<sup>7</sup>:T<sup>16</sup>, G<sup>8</sup>:C<sup>15</sup>, and A<sup>9</sup>:T<sup>14</sup>. The cross-peaks are T<sup>14</sup> N3H → G<sup>8</sup> N1H, G<sup>8</sup> N1H → T<sup>16</sup> N3H, T<sup>16</sup> N3H → T<sup>17</sup> N3H, T<sup>17</sup> N3H → G<sup>18</sup> N1H, G<sup>18</sup> N1H → T<sup>19</sup> N3H, and T<sup>19</sup> N3H → G<sup>3</sup> N1H. The spectrum was obtained at 600 MHz with a mixing time of 250 ms at 10 °C. (D) Interstrand NOEs between complementary bases. The cross-peaks are assigned as a, A<sup>10</sup> H2 → T<sup>13</sup> N3H; b, A<sup>7</sup> H2 → T<sup>16</sup> N3H; c, A<sup>9</sup> H2 → T<sup>14</sup> N3H; d, A<sup>6</sup> H2 → T<sup>17</sup> N3H; e, A<sup>4</sup> H2 → T<sup>19</sup> N3H; f, C<sup>21</sup> N<sup>4</sup>H1 → G<sup>2</sup> N1H; g, C<sup>21</sup> N<sup>4</sup>H2 → G<sup>2</sup> N1H; h, C<sup>20</sup> N<sup>4</sup>H1 → G<sup>3</sup> N1H; i, C<sup>20</sup> N<sup>4</sup>H2 → G<sup>3</sup> N1H; in C<sup>3</sup> N<sup>4</sup>H2 → G<sup>3</sup> N1H; in C<sup>3</sup> N<sup>4</sup>H2 → G<sup>3</sup> N1H; in C<sup>3</sup> N<sup>4</sup>H1 → G<sup>8</sup> N1H; in C<sup>3</sup> N<sup>4</sup>H1 → G<sup>8</sup> N1H; in C<sup>3</sup> N<sup>4</sup>H2 → G<sup>8</sup> N1H; in C<sup>3</sup> N<sup>4</sup>H2 → G<sup>8</sup> N1H; in C<sup>3</sup> N<sup>4</sup>H2 → G<sup>8</sup> N1H; in

low S/N, broadened, and/or highly overlapped cross-peaks, such as those in regions close to the water resonance or to the diagonal line of the spectrum. These were assigned 40% and 50% errors, respectively. An unmodified B-type DNA duplex was constructed using the program INSIGHT II (Accelrys Inc., San Diego, CA). The adenine at position  $A^6$  was replaced by the (2S)- $N^6$ -HB-dA adduct. Partial charges for the (2S)-N<sup>6</sup>-HB-dA base were calculated with the B3LYP/6-31G\* basis set in Gaussian.<sup>72</sup> These were employed in the parameter files prepared in the program XLEAP.<sup>73</sup> Table S1 in the Supporting Information provides the parametrization for the modified  $(2S)-N^6$ HB-dA nucleotide. The modified duplex was subjected to 1000 cycles of potential energy minimization. The integrated cross-peak intensities, separated into five classes, as described above, were combined by the program MARDIGRAS with volumes calculated from complete relaxation matrix analysis of the starting model to generate a hybrid intensity matrix.<sup>74,75</sup> The program MARDIGRAS was employed to refine the hybrid intensity matrix<sup>76</sup> and calculate interproton distance vectors, with upper and lower bounds to each distance vector. For methyl protons, the JUMP 3<sup>77</sup> model was employed. These calculations were performed at 2, 3, and 4 ns isotropic correlation

Restrained Molecular Dynamics Calculations. The interproton distance vectors calculated by MARDIGRAS were used to provide distance restraints used in the restrained molecular dynamics (rMD) calculations. The widths of the distance restraint potential energy wells corresponded to the upper and lower bounds on the inter proton distance vectors as calculated by MARDIGRAS. Additional phosphodiester backbone restraints and deoxyribose pseudorotation restraints were employed, derived from B-DNA. For the modified nucleotide Y<sup>6</sup>, the square potential energy wells for phosphodiester restraints were assigned as  $\pm 120^{\circ}$ . For unmodified nucleotides, the widths of the square potential energy wells for the phosphodiester restraints were assigned as  $\pm 60^{\circ}$ . The pseudorotation restraints were not employed for the modified nucleotide Y<sup>6</sup> or for the terminal bases C<sup>1</sup>, G<sup>11</sup>, G<sup>12</sup>,

and G<sup>22</sup>. Watson—Crick base pair restraints were employed for all base pairs.

