Sugar Radicals Formed by Photoexcitation of Guanine Cation Radical in Oligonucleotides

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Received: February 8, 2007; In Final Form: April 11, 2007

This work presents evidence that photoexcitation of guanine cation radical (G\(^{+\ast}\)) in dGpdG and DNA-oligonucleotides TGT, TGTT, TGGGT, TTGTT, TTGGTTG, AGA, and AGGGA in frozen glassy aqueous solutions at low temperatures leads to hole transfer to the sugar phosphate backbone and results in high yields of deoxyribose radicals. In this series of oligonucleotides, we find that G\(^{+\ast}\) on photoexcitation at 143 K leads to the formation of predominantly C5\(^{\ast}\) and C1\(^{\ast}\) with small amounts of C3\(^{\ast}\). Photoconversion yields of G\(^{+\ast}\) to sugar radicals in oligonucleotides decreased as the overall chain length increased. However, for high molecular weight dsDNA (salmon testes) in frozen aqueous solutions, substantial conversion of G\(^{+\ast}\) to C1\(^{\ast}\) (only) sugar radical is still found (ca. 50%). Within the cohort of sugar radicals formed, we find a relative increase in the formation of C1\(^{\ast}\) with length of the oligonucleotide, along with decreases in C3\(^{\ast}\) and C5\(^{\ast}\). For dsDNA in frozen solutions, only the formation of C1\(^{\ast}\) is found via photoexcitation of G\(^{+\ast}\), without a significant temperature dependence (77–180 K). Long wavelength visible light (>540 nm) is observed to be about as effective as light under 540 nm for photoconversion of G\(^{+\ast}\) to sugar radicals for short oligonucleotides but gradually loses effectiveness with chain length. This wavelength dependence is attributed to base-to-base hole transfer for wavelengths >540 nm. Base-to-sugar hole transfer is suggested to dominate under 540 nm. These results may have implications for a number of investigations of hole transfer through DNA in which DNA holes are subjected to continuous visible illumination.

Introduction

Sugar—phosphate free radicals are among the most damaging of DNA lesions because they are precursors to DNA strand breaks.\(^1\) We have found that irradiation of hydrated DNA samples with heavy-ion beams results in substantially increased amounts of sugar and phosphate backbone radicals over that found with low LET \(\gamma\)-irradiation.\(^2\) In these same ion beam samples, we also found evidence for the C3-dephosphorylated and phosphoryl radicals, resulting from immediate strand breaks.\(^2\) The usual mechanism suggested for formation of sugar radicals (hole deprotonation) is that of two sugar radicals nor account for the higher yield of sugar phosphate radicals observed in high LET-irradiated samples. Therefore, two new mechanisms are under serious consideration in our laboratory to explain these increases in sugar radical formation: (i) the role of low-energy electrons (LEE) as a potential source of prompt strand breaks, which we have suggested result in the immediate strand break radicals,\(^3\)–\(^12\) and (ii) the role of excited states of DNA base cation radicals.\(^13\)–\(^17\) This study continues our efforts to understand the role of excited states in the production of DNA sugar—phosphate radicals.

In our early work with \(\gamma\)-irradiated DNA, we found evidence suggestive of conversion of G\(^{+\ast}\) to sugar radicals at high irradiation doses.\(^18\) This led us to suggest a role for excited states of DNA base cation radicals as a source for sugar radical formation.\(^18\) The observation of relatively high yields of neutral sugar radicals in DNA irradiated with high-energy argon-ion beams, relative to that found in \(\gamma\)-irradiated samples, also led us to hypothesize that excited states in the densely ionized ion beam track core may lead to sugar radicals.\(^2\)

