The Influence of Single, Tandem, and Clustered DNA Damage on the Electronic Properties of the Double Helix: A Theoretical Study

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Abstract: Oxidatively generated damage to DNA frequently appears in the human genome as the effect of aerobic metabolism or as the result of exposure to exogenous oxidizing agents, such as ionization radiation. In this paper, the electronic properties of single, tandem, and clustered DNA damage in comparison with native ds-DNA are discussed as a comparative analysis for the first time. A single lesion—8-oxo-7,8-dihydro-2′-deoxyguanosine (G\textsubscript{oxo}), a tandem lesion—(5′S) and (5′R) 5′,8-cyclo-2′-deoxyadenosine (cdA), and the presence of both of them in one helix turn as clustered DNA damage were chosen and taken into consideration. The lowest vertical and adiabatic potential (VIP ≈ 5.9 and AIP ≈ 5.5 eV, respectively) were found for G\textsubscript{oxo}, independently of the discussed DNA lesion type and their distribution within the double helix. Moreover, the VIP and AIP were assigned for ds-trimers, ds-dimers and single base pairs isolated from parental ds-hexamers in their neutral and cationic forms. The above results were confirmed by the charge and spin density population, which revealed that G\textsubscript{oxo} can be considered as a cation radical point of destination independently of the DNA damage type (single, tandem, or clustered). Additionally, the different influences of cdA on the charge transfer rate were found and discussed in the context of tandem and clustered lesions. Because oligonucleotide lesions are effectively produced as a result of ionization factors, the presented data in this article might be valuable in developing a new scheme of anticancer radiotherapy efficiency.

Keywords: DNA damage; electronic properties; charge transfer; DFT; (5′R)/(5′S)-5′,8-cyclo-2′-deoxyadenosine; 8-oxo-7,8-dihydro-2′-deoxyguanosine

1. Introduction

DNA is a storage house of genetic information in each cell of a living organism [1]. This information is continuously exposed to different kinds of harmful endo- and exogenous factors, such as ionization radiation (UV, gamma, X-ray), metabolic byproducts, etc. Their interaction with cellular oligonucleotides can cause the formation of DNA lesions both directly and indirectly. Until now, more than 70 types have been identified [2]. It is generally recognized that in the human body, \(3 \times 10^{17}\) DNA damage events per hour take place [3]. On the other hand, DNA lesions can be formed by the activity of the reactive oxygen/nitric species (ROS, RNS) [4]. It has been estimated that approximately \(2 \times 10^4\) free radical events per cell per day take place. Moreover, their number can increase with physical activity by up to 50% [5]. Of the plethora of radical oxygen species, the hydroxyl radical (\(^{•}\)OH) has been found as the most reactive, with \(k = 2 \pm 10^9 \text{ M}^{-1}\text{s}^{-1}\) [6]. From the DNA damage distribution perspective, three main types of lesions can be distinguished: (a) Isolated—one lesion per one helix turn; (b) clustered—two or more per turn; and (c) tandem lesions as a result of a single DNA damage event in which a reactive nucleotide intermediate reacts with an adjacent one [7]. The DNA damage structures discussed in the article are shown in Figure 1.
Figure 1. Graphical representation of the structure of the discussed DNA damage.

Among all DNA lesions, 8-oxo-7,8-dihydro-2′-deoxyguanosine (dG\(^{\text{oxo}}\)) is recognized as the most abundant. Its frequency in a cell has been estimated at \(5.5 \times 10^8\) [8,9]. At the other end of the scale, (5′R)/(5′S)-5′,8-cyclo-2′-deoxyadenosine ((5′R)-cdA and (5′S)-cdA) exist, and their frequency in cellular environments has been assigned as unequal 0.07/0.93 \(\times 10^6\) R-cdA/S-cdA, respectively [10]. However, these results have yet to be verified scientifically and are still under discussion [11,12]. From the cellular point of view, these two diastereomers exhibit different biological/biochemical effects [13–16]. The stability of genetic information is crucial for the future generation of a species, and several repair systems are present in the cell, such as the base/nucleotide repair system (BER, NER), homologous and non-homologous end joining (HEJ, NHEJ), and nucleotide incision repair (NIR) [17]. Their correct activity guarantees the suitable nucleoside sequence in DNA. A failure, however, for example, in NER, can lead to different genetic disorders, cancer, or neurodegenerative disease [18–20].

For all the above repair processes to be effective, the recognition step is the most vital. The cascade of BER proteins starts from the glycosylases’ action. These enzymes can recognize and remove simple DNA damage, such as dG\(^{\text{oxo}}\), 2′-deoxyuridine [21]. To keep genetic material reproducible and stable, several specific glycosylases exist in cells, for example, OGG1 (8-oxo-guanine glycosylase 1), MutY (adenine DNA glycosylase), and UDG (uracil-DNA glycosylase) [22,23]. On the other hand, due to the additional C5′-C8 covalent bond, neither diastereomer of cdA is a substrate for the BER system—no cdA-specific glycosylases are known. The tandem lesions in both diastereomeric forms (5′R)-cdA and (5′S)-cdA are removed from the genome by the more complicated NER machinery. It is important to mention here that these small molecules, depending on the configuration on the 5′ carbon S or R, can significantly change the global structure of the DNA double helix, and as a result are removed from the genome at different rates [24]. The structure of the double helix and its changes are commonly described by a DNA standard reference frame. This analysis uses parameters that are useful for hydrogen bonding and base pairs’ stacking interaction description, which are fundamental for spatial DNA geometry pronunciation. For further details, please see the work of Olson et al. [25].

From an electronic point of view, ds-DNA can be perceived as a conductor of nanofibers [26], which has been shown by Shuster, Barton among others [27,28]. Recently, it was proposed that this phenomenon can allow MutY to scan the genome effectively with the electron transfer mode [29], even though the number of these protein copies is relatively low. MutY is able to verify/scan the E. coli genome \((5 \times 10^9\) base pairs\) within 10 s [30]. For details, please see the review Barton et al. [31].

In this paper, comparative studies between isolated, clustered, and tandem DNA lesions, contained in (5′R)-cdA, (5′S)-cdA and dG\(^{\text{oxo}}\), and their influence on the electronic properties and charge transfer (CT) process of the double helix were considered. It is worth noting that little data exists in the literature that is dedicated to the influence of DNA damage on the hole transfer in ds-DNA [32–34].
2. Results and Discussion

To elucidate the influence of different types of DNA damage on charge transfer induced by a one-electron oxidizing event, nine double-stranded (ds) hexamers were chosen (Table 1).

Table 1. Nucleobase sequence “structures” of double-stranded oligonucleotides taken into theoretical consideration. \(^{\text{oxo}}G\) - 8-oxo-7,8-dihydro-2'-deoxyguanosine, (5'S)-cA: (5'S)-5',8-cyclo-2'-deoxyadenosine, (5'R)-cA: (5'R)-5',8-cyclo-2'-deoxyadenosine.

| DNA Damage Type | Oligonucleotide | Oligonucleotide Base Sequence |
|-----------------|----------------|------------------------------|
| Undamaged Native ds-DNA | N-DNA | d[A1G2A3G4G5G6]d[T1C2G3C4A3C2T1] |
| Single | 3G\(^{\text{oxo}}\)-N-DNA | d[A1G2A3\(^{\text{oxo}}\)G4G5G6]d[T1C2G3C4A3C2T1] |
| | 5G\(^{\text{oxo}}\)-N-DNA | d[A1\(^{\text{oxo}}\)G2A3G4G5G6]d[T1C2G3C4A3C2T1] |
| Tandem | ScA-DNA | d[A1G2(5'S)cA3G4G5G6]d[T1C2G3C4A3C2T1] |
| Clustered | 3G\(^{\text{oxo}}\)-ScA-DNA | d[A1G2(5'S)cA3\(^{\text{oxo}}\)G4G5G6]d[T1C2G3C4A3C2T1] |
| | 5G\(^{\text{oxo}}\)-ScA-DNA | d[A1\(^{\text{oxo}}\)G2(5'S)cA3G4G5G6]d[T1C2G3C4A3C2T1] |
| Tandem | RcA-DNA | d[A1G2(5'R)cA3G4G5G6]d[T1C2G3C4A3C2T1] |
| Clustered | 3G\(^{\text{oxo}}\)-RcA-DNA | d[A1G2(5'R)cA3\(^{\text{oxo}}\)G4G5G6]d[T1C2G3C4A3C2T1] |
| | 5G\(^{\text{oxo}}\)-RcA-DNA | d[A1\(^{\text{oxo}}\)G2(5'R)cA3G4G5G6]d[T1C2G3C4A3C2T1] |