The simulated annealing protocol<sup>79</sup> used for the rMD calculations utilized the program AMBER<sup>80</sup> and the parm99 force field.<sup>81</sup> Force constants of 32 kcal  $\text{mol}^{-1}$  Å<sup>-2</sup> were applied for all restraints. The generalized Born model<sup>82</sup> was used for solvation. The salt concentration was 0.1 M. The molecule was coupled to the bath temperature to establish the temperature during calculations. Initially, calculations were performed for 20 ps (20,000 steps). For the first 1,000 steps, the system was heated from 0 to 600 K with a coupling of 0.5 ps, followed by 1,000 steps at 600 K, followed by 16,000 steps in which the system was cooled to 100 K with a coupling of 4 ps. In the last 2,000 steps, additional cooling was applied from 100 to 0 K with a coupling of 1 ps. Subsequently, a 100,000 step calculation was performed over 100 ps. For the first 5,000 steps, the system was heated from 0 to 600 K with a coupling of 0.5 ps, followed by 5,000 steps at 600 K, followed by 80,000 during which the system was cooled to 100 K with a coupling of 4 ps, followed by additional cooling for the last 10,000 steps with a coupling of 1 ps. Structure coordinates were saved after each cycle. Complete relaxation matrix analysis (CORMA)<sup>74,75</sup> was used to compare intensities calculated from these emergent structures with the experimentally measured distances. Nine structures were chosen, based on the lowest deviations from the experimental distance and dihedral restraints. These were subjected to potential energy minimization and used to obtain an average refined structure.

**Data Deposition.** The structure factors and coordinates were deposited in the Protein Data Bank (www.rcsb.org). The PDB ID code for the duplex containing the  $(2S)-N^6$ -HB-dA adduct is 2MNX.

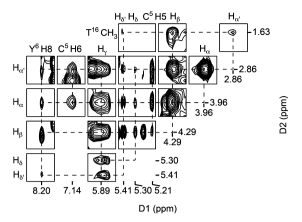
# RESULTS

**NMR Spectroscopy.** Base Proton Assignments. Figure 1 shows the regions of the NOESY spectra including the base aromatic proton resonances and deoxyribose H1' proton

resonances<sup>84,85</sup> for the modified duplex (panels A and B), in comparison with the corresponding unmodified duplex (panels C and D). The presence of the  $(2S)-N^6$ -HB-dA adduct induced small changes in the sequential pattern of NOEs between the aromatic base protons and the anomeric protons for either strand of the duplex, as compared to the unmodified duplex. For the modified strand, the  $C^5 H6 \rightarrow C^5 H1'$ ,  $C^5 H1' \rightarrow Y^6$ H8,  $Y^6$  H8  $\rightarrow$   $Y^6$  H1', and  $Y^6$  H1'  $\rightarrow$  A<sup>7</sup> H8 NOEs were of similar intensities as compared to the corresponding NOEs arising from distal nucleotides. The intensity of the Y<sup>6</sup> H8  $\rightarrow$  Y<sup>6</sup> H1' NOE was indicative of minimal change in the conformation of the glycosyl torsion angle at the site of modification. For the complementary strand, the  $T^{16}$  H6  $\rightarrow$   $T^{16}$ H1',  $T^{16}$  H1'  $\rightarrow$   $T^{17}$  H6,  $T^{17}$  H6  $\rightarrow$   $T^{17}$  H1', and  $T^{17}$  H1'  $\rightarrow$ G18 H8 NOEs were of similar intensities compared to the corresponding NOEs from nucleotides distal to the adduct. With the deoxyribose H1' assignments in hand, the remainder of the deoxyribose protons were assigned from a combination of NOESY and COSY data. The adenine H2 proton resonances were assigned based upon NOEs to the thymine N1H imino proton resonances of the respective A:T base pairs. The assignments of the nonexchangeable DNA protons are summarized in Table S2 of the Supporting Information.