In our ongoing efforts to delineate the role of involvement of excited states of one-electron-oxidized base cation radicals in the formation of sugar radicals, we have already observed that photoexcitation of guanine cation radical (G\(^{+\ast}\)) in aqueous (D\(_2\)O) glassy systems produced primarily C5\(^{\ast}\) and C3\(^{\ast}\) (in dGuo), C5\(^{\ast}\) and C1\(^{\ast}\) (in 3\(^{\prime}\)-dGMP), and predominantly C1\(^{\ast}\) (in 5\(^{\prime}\)-dGMP and in the dinucleoside phosphate TpdG).\(^13\)–\(^15\) Sugar radical formation was also found by photoexcitation of one-electron-oxidized adenine in deoxynucleosides and deoxynucleotides.\(^16\) Wavelengths of light from 320 to 650 nm are found to be effective.\(^14,16\) Photoexcitation of G\(^{+\ast}\) in \(\gamma\)-irradiated hydrated (\(I = 12 \pm 2\) D\(_2\)O/nucleotide) DNA in the UVA-vis range (310–480 nm) shows conversion of G\(^{+\ast}\) to C1\(^{\ast}\) in substantial yields.\(^14\) Photoexcitation of G\(^{+\ast}\) in DNA at wavelengths above 500 nm was not effective.\(^14\) These experiments along with theoretical studies using the time-dependent density functional theory (TD-DFT)\(^16\) have established that excitation of one-electron-oxidized purine base radical results in delocalization of a significant fraction of the spin and charge onto the sugar moiety, followed by rapid deprotonation, leading to the formation of a neutral sugar radical. Moreover, TD-DFT studies in TpdG\(^15\) and also in other dinucleoside phosphates\(^17\) show evidence for a competitive excitation process, base-to-base hole transfer, which was predicted to occur at low excitation energies in stacked DNA base systems.

Although the experimental and theoretical efforts described above have established the role of hole-excited states in the sugar radical formation in DNA, a number of points still remain unexplained:

(i) Why are wavelengths over 500 nm effective in sugar radical formation in the photoexcitation of G\(^{+\ast}\) in small DNA model systems, but not in dsDNA;
Materials and Methods

Model Compound Sample Preparation. The dinucleoside phosphate (monosodium salt) dGpdG and lithium chloride (99% anhydrous, SigmaUlta) were procured from Sigma (St. Louis, MO). The DNA oligonucleotides TGT, TTGGT, AGA, desalted and tested by MALDI-TOF mass spectrometry, were purchased from Synthegen, LLC. (Houston, TX). 5′-Phosphorylated and desalted 5′-p-TTGGT was obtained from IDT (Coralville, IA). The remaining oligonucleotides TTGGT, TTGGT, TGGGTTG, and AGGGA in this study were all obtained from IDT with standard desalting (Coralville, IA). Potassium persulfate (crystal) was from Mallinckrodt, Inc. (Paris, KY). All were employed without further purification.

About 0.5 mg of each of the dinucleoside phosphates or 1.5 mg of each of the oligonucleotides was dissolved in 0.35 mL of 7.5 M LiCl in D₂O in the presence of 2 mg of K₂S₂O₈. The solutions were deoxygenated by bubbling with nitrogen and as per our earlier works. The transparent glassy samples were prepared by drawing the solution into 4 mm Suprasil quartz tubes (Catalogue no. 734-PQ-8, Wilmad Glass Co., Inc., Buena, NJ), followed by cooling to 77 K. All samples are stored at 77 K in the dark.

DNA Sample Preparation. Salmon testes DNA (sodium salt, 57.3% AT and 42.7% GC, Sigma Chemical Company, St. Louis, MO), trichloration and deuterium oxide (99.9 atom % D, Aldrich Chemical Co. Inc., Milwaukee, WI) were used without any further purification.

Ice-like samples of DNA with Tl³⁺, at loading 1 Tl³⁺/10 bp in D₂O were prepared using procedures described earlier. This loading of Tl³⁺ thoroughly suppresses the reductive-damage pathway and allows observation of the formation of sugar radicals from photoexcited G⁺⁺ in DNA. All samples were stored at 77 K in the dark.

γ-Irradiation. γ-Irradiation of glassy samples of all oligonucleotides was performed using a model 109-GR 9 irradiator which contains a shielded 60Co source. A 400 mL Styrofoam Dewar containing the samples under liquid nitrogen enters the irradiation chamber via an elevator system that prevents exposure. Glassy samples of all the oligonucleotides containing a number of bases up to four were γ-irradiated (60Co) with an absorbed dose of 2.5 kGy at 77 K. For glassy samples of oligonucleotides containing a number of bases higher than four, the absorbed dose was 5 kGy at 77 K. Salmon testes DNA-Tl³⁺ ice samples were γ-irradiated (60Co) with an absorbed dose of 15.4 kGy at 77 K, as per our earlier work.