The damage of interest was positioned in the central part of ds-oigo. The initial geometry of each in neutral and radical cation forms was optimized at the M062x/D95*-UFF level of theory in the aqueous phase using our own n-layered integrated molecular orbital and molecular mechanics (ONIOM) strategy [35]. The M062x functional was chosen as being suitable for estimating noncovalent interactions. Although the optimization was done for structural studies; additionally, the D95* basis set was used due to its efficiency in calculating such complicated systems in a reasonable time frame [36,37]. The electronic properties of the discussed ds-hexamers were obtained at the M062x/6-31+G** level of theory in the aqueous phase. However, due to the nature of DNA solvation, the aqueous phase relaxation influences the vertical ionization potential, and the vertical electron attachment was omitted [38]. This choice was sanctioned by the fact that the double helix is solvated from its outer and not internal shape where the base pair aromatic rings stack. The formed scaffold is the highway for hole migration via the hole hopping or super-exchange mechanism [39]. From the structural point of view, although the optimization of spatial geometry was performed for hexamers, only the central part (tetramer) was given further theoretical consideration. It is well known, and indeed observed in this study too, that the nucleoside pairs located on the 3'- and 5'-ends of ds-DNA adopted a deformed spatial structure on account of the lack of stacking interaction from one of the sides. Their inclusion in the discussion can obscure the clear and correct view of DNA electronic properties as well as charge transfer.

2.1. Structural Analysis of Isolated, Tandem, and Closed DNA Damage

The stability of the double helix depends on three factors: The hydrogen bond (HB) energies between complementary bases, the stacking energy within the base pair (BP) dimers, and solvation (first shape water layer) [40]. Although the mutual BP geometry is rather rigid and sensitive to structural changes (for example, crosslink, alkylation, loss of bases, or part of the aromatic ring), the global spatial geometry of the double helix is to a greater or lesser extent similar. This phenomenon is derived from the high flexibility of the sugar-phosphate backbone, which keeps bases together in the oligonucleotide strands and prevents them from being scattered. Although the helix spine was taken for geometry optimization, due to its lack of significant meaning for hole transfer and electronic properties, it was removed and is not discussed further. The geometry analysis elucidated that G\(^{\text{oxo}}\) (the isolated lesion) appearing in the investigated double helix structures causes negligible h-rise parameter changes in comparison to the unmodified oligo, independently of its relative position to central A3. The h-rise...
parameters are presented in Table 2. The situation is different for the tandem lesion cdAs. These lesions forced h-rise increases, equal for both diastereomers, for the base pair dimers located on the 5′-end and 3′-end direction determined by cdA3. Subsequently, h-rise decreases between cdA4 and G4 were observed; however, a higher value was noted for (5′S)-cdA3 than (5′R)-cdA3, which can predict a different influence on the charge transfer process. These observations show that the rigidity of cdA cannot be eliminated by the sugar-phosphate backbone geometry rearmament in comparison to G\textsuperscript{oxo}.

### Table 2. The h-rise parameters of base pair dimers obtained for the discussed ds-oligonucleotides in their neutral (Neut.) and adiabatic radical cation (ARC) form geometries.

| ds-DNA Base Dimer h-rise Parameter [Å] | 3G\textsuperscript{oxo}-N-DNA | 5G\textsuperscript{oxo}-N-DNA |
|----------------------------------------|-------------------------------|-------------------------------|
| N-DNA                                  | Neut. | ARC | Neut. | ARC | Neut. | ARC | Neut. | ARC |
| G\textsubscript{2A}\textsubscript{3}   | 2.96  | 3.01 | G\textsubscript{2A}\textsubscript{3}   | 3.06  | 3.02 | o\textsuperscript{oxo}G\textsubscript{2A}\textsubscript{3} | 2.89  | 2.86 |
| A\textsubscript{3}G\textsubscript{4}   | 3.31  | 2.88 | A\textsubscript{3}G\textsubscript{4}\textsuperscript{oxo} | 3.33  | 3.25 | A\textsubscript{3}G\textsubscript{4} | 3.24  | 3.23 |
| G\textsubscript{4}G\textsubscript{5}   | 3.34  | 3.14 | o\textsuperscript{oxo}G\textsubscript{4}G\textsubscript{5} | 3.28  | 3.07 | G\textsubscript{4}G\textsubscript{5} | 3.34  | 3.38 |
| ScA-DNA                                | G\textsubscript{2}(5′S)cA\textsubscript{3} | 3.36  | 3.29 | G\textsubscript{2}(5′S)cA\textsubscript{3} | 3.36  | 3.39 | o\textsuperscript{oxo}G\textsubscript{2}(5′S)cA\textsubscript{3} | 3.26  | 3.11 |
| 5′S)cA\textsubscript{3}G\textsubscript{4} | 2.98  | 2.82 | (5′S)cA\textsubscript{3}G\textsubscript{4}\textsuperscript{oxo} | 3.05  | 2.87 | (5′S)cA\textsubscript{3}G\textsubscript{4} | 2.94  | 2.97 |
| G\textsubscript{4}G\textsubscript{5}   | 3.68  | 3.56 | o\textsuperscript{oxo}G\textsubscript{4}G\textsubscript{5} | 3.65  | 3.5 | G\textsubscript{4}G\textsubscript{5} | 3.68  | 3.68 |
| RcA-DNA                                | G\textsubscript{2}(5′R)cA\textsubscript{3} | 3.32  | 3.26 | G\textsubscript{2}(5′R)cA\textsubscript{3} | 3.45  | 3.41 | o\textsuperscript{oxo}G\textsubscript{2}(5′R)cA\textsubscript{3} | 3.37  | 3.24 |
| 5′R)cA\textsubscript{3}G\textsubscript{4} | 2.8   | 2.7 | (5′R)cA\textsubscript{3}G\textsubscript{4}\textsuperscript{oxo} | 3.15  | 3.12 | (5′R)cA\textsubscript{3}G\textsubscript{4} | 2.87  | 3.05 |
| G\textsubscript{4}G\textsubscript{5}   | 3.6   | 3.49 | o\textsuperscript{oxo}G\textsubscript{4}G\textsubscript{5} | 3.68  | 3.55 | G\textsubscript{4}G\textsubscript{5} | 3.63  | 3.67 |

The second structural parameter that strongly influences the hole migration process is the mutual base pair spatial arrangement. It is worth noting that the aromatic ring overlap (ARO) can be perceived as the outcome of the tilt, twist, slide, shift, and roll standard DNA reference frame parameters. The following differences between native N-DNA and ds-oligo containing single, tandem or clustered lesions in the aromatic ring overlapping BP dimers were found (Table 3):

(a) The appearance of G\textsuperscript{oxo} in the discussed system leads to ARO decreases in all the investigated BP dimers except G\textsubscript{2}A\textsubscript{3} of 3G\textsuperscript{oxo}-N-DNA. The above indicates by comparison with 5G\textsuperscript{oxo}-N-DNA that G\textsuperscript{oxo} forces BP flipping from the ideal/parent position in its 3′- and 5′-end directions.

(b) The tandem lesion appearing in the ds-oligonucleotide, i.e., R- or S- cdA, far more strongly disrupts the double helix structure than G\textsuperscript{oxo}. Surprisingly, the presence of (5′R) or (5′S) 5′,8-cyclo-2′-deoxyadenosine leads to G\textsubscript{4}G\textsubscript{5} aromatic ring overlapping increases by −1.6 Å. On the other hand, (5′R)-cdA affected ARO in the case of the cA\textsubscript{3}G\textsubscript{4} base pair dimer more strongly than (5′S)-cdA, with the following values found [in Å\textsuperscript{2}]: 0.69 and 1.79 for ScA-DNA and RcA-DNA, respectively. These differences indicate that (5′R)-cdA disrupts more strongly the spatial ds-DNA structure than the opposite diastereomer, and therefore the effect of this difference should be visible in the values of the charge transfer process parameters.