Imino and Amino Proton Assignments. Figure 2 shows the regions of the NOESY spectra yielding the assignments of the Watson-Crick hydrogen bonded guanine and thymine imino resonances, and the adenine  $N^6$  and cytosine  $N^4$  exocyclic amino resonances, for the modified duplex in comparison with the unmodified duplex. In the NOESY spectrum of the modified duplex collected at 10 °C, the T<sup>16</sup> and T<sup>17</sup> N3H imino resonances overlapped. For the modified duplex, the sequential pattern of cross-peaks between imino protons<sup>86</sup> was observed for base pairs  $G^3:C^{20} \to A^4:T^{19} \to C^5:G^{18} \to Y^6:T^{17} \to A^7:T^{16} \to G^8:C^{15} \to A^9:T^{14}$  (Figure 2, panel C). The NOEs between the base imino and amino protons for G:C base pairs showed cross-peaks for all base pairs, with the exceptions of the terminal base pairs  $C^1:G^{22}$  and  $G^{11}:C^{12}$  (Figure 2, panel B). These cross-peaks for the G<sup>2</sup>:C<sup>21</sup> base pair were weak (Figure 2, panel B, cross-peaks k and l, plotted at 4× the contour level). The NOEs between the base imino and adenine H2 protons for T:A base pairs showed cross-peaks for all base pairs. The crosspeak between T13 N3H and A10 H2 was of low intensity (Figure 2, Panel B, cross-peak a, plotted at 4× the contour level). The Y<sup>6</sup> H2  $\rightarrow$  T<sup>17</sup> N3H cross-peak was of similar intensity compared to the other cross-peaks (g, Figure 2, panel B). The chemical shifts of the imino and amino protons are provided in Table S3 of the Supporting Information.

(2S)-N<sup>6</sup>-HB-dA Proton Assignments. Six resonances were observed between 2.8 and 5.9 ppm (Figure 3). The H<sub>v</sub> resonance was identified at 5.89 ppm. The  $H_{\delta}$  proton, identified at 5.30 ppm, and the  $H_{\delta'}$  proton, identified at 5.41 ppm, were assigned from their coupling constants to the H<sub>v</sub> proton. The  $H_{\delta'}$  proton exhibited the larger coupling constant to the  $H_{\nu}$ proton (Figure 3). The diastereotopic  $H_{\alpha}$  and  $H_{\alpha'}$  resonances of the methylene group were assigned based on the intensities of NOE cross-peaks to the  $H_{\beta}$  proton and to the Y<sup>6</sup> H8 proton. The  $H_{\alpha}$  resonance was located at 3.96 ppm, while the  $H_{\alpha'}$ resonance was located at 2.86 ppm. The  $H_{\beta}$  resonance was located at 4.29 ppm. The  $H_{\alpha}$  proton gave a more intense NOE cross-peak to  $H_{\beta}$  than did  $H_{\alpha'}$ . Likewise, the  $H_{\alpha'}$  proton gave a more intense NOE cross-peak to the Y6 H8 proton than did  $H_{\alpha}$ . Both the  $H_{\alpha}$  and  $H_{\alpha'}$  protons exhibited strong NOEs to the  $C^5$  H5 and H6 protons. The H<sub> $\beta$ </sub> proton exhibited a strong NOE



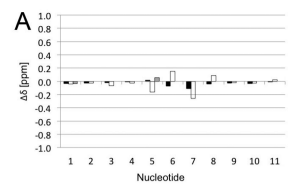
**Figure 3.** Expanded plot of the NOESY spectrum of the (2S)- $N^6$ -HB-dA modified duplex, showing assignments of the adduct protons and cross-peaks from the adduct protons to neighbor base protons. The chemical shifts for each proton are Y<sup>6</sup> H8, 8.20 ppm; C<sup>5</sup> H6, 7.14 ppm; Y<sup>6</sup> H $_{p}$ , 5.89 ppm; Y<sup>6</sup> H $_{\delta\prime}$ , 5.41 ppm; Y<sup>6</sup> H $_{\delta\prime}$ , 5.30 ppm; C<sup>5</sup> H5, 5.21 ppm; Y<sup>6</sup> H $_{\beta\prime}$ , 4.29 ppm; Y<sup>6</sup> H $_{\alpha\prime}$ , 3.96 ppm; Y<sup>6</sup> H $_{\alpha\prime}$ , 2.86 ppm; and T<sup>16</sup> CH $_{3}$ , 1.63 ppm. The dashed lines show the NOE connectivity for each proton. Cross-peak Y<sup>6</sup> H8–Y<sup>6</sup> H $_{\beta\prime}$  is overlapped with Y<sup>6</sup> H8–Y<sup>6</sup> H4′. The spectrum was obtained at 800 MHz, with a mixing time of 250 ms and at 15 °C.

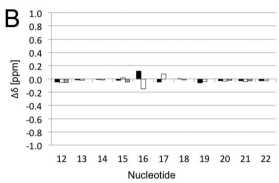
to the  $T^{16}$  CH<sub>3</sub> protons, located in the 3'-neighbor  $A^7:T^{16}$  base pair. Weak interstrand NOEs were observed between  $H_{\alpha}$ ,  $H_{\alpha'}$ , and  $H_{\beta}$  and the  $T^{17}$  N3H imino proton, as well as  $H_{\alpha}$   $H_{\alpha'}$ , and the  $G^{18}$  N1H imino proton (Figure 2, panel A).