Annealing and Illumination of Samples. As mentioned in our previous works, we have used a variable temperature assembly for carrying out the annealing of the samples. For monitoring the temperatures during annealing, we have used a copper-constantan thermocouple in direct contact with the sample. The glassy samples (7.5 M LiCl/D₂O) stored at 77 K were annealed to 155 K for 10–20 min, which results in the loss of Cl₂⁻ with the concomitant formation of only G⁺⁺. Note that we did not observe sugar radical formation by direct attack of Cl₂⁻ on the sugar moiety in these oligonucleosides/oligodeoxyribose radicals.

Photoexcitation of these glassy samples of oligonucleosides or dinucleoside phosphate containing the G⁺⁺ were then carried out at 143 K with a 250 W tungsten lamp with or without a cutoff filter (≤540 nm). The active visible light intensity at the sample was a small fraction of the total intensity (ca. 60 mW).

Following our earlier works, γ-irradiated DNA-Tl³⁺ ice samples were annealed to 130 K to remove the ESR signal from •OH. The •OH is in a separate ice phase, and annealing does not result in additional DNA radicals. These annealed samples were then illuminated using a 250 W tungsten lamp at 143 K with and without a cutoff filter (≤540 nm) and at 180 K without any filter.

Electron Spin Resonance. After γ-irradiation at 77 K, annealing to 155 K, and illumination at 143 K, samples were immediately immersed in liquid nitrogen, and an ESR spectrum was recorded at 77 K and 40 dB (20 μW). After γ-irradiation at 77 K, annealing to 130 K, and illumination at 143 K and at 180 K, DNA-Tl³⁺ ice samples were immediately immersed in liquid nitrogen, and an ESR spectrum was recorded at 77 K and 40 dB (20 μW). We have recorded these ESR spectra using a Varian Century Series ESR spectrometer operating at 9.2 GHz with an E-4531 dual cavity, a 9 in. magnet, and with a 200 mW klystron. Similar to our previous works, Femy’s salt (with g = 2.0056, ΑΝ = 13.09 G) was used for field calibration. In the glassy samples of oligonucleosides or dinucleoside phosphate, we have subtracted a small singlet "spike" from irradiated quartz at g = 2.0006 from spectra before analyses.

Analysis of ESR Spectra. The fraction that a particular radical contributes to an overall spectrum is estimated from doubly integrated areas of benchmark spectra. The doubly integrated areas are directly proportional to the number of spins of each radical species (moles of each radical). Least-squares fittings of benchmark spectra (Figure 1) were employed to determine the fractional composition of radicals in the experimental spectra using programs (ESRPLAY, ESRADSUB) written in our laboratory.

The benchmark spectra for the glassy samples of dinucleoside phosphate and oligonucleotides are shown in Figure 1. These are for G⁺⁺ from dGpdG (Figure 1A) and C5⁺ (Figure 1B) and C3⁺ (Figure 1C) from dGuo. We have used two benchmark spectra for C1⁺ (Figure 1D) for TGT and E (for dGpdG) because the two major hyperfine coupling constants for the two C2′–H atoms (β-proton couplings) vary slightly with compound (see Supporting Information, Figure S1). The origin of the benchmark spectra for C1⁺ in TGT and dGpdG as well as hyperfine couplings and the g-values are given in Supporting Information Figure S1.

The spectra obtained from DNA-Tl³⁺ ice samples were analyzed using the benchmark spectra of the guanine cation radical (G⁺⁺), of the one-electron reduced species, namely, T⁺⁺; and C(N3H)⁺, of a composite spectrum of neutral radicals found.
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Figure 1. Benchmark spectra used for computer analysis. (A) G** in dGpdG via one-electron oxidation by Cl₂•• (see Figure 2A also). (B) C5•, formed via photoexcitation of G* in 8-dT•-dGMP. (C) C3•, produced from G** in dGuo. (D) C1•, produced from G** in TGT. (E) C1•, produced from G** in dGpdG. (see Supporting Information Figure S1 for details regarding Figure 1D and E). The three reference markers in this figure and in the other figures in this work represent positions of Frenzy’s salt resonances (the central marker is at g = 2.0086, and each of three markers is separated from one another by 13.09 G).