(c) The geometrical analysis of the clustered lesion, composed of cdA and G\textsuperscript{oxo}, reveals that the (5′S)-cdA causes ARO increases independently of the relative G\textsuperscript{oxo} 3′ or 5′-end position of cdA. The situation is different in the case of (5′R)-cdA if G\textsuperscript{oxo} is present at the 3′ hydroxyl group site of cdA with a BPs ARO decrease being observed. Contrary to the above, G\textsuperscript{oxo} shifted to the 5′ site of (5′R)-cdA, causing aromatic ring overlapping increases within the A\textsubscript{3}G\textsubscript{4} base pair dimers. Based on the above, it can be expected that in the case of clustered lesions, (5′R)-cdA should more strongly affect hole migration than the opposite diastereomer.
The thorough stacking energy analysis (see Table 4) between BP involved directly in the dimer structure leads to melting temperature value decreases of 6 °C. The experimental data shows that (5′)-end site of R-cdA (3Goxo-RcA-DNA) leads to opposite results, with an ST energy decrease noted for the G2A3 dimer, while for the remaining two, rises in its value were assigned. The above results strongly indicate that the influence of DNA damage on stacking interaction strongly depends on the oligonucleotide base sequence. However, it should be pointed out that for all the discussed DNA lesions, the ST energy increases in the G4G5 dimer were observed in a range between 0.04 and 2.79 kcal/mol. These observations indicate that the hole transfer between GG is preferred, which is in good agreement with previous theoretical and experimental data [41].

As mentioned above, the stability of the DNA double helix is the result of stacking and hydrogen bond energies. From previous studies, it is known that base modification and DNA damage can strongly affect mutual complementary base interaction and therefore influence HB energy [34,42]. The results discussed below are presented in Table 5. The comparison of ds-DNA containing a single or clustered lesion with native DNA elucidated that the dGoxo appearing in the oligonucleotide causes increases in the HB energy of the dCoxo:dGoxo pair independently of other lesions present in the range between 0.38 and 0.87 kcal/mol. For a single DNA lesion, the HB energy increases in the dCoxo:dGoxo pair were almost the same for 3Goxo-N-DNA and 5Goxo-N-DNA, i.e., 0.46 and 0.54 kcal/mol, respectively. Moreover, for oligonucleotides containing only dGoxo, the HB energies calculated for other base pairs were almost unaffected in comparison with N-DNA, except the A3:T3 base pair of 3Goxo-N-DNA, for which a fall of 0.28 kcal/mol was noted. Opposite results were noted for other ds-DNA with tandem or clustered lesions in all cases other than for dC:dGoxo base pairs, where a decrease in HB energy was observed. The results presented above indicated that clustered or tandem lesions strongly affect the double helix structure and influence its stability. The experimental data shows that (5′)-cdA leads to melting temperature value decreases of 6 °C [43], while dGoxo affected these parameters only negligibly by 2 °C [44] in comparison to unmodified ds-oligo.
Table 4. Stacking energy interaction in kcal/mol within base pairs dimers of the discussed ds-oligonucleotides in their neutral (Neut.) and vertical neutral (after electron adoption by adiabatic radical cation) (VER.N) form geometries.

| ds-DNA Stacking Energy (kcal/mol) | N-DNA | 3Goxo-N-DNA | 5Goxo-N-DNA |
|-----------------------------------|-------|-------------|-------------|
|                                   | NEUT. | VER.N       | NEUT.       | VER.N       | NEUT. | VER.N |
| G₂A₃                             | -14.56 | -14.55      | G₂A₃        | -14.23      | -14.91 | G₂oxo-A₃ | -14.43 | -13.75 |
| A₃G₄                             | -13.59 | -14.88      | A₃G₄        | -14.69      | -14.63 | A₃G₄      | -13.38 | -14.55 |
| G₄G₅                             | -12.05 | -13.39      | G₄oxo-G₅    | -12.86      | -13.03 | G₄G₅      | -12.09 | -12.76 |
| ScA-DNA                          | G₂A₃  | -12.90      | G₂A₃        | -12.90      | -13.10 | oxo G₂A₃ | -12.82 | -12.31 |
| (5'5'S)cA₃G₄                    | -11.25 | -11.20      | (5'5'S)cA₃G₄ | -11.88      | -11.07 | (5'5'S)cA₃G₄ | -11.40 | -11.90 |
| oxoG₄G₅                         | -14.15 | -13.27      | oxoG₄G₅     | -14.68      | -13.86 | oxoG₄G₅  | -14.16 | -14.25 |
| RcA-DNA                          | G₂A₃  | -12.92      | G₂A₃        | -12.92      | -12.86 | oxo G₂A₃ | -13.05 | -10.58 |
| (5'5'R)cA₃G₄                    | -13.39 | -12.52      | (5'5'R)cA₃G₄ | -14.73      | -13.77 | (5'5'R)cA₃G₄ | -13.24 | -13.70 |
| oxoG₄G₅                         | -13.50 | -12.92      | oxoG₄G₅     | -14.84      | -14.28 | oxoG₄G₅  | -13.48 | -14.54 |

Table 5. Hydrogen bond energy in kcal/mol of base pairs included in the structure of the discussed ds-oligonucleotides in their neutral (Neut.) and vertical neutral (after electron adoption by adiabatic radical cation) (VER.N) form: (a) calculated for an ideal base pair model, (b) calculated for base pairs extracted/selected from 2lsf.pdb [45] and (c) 5iv1.pdb [46] structures.

| ds-DNA Hydrogen Bond Energy | N-DNA | 3Goxo-N-DNA | 5Goxo-N-DNA |
|-----------------------------|-------|-------------|-------------|
|                             | NEUT. | VER.N       | NEUT.       | VER.N       | NEUT. | VER.N |
| G₂C₂                        | -17.23 | -18.10      | G₂A₃        | -17.32      | -17.36 | oxoG₂C₂ | -17.69 | -18.16 |
|                             | -17.54 (a) | -14.36 (b) | -10.81 (a) | -9.75 (a) | -8.64 (b) | -10.53 (a) | -10.44 (a) | -10.80 (a) | -10.39 (a) |
| A₃T₃                        | -10.61 | -9.77 (a)   | A₃T₃        | -10.53      | -10.44 | A₃T₃      | -10.80 | -10.39 |
|                             | -10.95 (a) | -8.64 (b) | -10.74 (a) | -9.75 (a) | -8.64 (b) | -10.53 (a) | -10.44 (a) | -10.80 (a) | -10.39 (a) |
| G₄C₄                        | -17.20 | -18.04 (a)  | G₄C₄        | -17.74      | -17.97 | G₄C₄      | -17.26 | -17.14 |
|                             | -17.00 | -18.49 (a)  | oxoG₄C₄     | -18.04 (a)  | -18.49 (a) | oxoG₄C₄ | -17.26 | -17.14 |
|                             | -17.21 | -16.83 (c)  | oxoG₄C₄     | -18.04 (a)  | -18.49 (a) | oxoG₄C₄ | -17.26 | -17.14 |
| ScA-DNA                     | G₂C₂  | -17.30      | G₂C₂        | -17.25      | -17.30 | oxoG₂C₂  | -17.84 | -18.38 |
| (5'5'S)cA₃T₃                | -10.62 | -10.81 (a)  | (5'5'S)cA₃T₃ | -10.54      | -10.61 | (5'5'S)cA₃T₃ | -10.66 | -9.99 |
|                             | -10.98 (a) | -9.77 (a) | -10.54 (a) | -10.61 (a) | -9.77 (a) | -10.54 (a) | -10.61 (a) | -9.77 (a) | -10.54 (a) |
| G₄C₄                        | -17.10 | -17.58      | G₄C₄        | -17.23      | -17.32 | G₄C₄      | -17.12 | -17.04 |
|                             | -17.06 | -17.08      | oxoG₄C₄     | -17.23      | -17.32 | oxoG₄C₄ | -17.08 | -17.06 |
|                             | -17.19 | -17.12      | oxoG₄C₄     | -17.23      | -17.32 | oxoG₄C₄ | -17.08 | -17.06 |
| RcA-DNA                     | G₂C₂  | -16.94      | G₂C₂        | -17.03      | -17.03 | oxoG₂C₂  | -16.82 | -17.50 |
| (5'5'R)cA₃T₃                | -10.60 | -10.49 (a)  | (5'5'R)cA₃T₃ | -10.65      | -10.55 | (5'5'R)cA₃T₃ | -10.65 | -9.88 |
|                             | -10.98 (a) | -9.74 (a) | -10.65 (a) | -10.55 (a) | -9.74 (a) | -10.65 (a) | -10.55 (a) | -9.74 (a) | -10.55 (a) |
| G₄C₄                        | -16.77 | -18.07      | G₄C₄        | -16.82      | -17.50 | G₄C₄      | -16.82 | -17.50 |
|                             | -16.95 | -16.85      | oxoG₄C₄     | -16.82      | -17.50 | oxoG₄C₄ | -16.96 | -16.80 |
2.2. The Ionization Potential of Isolated, Tandem, and Closed DNA Damage