Chemical Shift Perturbations. A number of chemical shift perturbations were observed in the modified strand at the lesion site as compared to the unmodified duplex (Figures 1 and 4). The C<sup>5</sup> H1' resonance shifted 0.16 ppm downfield, the Y<sup>6</sup> H8 resonance shifted less than 0.1 ppm downfield, the Y<sup>6</sup> H2 resonance shifted 0.15 ppm upfield, and the A<sup>7</sup> H1' and H8 resonance shifted 0.25 and 0.1 ppm downfield. The Y<sup>6</sup> H1' resonance shifted 0.15 ppm upfield. For the complementary strand of DNA, the largest chemical shift changes were observed for the T<sup>16</sup> H1' resonance, which shifted 0.15 ppm downfield, and for the T<sup>16</sup> H6 resonance, which shifted 0.12 ppm upfield. In the imino proton region of the spectrum, the greatest chemical shift perturbation was observed for the T<sup>17</sup> N3H resonance, which shifted 0.24 ppm downfield as compared to the unmodified duplex (Figure 2).

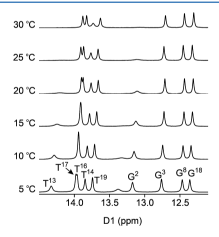
**Thermal Melting Studies.** The unfolding of the duplexes was examined by temperature-dependent UV spectroscopy monitored at 260 nm. Incorporation of the (2S)- $N^6$ -HB-dA adduct resulted in a 5 °C reduction in the  $T_{\rm m}$  value of the duplex. At the concentration of 1.24  $\mu$ M in 0.1 M NaCl at pH 7, the  $T_{\rm m}$  for the unmodified duplex was 44 °C, whereas the  $T_{\rm m}$  for the modified duplex was 39 °C. In  $^1$ H NMR experiments collected as a function of temperature from 5 to 30 °C, the T $^{16}$  and T $^{17}$  N3H and G $^{18}$  N1H imino proton resonances remained sharp as compared to the other imino proton resonances as temperature was increased, which suggested that over this range of temperatures, the presence of the modified (2S)- $N^6$ -HB-dA base did not increase the rates of exchange with solvent for the C $^5$ :G $^{18}$ , Y $^6$ :T $^{17}$ , and A $^7$ :T $^{16}$  base pairs (Figure 5).

**Structural Refinement.** A total of 284 distance restraints obtained from the analyses of the NOESY spectra of nonexchangeable protons were used for restrained molecular dynamics (rMD) calculations. Of these, 126 were internucleotide restraints, and 158 were intranucleotide





**Figure 4.** Chemical shift perturbations for the (2S)- $N^6$ -HB-dA modified duplex compared to those of the unmodified duplex. (A) Strands 1–11, (B) strands 12–22 for aromatic H6/H8 (shown in black), cytosine H5 (gray), and H1′ (white) protons, where  $\Delta\delta$  [ppm] =  $\delta_{\rm Unmodified}$  [ppm] -  $\delta_{\rm Modified}$  [ppm].



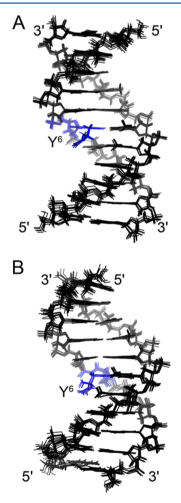
**Figure 5.** NMR spectra showing the imino proton resonances for the (2S)- $N^6$ -HB-dA duplex, as a function of temperature. The individual nucleotides are identified by superscripts. The spectra were obtained at 600 MHz, at temperatures 5, 10, 15, 20, 25, and 30 °C.

restraints. As the NMR data were consistent with a right-handed helical DNA duplex similar to that of the canonical B-form DNA,<sup>78</sup> a total of 90 backbone torsion angles, 45 hydrogen bondings, and 17 deoxyribose pseudorotations were included as empirical restraints in the rMD calculations. Table 1 summarizes the restraints that were used and the refinement statistics.