Figure 2. (A) ESR spectrum of one-electron-oxidized dGpdG (1 mg/mL) by Cl₂•• attack in the presence of K₂S₂O₈ (5 mg/mL) as an electron scavenger in 7.5 M LiCl (D₂O) glasses at low temperatures, the proton at N1 in G** is ~50% retained; thus, about half of the radicals one-electron-oxidized guanine are in the deprotonated form, G(−H)•. The ESR spectra of G* and G(−H)• are very similar. For ease of discussion, in this work, the one-electron-oxidized guanine will be referred to as G**.

Figure 2B shows formation of sugar radicals after photoexcitation of G** at 143 K. Prominent outer line components from C1• and an intense central ~19 G doublet from C5• are visible in the spectrum. We also observe, in low intensity, line components of C3• in the wings (arrows). Analysis using the benchmark spectra shown in Figure 1 indicates that the spectrum in Figure 2B contains contributions from C1• (33%), C5• (53%), C3• (4%) and a small residual amount of G** (ca. 10%).

Results

Photoexcitation of One-Electron-Oxidized dGpdG. In Figure 2A, we show the ESR spectrum of G** formed in the dinucleoside phosphate dGpdG after one-electron oxidation by Cl₂•• on annealing to 155 K. This spectrum in Figure 2A is found to be identical to that of the G** spectrum already reported in the literature. The presence of G** in this sample is also validated by its characteristic UV-vis absorption, which produces red-violet color. In other work, we have shown that in 7.5 M LiCl (D₂O) glasses at low temperatures, the proton at N1 in G** is ~50% retained; thus, about half of the radicals one-electron-oxidized guanine are in the deprotonated form, G(−H)•. The ESR spectra of G* and G(−H)• are very similar.

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Photoexcitation of One-Electron-Oxidized DNA-Oligonucleotide Radicals. In Figure 3A and F, the ESR spectra of one-electron-oxidized TGT and TTGTTT are shown. These two spectra and those of one-electron-oxidized TGGT, 5′-p-TGGT, TGCGT, TTGTTT, and TTGGTTGGTT (see Supporting Information Figure S2), as well as that of one-electron-oxidized AGGGAG (see Supporting Information Figure S3), are found to be similar. Using the benchmark spectrum of G** in Figure 1 A, we find that the spectra in Figure 3A and F and those of the one-electron-oxidized oligonucleotides mentioned above are all found to result predominantly from G**.

Although thymine is not found to be significantly oxidized in oligonucleotides containing T and G, we find that A was also oxidized in AGA and AGGGAG. Using the benchmark spectrum of deprotonated one-electron-oxidized adenine, that is, A(−H)• from our earlier work and the benchmark spectrum of G** in Figure 1A, we find that the spectrum of one-electron-oxidized AGA (see Supporting Information Figure S3) is a mixture of G** (ca. 60%) and A(−H)• (ca. 40%). The guanine base is preferentially oxidized by Cl₂•• due to its lower redox potential than adenine.

Formation of sugar radicals via photoexcitation of G** in oligonucleotides at 143 K is evidenced in the ESR spectra shown in Figure 3B–E and G–I, as well as in one-electron-oxidized AGA and AGGGAG (see Supporting Information Figure S3). Prominent line components from C1• and an intense central ~19 G doublet from C5• are visible in these spectra. We do also observe small amounts of the line components of C3• in the wings in these systems.

Analyses of the spectra shown in Figure 3B–E and G–I were obtained after photoexcitation using benchmark spectra (Figure 1). The percent conversion of one-electron-oxidized dinucleoside monophosphates and oligonucleotides to the sugar radicals, the initial rate of sugar radical formation, and relative percentages of various types of sugar radicals at 143 K in the presence and in absence of a cutoff filter (≤540 nm) are presented in Table 1 and are compared with the previous results obtained from samples of dGuo and its 3′- and 5′- nucleotides at 143 K as well as at 77 K. Initial rates shown in Table 1 represent the rate of sugar radical production during the first 20 min in percent per minute and are determined by analyses of the spectra for the fraction of sugar radicals produced after the initial 20 min photoexcitation.