During the genome one-electron oxidation process initiated by, for example, exposure to ionization radiation, radical cations can be formed randomly. The holes (radical cations) can migrate and become trapped at some preferred places of the oligonucleotide structure with the lowest ionization potential. The following order of nucleic base ionization potential (IP) has been noted: thymine ≈ cytosine > adenine > guanine; additionally, the following radical distribution has been noted during oligonucleotide γ-radiation: 35% G**, 5% A**, and about 45% of T** and C** [47]. Therefore, it can be concluded that pyrimidines have a higher ionization potential than purines. Moreover, 8-oxo-2'-deoxyguanine has a lower ionization potential than dG and is easiest to oxidize [48]. The above results are in good agreement with those presented in this article. Table 6 presents the adiabatic/vertical ionization potential of the isolated base, with base pairs calculated at the M062x/6-31+G** level of theory in the aqueous phase. The situation is a little bit more complicated when ds-DNA is taken into consideration. Independently, Senthilkumar and Voityuk calculated the vertical ionization potential (VIP) of all double-stranded tetramers [49,50]. From their study it is clear that the VIP of the tetramers depends on their sequences. In these studies, both the adiabatic and vertical IP of trimers contained within the tetramer structures (extracted as a central part of optimized ds-hexamers: Figure 2), as well as ds-dimers and isolated base pairs, were taken into theoretical investigation. In all the investigated ds-trimers, the lowest VIP and AIP values were found for (3Goxo-N-DNA) A3oxoG4G5 (5.37/5.79 eV), which are lower by 0.06 and 0.14 eV, respectively, than those noted for oxoG2A3G4. It is important to mention that the lack of oxidized guanosine in the structure eliminates the difference between the vertical and adiabatic state as observed for A3G4G5 extracted from 5Goxo-N-DNA. The VIP of this trimer was found at the same level as that of the corresponding one in N-DNA (VIP = 6.02 eV). As presented in Table 6, the same pattern of VIP and AIP was observed for the discussed tandem and clustered DNA lesions. It is important to mention that the presence of both cdA diastereomers causes slight ionization potential increases in clustered and tandem DNA damage in comparison to the corresponding trimers of native or single-lesioned ds-oligo in the following range: VIP: 0.01–0.12 eV and AIP: 0.02–0.05 eV. Based on the above, it can be postulated that Goxo is a crucial factor, which determines the sink of radical cations and is able to cover the cdA structural influence on the electronic properties of ds-trimers.

Figure 2. Graphical representation of ds-oligonucleotides divided into two ds-trimers, three ds-dimers, and four base pairs (indicated by dashed squares).
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Table 6. Electronic properties in eV: Vertical (VIP) and adiabatic ionization potential (AIP) of the discussed double-stranded trimers, dimers, as well as single base pairs isolated from their parent ds-tetramers, calculated at the M06-2x/6-31+G** level of theory in the aqueous phase.

|                | AIP   | VIP   | AIP   | VIP   | AIP   |
|----------------|-------|-------|-------|-------|-------|
| **ds-trimers** |       |       |       |       |       |
| N-DNA          |       |       |       |       |       |
| G2A3G4         | 5.72  | 6.10  | 5.51  | 5.85  | 5.45  | 5.90  |
| A1G3G5         | 5.64  | 6.03  | 5.37  | 5.79  | 6.01  | 6.02  |
| ScA-DNA        |       |       |       |       |       |
| G2(5'S)cAcG4  | 5.74  | 6.13  | 5.51  | 5.91  | 5.44  | 5.86  |
| (5'S)cAcG4    | 5.69  | 6.08  | 5.45  | 5.88  | 6.09  | 6.14  |
| RcA-DNA        |       |       |       |       |       |
| G2(5'R)cAcG4  | 5.72  | 6.08  | 5.50  | 5.94  | 5.43  | 5.87  |
| (5'R)cAcG4    | 5.66  | 6.03  | 5.47  | 5.91  | 6.13  | 6.03  |
| **ds-dimers**  |       |       |       |       |       |
| N-DNA          |       |       |       |       |       |
| G2A3           | 6.13  | 6.15  | 6.13  | 6.12  | 5.50  | 5.91  |
| A1G4           | 5.73  | 6.12  | 5.48  | 5.88  | 6.12  | 6.11  |
| G4G5           | 5.68  | 6.05  | 5.40  | 5.83  | 6.04  | 6.05  |
| ScA-DNA        |       |       |       |       |       |
| G2(5'S)cAcG4  | 5.75  | 6.14  | 5.52  | 5.93  | 6.20  | 6.15  |
| (5'S)cAcG4    | 5.78  | 6.12  | 5.48  | 5.89  | 6.13  | 6.13  |
| RcA-DNA        |       |       |       |       |       |
| G2(5'R)cAcG4  | 5.74  | 6.09  | 5.51  | 5.94  | 6.13  | 6.10  |
| (5'R)cAcG4    | 5.73  | 6.21  | 5.49  | 5.93  | 6.21  | 6.21  |
| **Single base pairs** |       |       |       |       |       |
| N-DNA          |       |       |       |       |       |
| G2C3           | 6.17  | 6.17  | 6.19  | 6.19  | 5.55  | 5.93  |
| A1T3           | 6.63  | 6.65  | 6.69  | 6.65  | 6.65  | 6.64  |
| G4C4           | 5.86  | 6.13  | 5.55  | 5.91  | 6.14  | 6.13  |
| G5C5           | 6.15  | 6.20  | 6.14  | 6.18  | 6.19  | 6.20  |
| ScA-DNA        |       |       |       |       |       |
| G2(5'S)cAcT3  | 6.65  | 6.68  | 6.67  | 6.68  | 6.83  | 6.69  |
| (5'S)cAcT3    | 5.84  | 6.14  | 5.55  | 5.94  | 6.19  | 6.18  |
| G5C3           | 6.19  | 6.22  | 6.19  | 6.22  | 6.22  | 6.23  |
| RcA-DNA        |       |       |       |       |       |
| G2(5'R)cAcT3  | 6.68  | 6.61  | 6.65  | 6.61  | 6.77  | 6.61  |
| (5'R)cAcT3    | 5.80  | 6.12  | 5.53  | 5.96  | 6.08  | 6.20  |
| G5C3           | 6.18  | 6.22  | 6.15  | 6.19  | 6.19  | 6.22  |
| **Ideal Base Pair Model** |       |       |       |       |       |
| G::C           | 5.58  |       |       | 6.13  |       |       |
| G::C'          | 5.90  |       |       | 6.14  |       |       |
| ScA::T         | 6.23  | 6.62  |       | 6.82  | 6.71  | 6.66  |
| Rca::A         | 6.35  | 6.62  |       | 6.66  | 6.66  | 6.66  |

Isolated from 5iv1pdb [43] and 21sf pdb [44] Structure
The investigated ds-tetramers divided into three base pairs ds-dimers show that in the structure of native ds-DNA, the lowest VIP and AIP were noted for G4G5 (6.05/5.68eV), which was expected. Only negligible differences between the adiabatic and vertical ionization potential were found for G2A3 moiety. The situation was similar in the case of 3G**oxo-N-DNA (single-lesioned ds-oligo), in which the **oxoG3G4 part becomes a hole trap. The 8-oxo-2’-deoxyguanosine shift into the G2 position changes the pattern of IP distribution. The lowest VIP and AIP was noted for **oxoG2A3 ds-dimer (5.91/5.50 eV), while for other ds-dimers, a difference between IPs was not observed. The corresponding results were found for clustered DNA damage in which A3 was changed by (5'R)- or (5'S)-cdA. It should be pointed out that in all the above-discussed cases, the lowest calculated value of the vertical ionization potential, among the isolated dimers, corresponds to the lowest adiabatic IP. The situation is the opposite in the case of the tandem lesion: A discrepancy between VIP and AIP was noted. The lowest VIP was found for the G2cA3 ds-dimer of ScA-DNA and the cA3G4 of RcA-DNA, while the lowest AIP was calculated for A3G4 and G3G5, respectively (Table 6). These observations indicate that cdA appearing in the double helix leads to structural changes, which can obscure the charge migration process. As focused on in Table 3, the geometry rearrangement and its energetical pronunciation (Tables 4 and 5) are more visible after one-electron oxidation. To confirm the above results, parent ds-tetramers were divided into four single base pairs for which the VIP and AIP were calculated at the M062x/6-31+G** level of theory in the aqueous phase. The obtained results show that in the case of dG**oxo being absent, the lowest VIP and AIP were assigned for G4C4 BP independently of which ds-tetramer (native, single, tandem, or cluster lesioned) was isolated. As can be expected for the rest of the discussed ds-oligos, the lowest value of vertical and adiabatic IP was found for base pairs that contained **oxoG in their moiety. Moreover, almost the same values of these parameters were noted for G4C4 and **oxoG3C3: 6.1/5.8eV and 5.9/5.5 eV VIP /AIP, respectively. These results are in good agreement with the experimental data, which shows that the 5’-end GC pair in the d[GG]*d[CC] dimer is the most easily oxidized (due to it having the lowest VIP and AIP) [51]. Additionally, in each discussed case, **oxoGC BP had a lower VIP and AIP by 0.3 eV than the parent GC pair as shown in Table 6. For the remaining base pairs, the assigned IP values fluctuated. However, what is surprising is that the VIP was mainly noted as lower or at the same level as the adiabatic IP. Based on an ionization potential and structural analysis, it can be concluded that the hole migrated through ds-DNA without each BP structural rearrangement, which is necessary for the VIP→AIP conversion. Therefore, the hole slides through the double helix until it settles in the “pleasant” part of ds-oligo, thanks to it having the lowest VIP and AIP.