The rMD calculations employed a simulated annealing protocol. Nine structures emergent from the calculations were subjected to potential energy minimization. Figure 6 shows these nine superimposed structures. Table 2 summarizes the structural statistics. A satisfactory convergence was

Table 1. NMR Restraints Used for the rMD Structural Refinement of the (2S)- $N^6$ -HB-dA Modified Duplex and the Refinement Statistics

NMR restraints	
NOE restraints	
internucleotide	126
intranucleotide	158
total	284
backbone torsion angle restraints	90
hydrogen bonding distance restraints	45
deoxyribose pseudorotation restraints	17
total number of restraints	436
refinement statistics	
number of distance restraint violations > 0.025 Å	7
number of torsion restraint violations	10
total distance penalty/maximum penalty [kcal mol-1]	0.776/0.289
total torsion penalty/maximum penalty [kcal mol <sup>-1</sup> ]	0.490/0.159
r.m.s. distances (Å)	0.012
r.m.s. angles (deg)	2.355
distance restraint force constant [kcal mol <sup>-1</sup> Å <sup>-2</sup> ]	32
torsion restraint force constant [kcal $mol^{-1} deg^{-2}$ ]	32



**Figure 6.** Superpositions of nine structures obtained from a series of rMD calculations for the (2S)- $N^6$ -HB-dA modified duplex. The modified base  $Y^6$  is shown in blue. (A) View from the major groove. (B) Side view.

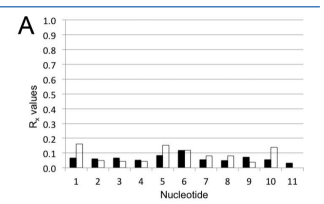
observed, with a maximum pairwise rmsd between the nine structures of 0.53 Å. These nine structures were averaged, and

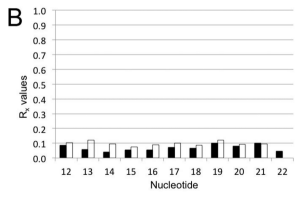
Table 2. Structural Statistics for the  $(2S)-N^6$ -HB-dA Modified Duplex

average structure (obtained from 9 structures)	
RMS pairwise difference between structures [Å]	0.53
RMS difference from average structure [Å]	0.35
CORMA analysis for average structure <sup>a</sup>	
$R_1^{xb}$	
intranucleotide	0.074
internucleotide	0.095
total	0.082
average error <sup>c</sup>	0.018

<sup>a</sup>The mixing time was 250 ms.  ${}^bR_1{}^x$  is the sixth root R factor:  $\Sigma[((Io)_i^{1/6}) - ((Ic)_i^{1/6})/\Sigma((Io)_i^{1/6})]$ . <sup>c</sup>Average error:  $\Sigma(I_c - I_o)/n$ , where  $I_c$  are NOE intensities calculated from refined structure,  $I_o$  are experimental NOE intensities.

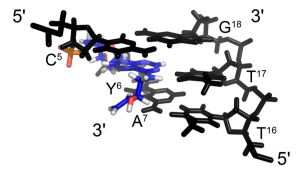
the resulting average structure was subjected to complete relaxation matrix analysis. <sup>74</sup> The results are shown in Figure 7. The sixth root residuals ( $R_1^x$  values) remained consistently below 15%, for both intranucleotide NOEs and internucleotide NOEs.





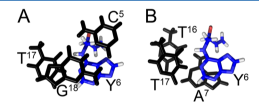
**Figure 7.** Complete relaxation matrix analysis (CORMA) results for internucleotide (shown in black) and intranucleotide (shown in white) NOEs for the (2S)- $N^6$ -HB-dA modified duplex, A (strands 1-11 and B (strands 12-22). $R_1^x$  is the sixth root R factor:  $\Sigma[((Io)_i^{1/6}) - ((Ic)_i^{1/6})/\Sigma((Io)_i^{1/6})]$ , where Ic are NOE intensities calculated from the refined structure, Io are experimental NOE intensities.

Structure of the (2S)- $N^6$ -HB-dA Modified Duplex. Figure 8 shows the average structure of the (2S)- $N^6$ -HB-dA modified duplex in the region of the  $C^5$ : $G^{18}$ ,  $Y^6$ : $T^{17}$ , and  $A^7$ : $T^{16}$  base pairs. The view is from the major groove. The (2S)- $N^6$ -HB-dA adduct was positioned in the major groove such that the butadiene moiety oriented in the 3' direction. The  $C_\alpha$  carbon



**Figure 8.** Average structure of the (2S)- $N^6$ -HB-dA modified duplex in the region of the  $C^5$ : $G^{18}$ ,  $Y^6$ : $T^{17}$ , and  $A^7$ : $T^{16}$  base pairs. The modified nucleotide  $Y^6$  is shown in blue.