Our analyses shown in Table 1 suggest the following salient points:

(i) Wavelength Dependence for Sugar Radical formation via Photoexcitation of G**. We find that the initial rate as well as overall extent of sugar radical formation via photoexcitation of G** at wavelengths ≥540 nm decreases as the overall chain length increases from dGuo to TTGGTTGGTT. In dsDNA ice samples, the overall conversion of sugar radicals is slightly lower than the largest oligonucleotide, and the initial rate of sugar radical formation via photoexcitation becomes negligible at wavelengths ≥540 nm.
Figure 3. ESR spectra of A and F of one-electron-oxidized TGT (4.5 mg/mL) and TTGGT (4.5 mg/mL) formed by attack of Cl₄⁺ in the presence K₂S₂O₅ (8 mg/mL) as an electron scavenger in 7.5 M LiCl glass/D₂O, in annealing to 155 K. These spectra in A and F are chiefly due to G⁺. Spectra B–E and G–I are obtained after photoexcitation of G⁺ in the oligonucleotides indicated with visible light at 143 K, showing considerable conversion of G⁺⁺ to sugar radicals, C₅⁺ (prominent doublet at the center) as well as to C1⁺ (prominent quartet). The samples in A and F are red-violet due to the absorption of G⁺⁺, whereas those in B–E and G–I are faded in color or are near colorless as conversion to sugar radical occurs. All spectra were recorded at 77 K.

**TABLE 1: Sugar Radicals Formed via Photoexcitation of One-Electron-Oxidized Dinucleoside Phosphates and Oligonucleotides at 143 K**

<table>
<thead>
<tr>
<th>compd</th>
<th>% converteda</th>
<th>init. rate (%/min)</th>
<th>C1⁺e</th>
<th>C3⁺e</th>
<th>C5⁺e</th>
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<td>dGuo/</td>
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<td>4.2</td>
<td>3.5</td>
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<td>35</td>
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<tr>
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<td>40</td>
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<td>(77 K)</td>
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<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>5'-dGMP/</td>
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<td>95</td>
<td>5</td>
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<tr>
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<td>(77 K)</td>
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<td>100</td>
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</table>

a Radical percentages expressed to ±10% relative error. Initial rates are based on the percent G⁺ converted to sugar radicals after the first 20 min of visible light exposure and should be considered as indicative of the relative rates of sugar radical formation. All glassy samples are at the native pH of 7.5 M LiCl (ca. 5). All samples were illuminated at 143 K unless indicated otherwise. Percentage of conversion of one-electron-oxidized dinucleoside phosphate or oligonucleotide to sugar radicals. The total spectral intensities before and after illumination were the same within experimental uncertainties. Each calculated as the percentage of total sugar radical concentration; these sum to 100%. Ref 14. Ref 15. The overall spectrum obtained after photoexcitation also has an underlying unidentified spectrum (ca. 40%). DNA (salmon testes) in a frozen aqueous (D₂O) solution (ice).

(ii) Effect of the sequence and length of the oligomers on the initial rate as well as overall production of sugar radicals via photoexcitation. (a) Photoexcitation of G⁺⁺ in both TpdG and 5'-dGMP at 143 K are unusual as they show conversion to C1⁺ and C5⁺ sugar radicals only, whereas all others show significant yields of C5⁺. We note that the C5⁺ formation did occur in 5'-dGMP when photoexcitation was carried out at 77 K (see Table 1). This observation leads us to suggest that conforma-

tion of the sugar phosphate ring in the excited state may control the sites of deprotonation.

(b) Photoexcitation of G⁺⁺ in dGuo and in 8-n-dGuo shows that C8 deuterium substitution in the guanine moiety has no effect on the initial rate or on the sugar radical cohort.