A comparative spatial geometry analysis of the discussed ds-tetramer between their initial geometry of adiabatic neutral and positively charged states shows that native N-DNA and 5G**oxo-RcA-DNA are the most sensitive to adiabatic radical cation formation (Table 7). In other cases, the presence of dG**oxo or cdA eliminate the structural changes forced by electron loss of ds-oligo. It can be predicted that DNA damage formation makes the hole transfer process much easier towards the radical cation sink formed by dG**oxo than in the case of unmodified ds-oligo, which required significant double helix changes for positive charge compensation. The rigidity of (5'R/5'S)-cdA (tandem lesion) makes the ds-DNA structure resistant to positive ionization. Additionally, the appearance of clustered damage formed by cdA and dG**oxo in the case of 5G**oxo-RcA-DNA leads to significant geometry changes (Table 7) in comparison to others.

In the ionization potential analysis presented above, differences between the discussed ds-oligo were observed forcing the comprehensive charge and spin analysis, presented in Table 8. As expected, independently of the type of ds-oligo, whether undamaged, isolated, tandem, or clustered lesioned, the charge and spin are mainly located on G**oxo or the 5’G4 of the G4G5 dimer in each case. Moreover, the difference between the vertical and adiabatic radical cation form of ds-DNA was negligible. These observations confirm the results (ionization potential) discussed above, which indicate that independently of the system after a complete dismantle of ds-DNAs into constituent base pairs, G**oxo or 5’G4 of G4G5 can be considered a suitable part of the double helix for positive charge accumulation.
Table 7. RMSD (root-mean-square deviation) in Å of the atomic positions calculated for ds-DNAs in neutral and cation radical forms.

| ds-oligo          | Backbone | Bases | All Nucleic Acid |
|-------------------|----------|-------|------------------|
| N-DNA             | 1.347    | 1.039 | 1.197            |
| \(3G^{\text{oxo}}\)-N-DNA | 0.299    | 0.225 | 0.261            |
| \(5G^{\text{oxo}}\)-N-DNA | 0.359    | 0.319 | 0.337            |
| ScA-DNA           | 0.699    | 0.193 | 0.516            |
| \(3G^{\text{oxo}}\)-ScA-DNA | 0.203    | 0.176 | 0.190            |
| \(5G^{\text{oxo}}\)-ScA-DNA | 0.247    | 0.214 | 0.231            |
| RcA-DNA           | 0.695    | 0.298 | 0.537            |
| \(3G^{\text{oxo}}\)-RcA-DNA | 0.119    | 0.082 | 0.103            |
| \(5G^{\text{oxo}}\)-RcA-DNA | 1.032    | 0.817 | 0.931            |

As mentioned above, the charge transfer through the double helix independently of the damage type can be described as a super-exchange or multistep hopping process [28,34]. Using the previously described strategy. the barrier (\(\Delta G\)) for hole transfer within interlaced trimers was assigned in vertical and adiabatic modes (Figure 2, Table 9) [42]. It was found that in all the discussed ds-oligonucleotides. the “hole” appearing in the double-helix structure preferably migrated to \(G_4\) or \(G^{\text{oxo}}\) independently of its position in the ds-DNA. These results are in excellent agreement with the experimental data obtained by Schuster [41].

The charge transfer migration through the double helix can be described according to the Marcus theory, which states that the rate constant (\(k_{\text{ET}}\)) of charge transfer (CT) depends on several factors: The structure of \(\pi\)-stacks, i.e., BP's. the driving force (\(\Delta F\)). nuclear reorganization (\(\lambda\)). activation (\(E_a\)) and the electron-coupling (\(V_{12}\)) energies [52]. \(V_{12}\) was calculated according to the GMH (generalized Mulliken–Hush) strategy within the terms of the occupied Kohn–Sham orbital method [53,54]. The charge transfer, which passes through the adiabatic states of the donor and acceptor, is associated with the movement of internal geometries (atoms), expressed by \(\lambda\) in the Marcus theory. All the above parameters calculated for the systems discussed in this article are presented in Table 10.

An analysis of the reorganization energies reveals a significant rise in the \(A_3G_4\) \(G_4G_5\) dimer in the case of native unmodified N-DAN and for cdA\(_3\)G\(_4\). G\(_4\)G\(_4\) of ds-DNA containing a tandem lesion. Moreover. the same was noted when \(G_4\) was converted into \(G^{\text{oxo}}\) (\(3G^{\text{oxo}}\)-N-DNA, \(3G^{\text{oxo}}\)-RcA-DNA, \(3G^{\text{oxo}}\)-ScA-DNA). It is important to mention that in the case of damage being present in the double helix. the reorganization energy of the \(A_3G_4\) dimer is almost equal to that found for \(G_4G_5\), while for unmodified ds-oligo (N-DNA). the \(\lambda\) of \(G_4G_5\) was two times higher than for \(A_3G_4\). For the oligonucleotides where \(G^{\text{oxo}}\) changed to the \(G_2\) position. the highest \(\lambda\) was denoted for the \(G_2^{\text{oxo}}A_3\) dimer (approximately 0.40 eV) while for the remaining dimers. the value was significantly lower (0.01–0.04 eV). However, for \(5G^{\text{oxo}}\)-RcA-DNA. the \(\lambda\) of \(A_3G_4\) and \(G_4G_5\) should be noted as follows: 0.16 and 0.12 eV respectively. This strongly indicates that \(G^{\text{oxo}}\) plays an invaluable role in genome protection. taking the role of radical slope/trash instead of both diastereomers of cdA (tandem lesion). Due to the fact that \(k_{\text{ET}}\) is strongly dependent on the distance and aromatic ring overlapping between the donor and acceptor, an influence of the single, tandem, and clustered DNA lesion on charge transfer in the double helix shape can be expected in comparison with unmodified ds-DAN. Table 9 presents the discussed parameters of the charge transfer process.
Table 8. Hirshfeld charge (Q) and spin (S) in [au] distribution in the shape of ds-oligonucleotides only nucleosides bases were taken into consideration, calculated at the M062X(D95°)//M062x/6-31+G** level of theory in the aqueous phase. A—neutral, VC—vertical cation, C—adiabatic cation.