remained in plane with the modified nucleobase  $Y^6$ , with the  $H_{\alpha}$  and  $H_{\alpha}'$  methylene protons facing the 5' direction, which placed the  $C_{\beta}$  carbon in the 3' direction. The  $C_{\beta}$  hydroxyl group faced the solvent, as did carbons  $C_{\gamma}$  and  $C_{\delta}$ . There was no indication of hydrogen bond formation between the hydroxyl group at  $C_{\beta}$  with either  $T^{16}$   $O^4$  or  $T^{17}$   $O^4$ . The (2S)- $N^6$ -HB-dA nucleoside maintained the *anti* conformation about the glycosyl bond. Figure 9 shows the base pairing



**Figure 9.** Stacking interactions for the (2S)- $N^6$ -HB-dA modified duplex. (A) Stacking of the  $C^5$ : $G^{18}$  base pair (black) above  $Y^6$  (blue) and  $T^{17}$  (black). (B) Stacking of the  $Y^6$ : $T^{17}$  pair (in blue and black, respectively) above base pair  $A^7$ : $T^{16}$  (black).

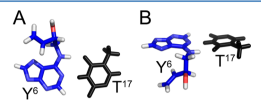
interactions at the lesion site. The modified base  $Y^6$  maintained Watson–Crick base pairing with the complementary base  $T^{17}$ . The (2S)- $N^6$ -HB-dA adduct perturbed stacking interactions at base pairs  $C^5$ : $G^{18}$ ,  $Y^6$ : $T^{17}$ , and  $A^7$ : $T^{16}$  (Figure 9). Thus, the  $Y^6$  base did not stack with its S' neighbor  $C^5$ , but it did with its S' neighbor  $S^7$ . The complementary thymine  $S^{17}$  stacked well with both S' and  $S^7$  neighbors  $S^{16}$  and  $S^{18}$ .

## DISCUSSION

 $N^6$ -HB-dA adducts induced by epoxybutene (EB, Chart 1) are of significant interest due to the ability of the BD lead to induce large numbers of mutations at A:T base pairs in DNA,  $^{30-34}$  even though alkylation of guanines by EB is more prevalent. The origins of these adenine-specific mutations are not well understood,  $^{32,37,38}$  but their presence implies that one or more adenine-specific lesions contribute to BD-induced genotoxicity. The  $N^6$ -HB-dA adducts can be formed via direct alkylation by EB of the  $N^6$ -dA position or they can arise via Dimroth rearrangement of the corresponding N1-dA adducts. The  $N^6$ -HB-dA adducts potentially interfere with Watson—Crick base pairing due to the presence of the hydroxybutenyl moiety at the  $N^6$  nitrogen, which is normally involved in hydrogen bonding with the complementary adenine. In the present study, we have prepared a DNA duplex containing the site- and stereospecific (2S)- $N^6$ -HB-dA adduct

opposite dT, and we have examined its effect on DNA structure and stability.

**Major Groove Orientation of the (2S)-** $N^6$ -**HB-dA Adduct.** The (2S)- $N^6$ -HB-dA adduct is positioned in the major groove of DNA. The *anti* conformation about the glycosyl bond of  $N^6$ -HB-dA is confirmed by NOE data showing that the intensity of the NOE between  $Y^6$  H8 and  $Y^6$  H1' is small as compared to the NOE between the cytosine H5 and H6 protons (Figure 1). The (2S)- $N^6$ -HB-adenine base ( $Y^6$ ) maintains Watson—Crick base pairing with the complementary  $T^{17}$  base (Figures 8 and 10), which is confirmed by the NOE



**Figure 10.**  $Y^6:T^{17}$  base pair in the (2*S*)- $N^6$ -HB-dA duplex. (A) View from the top. (B) View from the major groove.  $Y^6$  forms a Watson–Crick base pair with the complementary  $T^{17}$  base.