(c) Phosphorylation at 3' reduces the initial rate of formation of sugar radicals by a factor of 2, whereas at 5', substitution does not reduce the initial rate. For example, compare the data of dGuo and 3'-dGMP as well as TpdG and TGT in Table 1, which show the factor of 2 in the initial rates, whereas dGuo and 5'-dGMP as well as TTGGT and 5'-p-TTGGT show the same initial rate.

(d) Photoconversion yields of G⁺⁺ to sugar radicals in oligonucleotides decreases as the overall chain length increases. However, for high molecular weight dsDNA (salmon testes) in frozen aqueous solutions, substantial conversion of G⁺⁺ to C1⁺ (only) is still found.

(e) Within the cohort of sugar radicals formed, we find a relative increase in the formation of C1⁺ with length of the oligonucleotide along with a decrease in C3⁺ production while C5⁺ also decreases, but to a lesser extent. In dsDNA, similar to our earlier work, formation of only C1⁺ via photoexcitation is observed.

(f) In earlier work, we showed that phosphate substitution at a 3' or 5' position in nucleosides tended to deactivate radical formation at that site, presumably by increasing the bond strength of the C=H bond at that site. Thus, one hypothesis for the large extent of C5⁺ formation in these systems was that the hole was transferred from G⁺⁺ to the 5'-nucleotide, which has a 5'-OH. Therefore, expecting a decrease in C5⁺ formation via photoexcitation of G⁺⁺, we tested 5'-p-TGTT in addition to TTGGT. However, the initial rate and the final sugar radical cohort remained identical in TTGGT and in 5'-p-TTGGT. Thus, the presence of the phosphate moiety at the 5'-end in 5'-p-TTGGT apparently does not have a significant influence on the rate or the type of sugar radical formed via photoexcitation.

Temperature Dependence of These Photoexcitation Reactions in DNA. In Figure 4A, from our earlier work, we show the spectrum obtained after photoexcitation of G⁺⁺ in the ice samples of high molecular weight dsDNA (50 mg salmon testes DNA/mL in D₂O) with 1 Tl⁺/10 base pairs for 110 min using
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Figure 4. Ice samples of DNA (50 mg/mL in D₂O) with 1 Tl⁺/10 base pairs were γ-irradiated to 15.4 kGy dose at 77 K and were then annealed to 130 K to remove the •OH signal. (A) ESR spectrum obtained after illumination with the aid of a high-pressure Xe lamp (Oriel Corporation), at 77 K for 110 min with 380–480 nm band-pass filter.14 (B) After photoexcitation using a 250 W photoflood tungsten lamp of an identically prepared and handled sample as in A at 143 K for 90 min without any filter. (C) After illumination by the same photoflood lamp of another identically prepared and handled sample as in A and B at 180 K for 90 min. Spectrum C also shows outer line component of peroxy radical, which decreases the intensity of the sugar components slightly. All spectra were recorded at 77 K.

A 380–480 nm band-pass filter. In this case, photoexcitation was carried out using a high-pressure Xe lamp at 77 K. This spectrum shows prominent line components of C1′ at the wings.14 In Figure 4B and C, ESR spectra are presented of identically prepared and handled samples as in Figure 4A after photoexcitation using a 250 W photoflood lamp at 143 K for 90 min (4B) and 180 K for 90 min (4C). In Figure 4A, B, and C, we observe prominent outer line components of C1′ in the wings in spectra. In dsDNA in ices, unlike the deoxyribonucleoside/-tide systems in glasses, we do not observe an increase in the conversion of G⁺⁺ to C1′ at 143 K, as opposed to 77 K (see Table 1). The spectrum in Figure 4C for photolysis at 180 K shows evidence of peroxy radicals in the wings, which result from reaction of residual oxygen in the sample with the sugar radicals formed. However, we still observe the prominent line components of C1′. Thus, we conclude that the mechanisms in the formation of C1′ via photoexcitation of G⁺⁺ are functional up to at least 180 K. From the lack of temperature dependence observed from 77 to 180 K, it is likely that at temperatures which more closely correspond to biological conditions, this mechanism will be operative.