|        | ds-DNA Hirshfeld Charge and Spin Density Population | 3G<sub>oxo</sub>-N-DNA | 5G<sub>oxo</sub>-N-DNA |
|--------|---------------------------------------------------|-------------------------|-------------------------|
|        | N-DNA                                             | 3G<sub>oxo</sub>-N-DNA  | 5G<sub>oxo</sub>-N-DNA  |
|        | N        | VC       | C        | N        | VC       | C        | N        | VC       | C        | N        | VC       | C        |
|        | Q        | Q        | S        | Q        | Q        | S        | Q        | Q        | S        | Q        | Q        | S        |
| T<sub>6</sub> | -0.05   | -0.05   | -0.04   | T<sub>6</sub> | -0.05 | -0.05 | -0.04 | T<sub>6</sub> | -0.05 | -0.05 | -0.05 |
| C<sub>5</sub> | 0.18    | 0.19    | 0.18    | C<sub>5</sub> | 0.17 | 0.17 | 0.17 | C<sub>5</sub> | 0.18 | 0.18 | 0.17 |
| C<sub>4</sub> | 0.19    | 0.22    | 0.32    | C<sub>4</sub> | 0.21 | 0.24 | 0.29 | C<sub>4</sub> | 0.19 | 0.19 | 0.20 |
| T<sub>3</sub> | -0.07   | -0.05   | -0.02   | T<sub>3</sub> | -0.07 | -0.05 | -0.05 | T<sub>3</sub> | -0.06 | -0.06 | -0.05 |
| C<sub>2</sub> | 0.21    | 0.22    | 0.22    | C<sub>2</sub> | 0.20 | 0.21 | 0.21 | C<sub>2</sub> | 0.21 | 0.25 | 0.31 |
| T<sub>1</sub> | -0.08   | -0.07   | -0.07   | T<sub>1</sub> | -0.08 | -0.07 | -0.07 | T<sub>1</sub> | -0.08 | -0.05 | -0.02 |
| A<sub>1</sub> | 0.01    | 0.01    | 0.00    | A<sub>1</sub> | 0.01 | 0.01 | 0.01 | A<sub>1</sub> | 0.02 | 0.06 | 0.01 | 0.02 |
| G<sub>2</sub> | -0.14   | -0.13   | -0.14   | G<sub>2</sub> | -0.14 | -0.13 | -0.13 | G<sub>2</sub> | -0.16 | 0.69 | 0.97 | 0.59 | 0.97 |
| A<sub>3</sub> | 0.03    | 0.06    | 0.02    | A<sub>3</sub> | 0.03 | 0.07 | 0.02 | A<sub>3</sub> | 0.03 | 0.06 | 0.02 | 0.05 | 0.01 |
| G<sub>4</sub> | -0.18   | 0.67    | 0.96    | G<sub>4</sub> | 0.97 | 0.59 | 0.97 | G<sub>4</sub> | -0.18 | -0.17 | -0.16 |
| G<sub>5</sub> | -0.14   | -0.10   | 0.02    | -0.09 | 0.02 | G<sub>5</sub> | -0.13 | -0.10 | 0.01 | -0.10 | 0.01 | G<sub>5</sub> | -0.14 | -0.14 | -0.13 |
| A<sub>6</sub> | 0.03    | 0.04    | 0.03    | A<sub>6</sub> | 0.04 | 0.04 | 0.05 | A<sub>6</sub> | 0.04 | 0.04 | 0.04 |
|        | ScA-DNA  | 3G<sub>oxo</sub>-ScA-DNA | 5G<sub>oxo</sub>-ScA-DNA |
| T<sub>6</sub> | -0.06   | -0.06   | -0.05   | T<sub>6</sub> | -0.06 | -0.06 | -0.05 | T<sub>6</sub> | -0.06 | -0.06 | -0.06 |
| C<sub>5</sub> | 0.21    | 0.22    | 0.22    | C<sub>5</sub> | 0.21 | 0.22 | 0.22 | C<sub>5</sub> | 0.21 | 0.21 | 0.22 |
| C<sub>4</sub> | 0.13    | 0.16    | 0.23    | C<sub>4</sub> | 0.13 | 0.16 | 0.23 | C<sub>4</sub> | 0.12 | 0.12 | 0.13 |
| T<sub>3</sub> | -0.08   | -0.07   | -0.06   | T<sub>3</sub> | -0.08 | -0.07 | -0.07 | T<sub>3</sub> | -0.08 | -0.07 | -0.07 |
| C<sub>2</sub> | 0.22    | 0.22    | 0.23    | C<sub>2</sub> | 0.22 | 0.22 | 0.23 | C<sub>2</sub> | 0.22 | 0.25 | 0.31 |
| T<sub>1</sub> | -0.07   | -0.07   | -0.05   | T<sub>1</sub> | -0.06 | -0.06 | -0.06 | T<sub>1</sub> | -0.05 | -0.03 | -0.01 |
| A<sub>1</sub> | 0.00    | 0.00    | 0.00    | A<sub>1</sub> | -0.01 | -0.01 | -0.01 | A<sub>1</sub> | 0.06 | 0.11 | 0.02 | 0.10 |
| G<sub>2</sub> | -0.16   | -0.16   | -0.15   | G<sub>2</sub> | -0.16 | -0.16 | -0.14 | G<sub>2</sub> | -0.24 | 0.63 | 0.97 | 0.53 | 0.97 |
| ScA<sub>3</sub> | 0.04    | 0.08    | 0.03    | ScA<sub>3</sub> | 0.03 | 0.07 | 0.03 | ScA<sub>3</sub> | 0.03 | 0.06 | 0.01 | 0.06 | 0.03 |
| G<sub>4</sub> | -0.15   | 0.72    | 0.96    | G<sub>4</sub> | -0.16 | 0.71 | 0.96 | G<sub>4</sub> | -0.14 | -0.14 | -0.13 |
Table 8. Cont.

|          | ds-DNA Hirshfeld Charge | Spin Density Population |
|----------|-------------------------|-------------------------|
| G5       | -0.11                   | -0.08                   |
| A6       | 0.03                    | 0.04                    |
|          |                         |                         |
| A6       | 0.03                    | 0.04                    |
|          |                         |                         |
| Rca-DNA  | 0.05                    | 0.05                    |
|          |                         |                         |
| C5       | -0.06                   | -0.04                   |
|          |                         |                         |
| C4       | 0.13                    | 0.16                    |
|          |                         |                         |
| T3       | -0.03                   | -0.03                   |
|          |                         |                         |
| C2       | 0.22                    | 0.23                    |
|          |                         |                         |
| T1       | -0.02                   | -0.02                   |
|          |                         |                         |
| A1       | 0.05                    | 0.15                    |
|          |                         |                         |
| G4       | -0.17                   | -0.17                   |
|          |                         |                         |
| G5       | -0.12                   | -0.12                   |
|          |                         |                         |
| A6       | 0.03                    | 0.04                    |
Table 9. Energy barriers (in eV) for radical cation migration between base pairs within trimers. The vertical modes, i.e., the energies of each base pair’s radical cation, were calculated for their neutral geometry. The adiabatic modes, i.e., the energies of each base pair’s radical cation, were calculated for their cation geometry. Arrows indicate the direction of the hole transfer from one base pair to another, e.g., A$^+$ → G, calculated at the M062x/6-31+G** level of theory in the aqueous phase [42].