cross-peak between Y6 H2 and T17 N3H (cross-peak g, Figure 2B). Furthermore, there is no break in the pattern of sequential NOEs between the base paired imino protons (Figure 2C), indicating that the Y<sup>6</sup> base remains stacked into the DNA helix. The T<sup>16</sup> and T<sup>17</sup> N3H and G<sup>18</sup> N1H imino proton resonances remain sharp as compared to the other imino proton resonances as temperature is increased (Figure 5), suggesting that the (2S)-N<sup>6</sup>-HB-dA lesion does not increase the rates of exchange with solvent for the C5:G18, Y6:T17, and A7:T16 base pair imino protons. The lower intensities of the NOE crosspeaks for base pairs G<sup>2</sup>:C<sup>21</sup> and A<sup>10</sup>:T<sup>13</sup> as compared to other base pair imino proton cross-peaks (Figure 2B) are attributed to their increased rate of exchange with water. Furthermore, the (2S)-N<sup>6</sup>-HB-dA adduct perturbs the stacking of base pair A<sup>7</sup>:T<sup>16</sup> with the neighboring base pairs C<sup>5</sup>:G<sup>18</sup> and Y<sup>6</sup>:T<sup>17</sup> (Figure 9). The Y<sup>6</sup> base does not stack well with its 5' neighbor C<sup>5</sup> but does stack with its 3' neighbor A<sup>7</sup>, which is in the agreement with chemical shift changes for the Y<sup>6</sup> H8 and H2 protons. Y<sup>6</sup> H8 is shifted downfield by 0.1 ppm and Y<sup>6</sup> H2 is shifted upfield by 0.15 ppm (Figure 1). The complementary thymine, T17 stacks well with both 5' and 3' neighbors T16 and  $G^{18}$ . The 1.1 ppm chemical shift difference between the  $H_{\alpha}$  and  $H_{\alpha'}$  resonances (Figure 3) is attributed to a stacking interaction with the  $C^5$  base, in which  $H_a$  is less shielded compared to that of  $H_{\alpha'}$  (Figure 9). Collectively, these structural perturbations may account for the 5 °C reduction in the  $T_{\rm m}$  of the duplex in 0.1 M NaCl at pH 7.

Structural Comparisons to Other  $N^6$ -dA Adducts. The structure of the (2S)- $N^6$ -HB-dA adduct shows significant similarities to those of  $N^6$ -(2,3,4-trihydroxybutyl)-2'-dA  $(N^6$ -THB-dA) adducts 1 (Chart 2) arising from another epoxide metabolite of BD, 1,2-dihydroxy-3,4-epoxybutane (EBD) (Scheme 1). $^{55,56,88}$  Like (2S)- $N^6$ -HB-dA, the (2R,3R)-1 and (2S,3S)-1  $N^6$ -THB-dA adducts  $^{37,89}$  are accommodated in the major groove and maintain Watson—Crick base pairing. $^{90,91}$  This is consistent with the facile bypass of the (2R,3R)-1 and (2S,3S)-1  $N^6$ -THB-dA adducts by  $E.\ coli$  DNA polymerases and their low mutagenicities. $^{37}$  Structurally, the (2S)- $N^6$ -HB-dA adduct differs from the  $N^6$ -THB-dA adducts 1 due to the presence of the carbon—carbon double bond and the

Chart 2. Structures of Additional Adducts Arising from Alkylation at  $N^6$ -dA by EBD, Styrene Oxide, and DEB

corresponding loss of the two additional hydroxyl groups of the  $N^6$ -THB-dA adduct. For  $N^6$ -THB-dA adducts 1, stereospecific differences in hydrogen bonding patterns are observed for the R,R and the S,S adducts, $^{37,89}$  which may explain the ability of the R,R adduct to cause  $A \rightarrow T$  transversions, while the S,S adduct induces  $A \rightarrow G$  transitions. $^{37}$ 

Our structural results for the (2S)-N6-HB-dA adduct are consistent with published data for N<sup>6</sup>-dA adducts of styrene oxide, which are also accommodated in the major groove and maintain Watson-Crick base pairing. 92 Styrene-induced DNA alkylation can involve either the  $\alpha$  or  $\beta$  carbons of styrene oxide. Feng et al.  $^{93,94}$  showed that stereochemistry at the  $\alpha$ carbon modulates the structure of  $\alpha$ -N<sup>6</sup>-dA adducts **2** of styrene oxide. For the *R* stereoisomer, the styrene ring orients in the 5'direction in the major groove, whereas for the S stereoisomer, the styrene ring orients in the 3'-direction. In contrast, Hennard et al. 95 showed that the R- and S- $\beta$ -N<sup>6</sup>-adenyl-styrene adducts 3 (Chart 2) exhibit similar major groove orientations of the styrene ring, which was attributed to the longer tether of the  $\beta$  adducts. Site-specific mutagenesis studies of these regioisomeric N<sup>6</sup>-dA styrene oxide adducts also indicate low levels of mutations.<sup>96</sup>