Discussion

Wavelength and DNA Strand Length Influence on Sugar Radical Formation with Photo-Oxidation of G⁺⁺: The initial yield found for sugar radical formation was highest in nucleosides or dinucleoside phosphates and decreased gradually to 55% yield as the oligonucleotide increased in length. A definitive wavelength dependence on yield was observed for the longest oligonucleotide TTGGTGTGGT as well as high molecular weight dsDNA. TD-DFT calculations for excited G⁺⁺ in dGuo predict allowed electronic transitions throughout the whole visible spectral region that show delocalization of spin and positive charge on the sugar moiety. These calculations agree with experimental findings that for excited G⁺⁺ in dGuo, radical yields and radical identities are relatively independent of wavelength. When these TD-DFT calculations were extended to excited G⁺⁺ in TpdG and to other dinucleoside phosphates, the calculated transition energies indicated that base-to-base hole transfer is predicted at the longest wavelengths and that hole transfer to the sugar—phosphate moiety is predicted in the UVA-vis wavelengths. These theoretical results offer an explanation for our experimental finding regarding decline in the yield of sugar radical formation at longer wavelengths with oligonucleotide length; that is, at longer wavelengths (>500 nm), base-to-base hole transfers occur which do not transfer the hole to the sugar—phosphate portion and thereby prevent DNA-sugar damage.

Although base-to-base hole transfer in the oligonucleotides studied is expected to place holes on thymine on the basis of the recent TD-DFT calculations, experimentally, no significant line component of thymine radicals was observed during formation of C1′ and C3′ via photoexcitation of G⁺⁺ in TpdG.15 Similarly, we did not observe line components of thymine radicals in oligonucleotides studied here (see Figure 3B–E and G–I and also Supporting Information Figure S3) or in DNA (see Figure 4 in this study and Figure 3 in ref 14) during or after photoconversion of G⁺⁺ to sugar radicals. It is well-known that base stacking in oligonucleotides and in dsDNA allows photoinduced transfer of spin and charge to nearby bases, as suggested in this work.25–27 We note that for double-stranded oligonucleotides lacking guanine bases, formation of the allylic thymyl, that is, UCH· radical, has been observed by Schuster and his co-workers.28 Initial ongoing studies on our part (not shown) regarding photooxidation of one-electron-oxidized TpdA and dApT point toward slightly higher yields of thymine radical components on photooxidation of the one-electron-oxidized dinucleoside phosphates.

Why Does Photooxidation of G⁺⁺ in dGuo Result in C3′, C5′, and C1′ but Only C1′ in dsDNA? Among all the model compounds including mononucleosides/-tides, dinucleoside phosphates, and oligonucleotides listed in Table 1, G⁺⁺ in 5′-dGMP and in TpdG show very high initial rates of conversion as well as the near complete conversion to C1′ (90–95% of sugar radicals) upon photooxidation at 143 K. However, for 5′-dGMP at 77 K, the results in Table 1 show less overall conversion to sugar radicals, with little C1′ formation (15% of sugar radicals) but with substantial amounts of C3′ and C5′. Earlier, we showed that warming of the glassy samples of 5′-dGMP (already photoexcited at 77 K) from 77 to 143 K in the dark without further photooxidation results in only small changes in hyperfine splittings, with no changes in radical distribution. Thus, the C5′ and C3′ do not convert to C1′ on annealing.14 Interestingly, while we find predominantly C1′ and no C5′ formation in TpdG, we find substantial C5′ formation in TGT (50%) and in TGGT (60%) (Table 1). The presence of the central doublet found in spectra shown in Figure 3B and C clearly and unequivocally point to the formation of C5′ in the one-electron-oxidized TGT and TGGT samples. However, it is not clear whether the C5′ radical is formed at the C5′ position on G or at the C5′ position on the 5′-terminal T. TD-DFT calculations of one-electron-oxidized TpdG predicts that during photooxidation (1.5 eV), hole transfer to the sugar moiety attached to the 5′-thymine will occur.15 Therefore, theory does allow for C5′ formation at several sites on TGGT. In TGT, if the 5′-terminal T site were the site for C5′ formation, we would expect that the presence of a phosphate moiety at the 5′-end in TGGT (i.e., in 5′-p-TGGT) would hinder C5′ formation at that site. However, no such effect was observed experimentally (see Table 1 and compare spectra shown in Figure 3C and D).
Many of the above observations suggest that it is more than just gross structure which determines the deprotonation site. These results strongly suggest that conformation of the one-electron-oxidized radicals in the excited state during photoexcitation is critical. It is likely that conformation alters the hole distribution on the sugar ring and, thus, the final sugar radical cohort during photoexcitation at 143 K in the excited state. We also note here that recent ion-imaging studies regarding photodissociation of propanal cation\textsuperscript{29} show that fast intersystem crossing occurs from the excited state to a hot ground state and that this results in the products observed. Furthermore, the authors find that the specific pathway and the type of products formed depends critically on the conformation of the cation radical. Thus, although this work is in accord with our suggestion that the type of sugar radical formed via photoexcitation of G\textsuperscript{+} depends upon its conformation, this work also suggests that in our work, the deprotonation from the sugar moiety leading to the sugar radical formation may not occur directly from the excited cation state, but rather, from a “hot” cation ground state.