| ds-oligo | Mode     | Discussed Trimmers |         |         |         |         |         |         |         |         |         |         |
|----------|----------|--------------------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
|          |          | G2→A3 G4 A3→G4 G2→G4 A3→G4 G4→G5 A3→G5 |         |         |         |         |         |         |         |         |         |         |
| N-DNA    | Vertical | 0.49              | −0.49   | −0.03   | −0.49   | 0.73    | −0.04   |         |         |         |         |         |
|          | Adiabatic| 0.46              | −0.77   | −0.31   | −0.77   | 0.29    | −0.48   |         |         |         |         |         |
|          | G2→A3   | A3→G4             | G2→G4   | A3→G4   | G4→G5   | A3→G5   |         |         |         |         |         |         |
|          | Vertical | −0.46             | 1.18    | 0.31    | 1.18    | −0.01   | 0.89    |         |         |         |         |         |
|          | Adiabatic| −0.46             | 0.77    | 0.31    | 0.77    | −0.29   | 0.48    |         |         |         |         |         |
| 3Goxo-N-DNA | Vertical | G2→A3             | A3→oxoG4| G2→oxoG4| A3→oxoG4| oxoG4→G5| A3→G5   |         |         |         |         |         |
|          | Adiabatic| 0.50              | −1.14   | −0.64   | −1.14   | 0.59    | −0.54   |         |         |         |         |         |
|          | G2→A3   | A3→oxoG4          | G2→oxoG4| A3→oxoG4| oxoG4→G5| A3→G5   |         |         |         |         |         |         |
|          | Vertical | −0.49             | 1.44    | 0.64    | 1.44    | −0.23   | 0.85    |         |         |         |         |         |
|          | Adiabatic| −0.50             | 1.14    | 0.64    | 1.14    | −0.59   | 0.54    |         |         |         |         |         |
| 5Goxo-N-DNA | Vertical | oxoG2→A3          | A3→G4   | oxoG2→G4| A3→G4   | G4→G5   | A3→G5   |         |         |         |         |         |
|          | Adiabatic| 1.45              | −0.50   | 0.61    | −0.50   | 0.06    | −0.45   |         |         |         |         |         |
|          | oxoG2→A3| A3→oxoG4         | oxoG2→G4| A3→oxoG4| oxoG4→G5| A3→G5   |         |         |         |         |         |         |
|          | Vertical | −0.69             | 0.50    | −0.18   | 0.50    | −0.05   | 0.46    |         |         |         |         |         |
|          | Adiabatic| −1.10             | 0.51    | −0.59   | 0.51    | −0.05   | 0.46    |         |         |         |         |         |
| ScA-DNA  | Vertical | G2→ScA3           | ScA3→G4 | G2→ScA3 | ScA3→G4 | G4→G5   | ScA3→G5 |         |         |         |         |         |
|          | Adiabatic| 0.55              | −0.51   | 0.02    | −0.51   | 0.71    | −0.11   |         |         |         |         |         |
|          | G2→ScA3 | ScA3→G4          | G2→ScA3 | ScA3→G4 | G4→G5   | ScA3→G5 |         |         |         |         |         |         |
|          | Vertical | −0.46             | 1.16    | 0.35    | 1.16    | −0.05   | 0.82    |         |         |         |         |         |
|          | Adiabatic| −0.52             | 0.82    | 0.29    | 0.82    | −0.35   | 0.47    |         |         |         |         |         |
| ds-oligo | Mode      | Discussed Trimmers |
|----------|-----------|--------------------|
|          |           | G2→A3 | A3→G4 | G2→G4 | A3→G4 | G4→G5 | A5→G5 | A3→G3 |
| 3Goxo-ScA-DNA | Vertical | 0.55  | -0.73 | -0.20 | -0.73 | 1.01  | -0.11 | |
|           | Adiabatic | 0.53  | -1.12 | -0.59 | -1.12 | 0.64  | -0.48 | |
| 5Goxo-ScA-DNA | Vertical | 0.53  | -0.50 | 0.55  | -0.88 | 0.38  | -0.50 | |
|           | Adiabatic | 0.53  | 1.12  | 0.59  | -0.64 | 0.48  | -0.64 | |
| RcA-DNA  | Vertical | 0.49  | -0.52 | 0.02  | -0.52 | 0.69  | -0.18 | |
|           | Adiabatic | 0.55  | -0.88 | -0.33 | -0.88 | 0.38  | -0.50 | |
| 3Goxo-RcA-DNA | Vertical | 0.45  | -0.69 | -0.19 | -0.69 | 0.96  | -0.16 | |
|           | Adiabatic | 0.50  | -1.12 | -0.63 | -1.12 | 0.62  | -0.50 | |
| 5Goxo-RcA-DNA | Vertical | 1.36  | -0.53 | 0.67  | -0.53 | 0.08  | -0.61 | |
|           | Adiabatic | 1.19  | -0.53 | 0.47  | -0.53 | 0.08  | -0.61 | |

Table 9. Cont.
Table 10. Nuclear relaxation energy $\lambda$ [eV] and hole transfer rate constant $k_{HT}$ [s$^{-1}$], energy barrier $\Delta G$ [eV], activation energy $E_a$ [eV], and electron coupling energies $V_{12}$ [eV] of hole transfer between base pairs, calculated at the M062x/6-31+G** level of theory in the aqueous phase ([42] and references cited therein).

| System | $\lambda$ | $\Delta G$ | $E_a$ | $V_{12}$ | $k_{HT}$ | $\lambda$ | $\Delta G$ | $E_a$ | $V_{12}$ | $k_{HT}$ |
|--------|------------|-------------|-------|--------|--------|------------|-------------|-------|--------|--------|
| N-DNA  |            |             |       |        |        |            |             |       |        |        |
| $G_2\leftarrow A_3$ | 0.00 | -0.46 | 18.60 | 0.221 | 0.00 | $G_2\leftarrow A_3$ | 0.01 | -0.50 | 10.09 | 0.220 | 0.00 |
| $A_3\rightarrow G_4$ | 0.28 | -0.77 | 0.22 | 0.246 | $3.8 \times 10^{11}$ | $A_3\rightarrow 3G_{oxo}G_4$ | 0.37 | -1.14 | 0.41 | 0.363 | $5.2 \times 10^8$ | $A_3\rightarrow G_4$ | 0.01 | -0.51 | 4.31 | 0.246 | 0.00 |
| $G_{4}\leftarrow G_5$ | 0.44 | -0.29 | 0.01 | 0.051 | $4.0 \times 10^{13}$ | $G_{4}\leftarrow 3G_{oxo}G_5$ | 0.38 | -0.59 | 0.03 | 0.113 | $1.1 \times 10^{14}$ | $G_{4}\leftarrow G_5$ | 0.02 | -0.05 | 0.01 | 0.048 | $1.7 \times 10^{14}$ |
| ScA-DNA |            |             |       |        |        |            |             |       |        |        |
| $G_2\leftarrow ScA_3$ | 0.06 | -0.52 | 0.86 | 0.263 | 14.58 | $G_2\leftarrow cSA_3$ | 0.02 | -0.53 | 4.30 | 0.271 | 0.00 |
| $ScA_3\rightarrow G_4$ | 0.31 | -0.82 | 0.21 | 0.264 | $6.4 \times 10^{13}$ | $ScA_3\rightarrow 3G_{oxo}G_4$ | 0.39 | -1.12 | 0.35 | 0.378 | $5.5 \times 10^8$ | $ScA_3\rightarrow G_4$ | 0.04 | -0.64 | 2.62 | 0.271 | 0.00 |
| $G_{4}\leftarrow G_5$ | 0.36 | -0.35 | 0.00 | 0.035 | $3.4 \times 10^{13}$ | $G_{4}\leftarrow 3G_{oxo}G_5$ | 0.37 | -0.64 | 0.05 | 0.157 | $1.0 \times 10^{14}$ | $G_{4}\leftarrow G_5$ | 0.01 | -0.03 | 0.01 | 0.038 | $1.5 \times 10^{14}$ |
| RcA-DNA |            |             |       |        |        |            |             |       |        |        |
| $G_2\leftarrow RcA_3$ | 0.13 | -0.55 | 0.33 | 0.260 | $9.1 \times 10^9$ | $G_2\leftarrow RcA_3$ | 0.03 | -0.50 | 1.91 | 0.268 | 0.00 |
| $RcA_3\rightarrow G_4$ | 0.35 | -0.88 | 0.19 | 0.297 | $1.3 \times 10^{12}$ | $RcA_3\rightarrow 3G_{oxo}G_4$ | 0.43 | -1.12 | 0.27 | 0.349 | $7.8 \times 10^{10}$ | $RcA_3\rightarrow G_4$ | 0.16 | -0.69 | 0.43 | 0.291 | $1.75 \times 10^8$ |
| $G_{4}\leftarrow G_5$ | 0.31 | -0.38 | 0.00 | 0.086 | $1.9 \times 10^{14}$ | $G_{4}\leftarrow 3G_{oxo}G_5$ | 0.34 | -0.62 | 0.06 | 0.131 | $5.3 \times 10^{13}$ | $G_{4}\leftarrow G_5$ | 0.12 | -0.11 | 0.00 | 0.085 | $3.5 \times 10^{14}$ |
which the directional e
with the position of all other atoms frozen.

This was confirmed by the results obtained for 5G
These observations indicate that the charge transfer within the double helix can be disturbed by
Presented in Table 9. the transfer between G
G values: 0.00 for
atoms added for saturation were optimized at the M06-2X
the atoms were saturated, if necessary, with hydrogen atoms. The spatial location of the hydrogen
group were neutralized by the addition of protons. The full structure optimized
calculation using ONIOM in the aqueous phase [55]. The solvent e
3.1. Computation Methodology of QM/MM Studies [42,43]

The calculated k_{HTS} value between the base pair dimers of the reference ds-oligo gives the following
values: 0.00 for G2→A3, 3.8 × 10^{11} for A3→G4 4.0 × 10^{13}, and G4→G5. The obtained higher value for
G4→G5 is in good agreement with recent theoretical studies, which have postulated that the hole
migrated in the 5′-end direction of GG dimers [32]. The single lesion formation in the double helix
influence on the CT process depends on its place of settlement. The presence of G^{oxo} as part of the
G4G5 dimer (on its 5′-end) leads to a greater CT rate increase by one order of magnitude than for
N-DNA, with subsequent significant k_{HT} decreases for A3→G4 transfer (Table 9). The G^{oxo} shift to
the G2 position causes the CT rate to increase between A3→G2′ in comparison to native DNA up
to 3.2 × 10^{10}. However, the k_{HT} assigned for G4→G5 transfer was at the same level as observed for
N-DNA while ΔG decreases were noted.