The observed structural similarities between the (2S)- $N^6$ -HB-dA adduct investigated here and other known  $N^6$ -dA adducts predict that the (2S)- $N^6$ -HB-dA adduct will be weakly mutagenic. Detailed site-specific mutagenesis studies have not been reported for (2S)- $N^6$ -HB-dA. However, Carmical et al. <sup>37</sup> reported that regioisomeric (1R)- $N^6$ -HB-dA and (1S)- $N^6$ -HB-dA adducts 4 (Chart 2) were nonmutagenic in *Escherichia coli*. <sup>37</sup>

The minor effects of (2S)- $N^6$ -HB-dA adducts on DNA structure are in contrast with the more pronounced distortions induced by the bis-alkylation products at  $N^6$ -dA induced by diepoxybutane (DEB):  $N^6$ , $N^6$ -(2,3-dihydroxybutan-1,4-diyl)-2′-deoxyadenosine ( $N^6$ , $N^6$ -DHB-dA in Chart 2). Two enantiomers of  $N^6$ , $N^6$ -DHB-dA have been identified in DNA: R,R-5 and S,S-5. <sup>97,98</sup> NMR structures of R,R- and S,S-5 in the ras61 sequence  $N^6$  revealed that the 3,4-dihydroxypyrrolidine ring is localized in the major groove but rotates around the C6- $N^6$  bond, allowing for the complementary thymine to remain inserted in the DNA duplex. In contrast to the (2S)- $N^6$ -HB-dA adduct, R,R- and S,S-5 form only one Watson—Crick hydrogen

bond to the complementary thymine between the adenine N1 imino nitrogen and the T<sup>17</sup> N3H imino proton of the complementary strand. <sup>98</sup> As compared to the (2S)- $N^6$ -HB-dA adduct, the R,R- and S,S-S adducts significantly destabilize the duplex, evidenced by a 16–17 °C decrease in  $T_{\rm m}$ . <sup>99</sup> Replication studies conducted *in vitro* have revealed that the DHB-dA adducts block human DNA polymerase  $\beta$  and are bypassed in an error-prone manner by human translesion synthesis (TLS) polymerases  $\kappa$  and  $\eta$ , leading to both base substitution and deletion mutations. <sup>100</sup> Studies are in progress to evaluate polymerase bypass and mutagenicity of the (2S)- $N^6$ -HB-dA adduct.

# SUMMARY

Detailed solution NMR studies reveal that the (2S)- $N^6$ -HB-dA adduct is positioned in the major groove of DNA. This adduct maintains Watson—Crick base pairing with the complementary  $T^{17}$  base and is stacked into the helix. The (2S)- $N^6$ -HB-dA base does not increase the rates of exchange with solvent for the  $C^5$ : $G^{18}$ ,  $Y^6$ : $T^{17}$ , and  $A^7$ : $T^{16}$  base pairs. It modestly perturbs base stacking interactions at base pairs  $C^5$ : $G^{18}$ ,  $Y^6$ : $T^{17}$ , and  $A^7$ : $T^{16}$ , which when combined with the accommodation of the adduct in the major groove, may account for the 5 °C reduction in the  $T_m$  of the duplex in 0.1 M NaCl at pH 7.

#### ASSOCIATED CONTENT

# Supporting Information

Parameters for the (2S)- $N^6$ -HB-dA adduct used in rMD calculations; chemical shifts of the nonexchangeable protons of the (2S)- $N^6$ -HB-dA modified duplex; and chemical shifts of the nonexchangeable protons in the (2S)- $N^6$ -HB-dA modified duplex. This material is available free of charge via the Internet at http://pubs.acs.org.

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#### Notes

The authors declare no competing financial interest.

# ABBREVIATIONS

dA, 2'-deoxyadenosine; BD, 1,3-butadiene; BDT, butadiene triol; CORMA, correlated matrix analysis; COSY, correlated spectroscopy; DIPEA, N,N-diisopropylethyl amine; DMSO, dimethyl sulfoxide; dR, 2'-deoxyribose; DSS, 4,4-dimethyl-4-silapentane-1-sulfonic acid; DEB, 1,2,3,4-diepoxybutane; EB, 1,2-epoxy-3-butene; EBD, 1,2-dihydroxy-3,4-epoxybutane; HPLC, high pressure liquid chromatography; HPLC-ESI-MS, high pressure liquid chromatography—electrospray ionization—mass spectrometry; MALDI-TOF-MS, matrix-assisted laser desorption/ionization time-of-flight mass spectrometry;

 $N^6$ -HB-dA,  $N^6$ -(2-hydroxy-3-buten-1-yl)-2'-deoxyadenosine;  $R_1^x$ , sixth root residual; rMD, restrained molecular dynamics; rmsd, root-mean-square deviation; NOESY, nuclear Overhauser effect spectroscopy

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