Is the Mechanism of Sugar Radical Formation via Photo-Oxidation of One-Electron-Oxidized Base Radical Found Applicable to Biologically Relevant Temperatures? The formation of C1′ due to photoexcitation of G\textsuperscript{+} in dsDNA ice samples is shown in Figure 4 to occur at 180 K as well as at 77 and 143 K. Although these temperatures still do not correspond to biological conditions, the lack of a significant temperature dependence in the initial rate of C1′ formation upon photoexcitation of G\textsuperscript{+} in dsDNA from 77 to 180 K suggests applicability at higher temperatures. The availability of multiple local sites on the DNA itself as proton acceptors as well as a structure that inherently allows for facile proton transfer, we believe, accounts for the lack of significant temperature dependence of C1′ formation upon photoexcitation of G\textsuperscript{+} in frozen samples of highly polymerized dsDNA.

Conclusions

This work shows evidence that photoexcitation of one-electron-oxidized DNA-dinucleoside phosphates and DNA-oligonucleotides (no. of bases \( \leq 5 \)) lead to the formation of high yields (ca. 75–90\%) of deoxyribose-sugar radicals. The extent of photoconversion to sugar radicals as well as its initial rate decreases with increasing size of the oligonucleotides but remains substantial (50\%), even for DNA of thousands of base pairs in length (salmon testes DNA). Specific sugar radicals have been identified predominantly from deprotonation at the C5′ and C1′ positions in the deoxyribose moiety. In our earlier experimental and theoretical works,\textsuperscript{13–17} we have shown that photoexcited one-electron-oxidized adenine and guanine base radicals are quenched by formation of sugar radicals. These results may have implications for a number of investigations of hole transfer through DNA\textsuperscript{25–27} in which DNA holes are subjected to continuous visible illumination. These investigations rely on the competition of hole transfer with the quenching reaction of the hole (e.g., G\textsuperscript{+}) with water, forming 8-HO-G and ultimately 8-oxo-G\textsuperscript{25–27}. Therefore, from this work and also from our earlier works,\textsuperscript{13–17} we note that photoinduced DNA-hole transfer and its conversion to sugar radicals are also expected along with the natural DNA-hole transfer processes, such as tunneling and thermally activated hopping reported in the literature.\textsuperscript{25–27} and references therein

Acknowledgment. This work was supported by the NIH NCI under Grant No. R01CA045424. A.A. is grateful to the authorities of the Rajdhani College and the University of Delhi for leave to work on this research program.

Supporting Information Available: Supporting information is available and it contains the following: (1) Figure S1, showing comparison of the C1′ spectra obtained experimentally in TGT and in dGpdG via subtraction and via simulation. (2) Figure S2, representing ESR spectra of one-electron-oxidized TGT, TGGT, 5′,p-TGGT, TGGGT, TTGGT, and TTGGTTGGTT and showing that each one of these spectra is mainly due to G\textsuperscript{+}. (3) ESR spectra of one-electron-oxidized AGA and AGGGA, followed by photo-oxidation leading to the sugar radical formation. This material is available free of charge via the Internet at http://pubs.acs.org.

References and Notes


Sugar Radicals