The formation of 5′,8-cyclo-2′-deoxyadenosine in the double helix leads to different results
depending on the C5′ chirality. The (5′S)-cdA force the same effect as discussed for a native ds-oligo
(N-DNA), when it has been considered as a tandem lesion. The configuration inversion on the C5′
of cdA forces k_{HTS} increases in all the discussed CTs ((5′R)cA3→G2; (5′R)cA3→G4; G5→G4) (Table 9).
These observations indicate that the charge transfer within the double helix can be disturbed by
structural changes forced by (5′R)-cdA (Tables 2–4). Moreover, based on the energy barrier analysis
presented in Table 9, the transfer between G2→G4 can take place in the adiabatic mode (−0.33 eV).
More details on calculated energy levels can be found in the Supplementary Materials.

The presence of G^{oxo} and (5′R)- or (5′S)-cdA in the same helix turn leads to clustered damage
formation. In the case when G^{oxo} is part of the oxoG4G5 dimer (3G^{oxo}-ScA-DNA and 3G^{oxo}-RcA-DNA),
the charge transfer is allowed for cA3→G4^{oxo} and G5→G4^{oxo}. However, (5′S)-cdA increases the hole
migration from (5′S)cA3→G4 by one order of magnitude in comparison with N-DNA, while for the
opposite diastereomer, this value was two orders of magnitude higher. Subsequently, (5′S)-cdA left
k_{ET} of G5→G4^{oxo} at the same level as was assigned for native N-DNA for (5′R)-cdA; this value was
found to be one order of magnitude lower. The above indicates that depending on the C5′ chirality,
cdA can modulate the charge transfer in its 3′- or 5′-end direction in the case of a clustered DNA lesion.
This was confirmed by the results obtained for 5G^{oxo}-ScA-DNA and 5G^{oxo}-RcA-DNA, where G^{oxo} was
shifted to the G2 position (Figure 2, Table 1). As presented in Table 10, the presence of (5′S)-cdA slows
down the (5′S)cA→G2^{oxo} charge transfer by three orders of magnitude, while (5′R)-cdA is only by two
in comparison with 5G^{oxo}-N-DNA. Subsequently, both cdA diastereomers left G5→G4 at the same
level as forced by dA in suitable single-lesioned DNA. Surprisingly, in the case of 5G^{oxo}-RcA-DNA.
the (5′R)cA3→G4 was found to be allowed/possible—k_{ET} = 1.75 × 10^{8}—and was at the same level as
that assigned for (5′R)cA3→G2^{oxo}, which indicates that the 5′R diastereomer is able to disturb
the charge transfer process. These observations are in good agreement with previous theoretical studies in
which the directional effect of cdAs was noted [34].

3. Materials and Methods

3.1. Computation Methodology of QM/MM Studies [42,43]

The geometry optimizations of ds-hexamers presented in Table 1 were performed using the
QM/MM strategy [35,36]. The structures of the double-stranded oligonucleotides were divided
into high- HL (nucleobases, M06-2X/D95*), and low- LL (sugar-phosphate backbone, UFF) levels of
calculation using ONIOM in the aqueous phase [55]. The solvent effect was described for an aqueous
medium, applying Tomasi’s polarized continuum model [56]. The negative charges of each phosphate
group were neutralized by the addition of protons. The full structure optimized ds-hexamers were
converted to base pairs by sugar-phosphate backbone removal. In the formed base pair systems,
the atoms were saturated, if necessary, with hydrogen atoms. The spatial location of the hydrogen
atoms added for saturation were optimized at the M06-2X/D95* level of theory in the aqueous phase,
with the position of all other atoms frozen.
3.2. Computation Methodology of Density Functional Theory (DFT) Study

All energy calculations were performed in the aqueous phase by the density functional theory (DFT) using the M06-2X functional with a double-ζ 6-31+G** basis set [57,58]. The characterization of the transition dipole moment of excited states and the single point calculation at the M06-2X/6-31+G** level of theory were performed using time-dependent DFT (TD-DFT) methodology [59]. For all the optimized structures, a charge and spin analysis was achieved using Hirshfeld methodology at the M06-2X/6-31+G** level [60]. The electron coupling was calculated using generalized Mulliken–Hush methodology [61]. The electronic properties were calculated as previously described [62]. All calculations were performed in the aqueous phase with the Gaussian 09 (revision A.02) software package [63]. The three-dimensional structural analyses of the mentioned ss- and ds-DNAs, based on a standard reference frame, were obtained with by a 3DNA software package using the web-based interface w3DNA (web 3DNA) [64].

4. Conclusions

The appearance of different types of single, tandem, or clustered DNA lesions in the oligonucleotide sequence gives rise to various consequences of charge transfer in comparison with native ds-oligo (N-DNA). In this article, for the first time, a comparative analysis was made between unmodified ds-oligo and one which contains G\textsuperscript{\textbf{oxo}}, cdA, or both. Both types of lesions taken into consideration can be formed by hydroxyl radical activity. However, the dG\textsuperscript{\textbf{oxo}} by •OH addition to the C8 moiety of dG while the 5′R and 5′S diastereomers of 5′,8-cyclo-2′deoxyadenosine can occur in ds-DNA as a result of a two-step cyclization reaction induced by hydrogen atom abstraction from the C5′ position by a hydrogen radical [65]. These unusual tandem lesions can lead to different local spatial geometry changes in the double helix, next to their place of formation [24]. Probably, as a result, (5′R)-cdA and (5′S)-cdA(S) had a disparate influence on BER enzyme activities, as well as on the electronic properties of the ds-DNA part, next to its appearance. The results presented above indicate that dependent on C5′ chirality, cdA can modulate the charge transfer toward its 3′- or 5′-end direction in the case of a clustered DNA lesion. However, in all the discussed DNA lesions, the appearance of dG\textsuperscript{\textbf{oxo}} in the double helix structure constitutes the final destination of radical cation migration.

Supplementary Materials: The following materials can be online: Table S1. The energies (in Hartree) of Neutral, Vertical Cation, Adiabatic Cation and Vertical Neutral forms of ideal base pairs, base pairs extracted from 3lsf.pdb [1] and 5iv1.pdb [2] files calculated at the M062x/6-31+G** level of theory in the aqueous phase. Table S2. The energies (in Hartree) of Neutral. Vertical Cation. Adiabatic Cation and Vertical Neutral forms of base pairs extracted from ds-oligonucleotides calculated at the M062x/6-31+G** level of theory in the aqueous phase. Table S3. The energies (in Hartree) of Neutral. Vertical Cation. Adiabatic Cation and Vertical Neutral forms of base pairs dimers extracted from ds-oligonucleotides calculated at the M062x/6-31+G** level of theory in the aqueous phase. Table S4. Energies (in Hartree) of Neutral. Vertical Cation. Adiabatic Cation and Vertical Neutral forms of ds-trimers extracted from ds-oligonucleotides calculated at the M062x/6-31+G** level of theory in the aqueous phase. Table S5. The Ground and Excitation state energies and Excitation and HOMO Energies as well as corresponding Dipole Moments (Ground Excitation and Transition) of base pair dimers extracted from ds-oligonucleotides. calculated at the M062x/6-31+G** level of theory in the aqueous phase using the DFT or TD-DFT methodology.

Eighteen pdb files of discussed ds-oligo structures: ScA\_DNA\_Neutral.pdb, ScA\_DNA\_Cation.pdb, RaA\_DNA\_Neutral, RaA\_DNA\_Cation.pdb, N\_DNA\_Neutral.pdb, N\_DNA\_Cation.pdb, 5Goxo\_ScA\_DNA\_Neutral.pdb, 5Goxo\_ScA\_DNA\_Cation.pdb, 5Goxo\_RaA\_DNA\_Neutral.pdb, 5Goxo\_RaA\_DNA\_Cation.pdb, 5Goxo\_N\_DNA\_Neutral.pdb, 5Goxo\_N\_DNA\_Cation.pdb, 3Goxo\_ScA\_DNA\_Neutral.pdb, 3Goxo\_ScA\_DNA\_Cation.pdb, 3Goxo\_RaA\_DNA\_Neutral.pdb, 3Goxo\_RaA\_DNA\_Cation.pdb, 3Goxo\_N\_DNA\_Neutral.pdb, 3Goxo\_N\_DNA\_Cation.pdb.

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Sample Availability: Samples of the compounds are not available from the author.