Understanding DNA under oxidative stress and sensitization: the role of molecular modeling

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DNA is constantly exposed to damaging threats coming from oxidative stress, i.e., from the presence of free radicals and reactive oxygen species. Sensitization from exogenous and endogenous compounds that strongly enhance the frequency of light-induced lesions also plays an important role. The experimental determination of DNA lesions, though a difficult subject, is somehow well established and allows to elucidate even extremely rare DNA lesions. In parallel, molecular modeling has become fundamental to clearly understand the fine mechanisms related to DNA defects induction. Indeed, it offers an unprecedented possibility to get access to an atomistic or even electronic resolution. Ab initio molecular dynamics may also describe the time-evolution of the molecular system and its reactivity. Yet the modeling of DNA (photo-)reactions does necessitate elaborate multi-scale methodologies to tackle a damage induction reactivity that takes place in a complex environment. The double-stranded DNA environment is first characterized by a very high flexibility, but also a strongly inhomogeneous electrostatic embedding. Additionally, one aims at capturing more subtle effects, such as the sequence selectivity which is of critical important for DNA damage. The structure and dynamics of the DNA/sensitizers complexes, as well as the photo-induced electron- and energy-transfer phenomena taking place upon sensitization, should be carefully modeled. Finally the factors inducing different repair ratios for different lesions should also be rationalized. In this review we will critically analyze the different computational strategies used to model DNA lesions. A clear picture of the complex interplay between reactivity and structural factors will be sketched. The use of proper multi-scale modeling leads to the in-depth comprehension of DNA lesions mechanisms and also to the rational design of new chemo-therapeutic agents.

Keywords: DNA, photosensitization, photodynamic therapy, photochemistry, molecular modeling, energy/electron transfer

1. Introduction

DNA stability is essential to maintain cellular integrity of living organisms and avoid genetic mutations. Threats to DNA stability can be triggered by oxidative stress induced by the presence of metabolic reactive radical and oxygen species (ROS) or by photoreactivity. Oxidative and light-induced stress can have a very strong influence on the biological processes governed by DNA, as
well as in the indirect activation or transduction of signal cascades, that may result in different malignant outcomes for the life of the involved cells (Salmon et al., 2004; Kujoth et al., 2005; Klausig et al., 2010). Although, in the following we will focus more on OH• induced pathways, it is worth mentioning that superoxide radical (O2−) is also connected to the induction of DNA lesions (Fridovich, 1995; Cadenas and Davies, 2000; Valko et al., 2007), as well as reactive nitrogen species (RNS) that are known for their potential carcinogenic activity (Pryor and Squadrito, 1995; Pacher et al., 2007; Sainz et al., 2012).

In contrast to photochemical degradation mechanisms, oxidative stress sources can also derive from endogenous processes and mechanism. Inflammatory conditions are a known source of ROS (Kamp et al., 2011), such as OH•, and remarkably ROS and free radicals are also exploited by the cells for signal transduction (Hamanaka and Chandel, 2010; Finkel, 2011), and hence are a key factor of cell metabolism. Moreover, ROS ultimately attacking DNA, may also derive as byproducts of the degradation of oxidized lipids or proteins, via the endocytosis process (Lim et al., 2004; Miyamoto and Di Mascio, 2014). In addition an other important source of endogenous ROS production, also related to neurodegenerative diseases, is due to the metabolism of dietary and biogenic amines (Chaiyen et al., 2012; Ramsay, 2012; Vianello et al., 2012; Repic et al., 2014). Indeed, this metabolic pathway performed by monoamide oxidase enzymes leads to the production of H2O2 and subsequently of OH•.

DNA is also constantly exposed to light, and hence efficient dissipative channels exist to hamper potentially dangerous photochemical pathways. Nevertheless, inherently stable DNA photoproducts have been reported and their photochemistry has been deeply analyzed (Sinha and Hader, 2002; Brash, 2015). Indeed, the correlation between DNA photo-lesions and the emergence of threatening mutations is now well accepted, and in particular the correlation between some types of skin cancers and uncontrolled sun exposure (Sage et al., 2005). Furthermore, the absorption range of native DNA can be significantly extended from the UVB region (where the constitutive nucleobases individually absorb) to the less energetic UVA because of the excitonic coupling within the macromolecular DNA structure, or thanks to the interaction with endogenous or exogenous chromophores, i.e., the photosensitization (Epe, 2012).

The cells response to the oxidative and photochemical stress normally results in an acceleration of protective mechanisms aimed to expel the stress enhancing factors. In the case of DNA lesions the activation of “repair” proteins, that may vary greatly in terms of specificity and efficiency, is invoked. The latter enzymes usually excise the damaged DNA area (Sancar and Sancar, 1988; Radzimanowksi et al., 2013), therefore hampering any mismatch in the replication process. However some situations exist where the repair mechanisms fail to efficiently eradicate the lesions. In this case two scenarios are possible. On the one hand mismatches can occur during transcription, ultimately leading to mutations and eventually to carcinogenesis. On the other hand if the stress is too strong or the replication is impossible, for instance because it is blocked by interstrand cross-links, cells may induce their programmed death (apoptosis).

Apoptosis induction favored by the controlled production of DNA lesions has a very important therapeutic significance. Indeed, the formation of irreversible DNA lesions, and the interference with the replication process, has been and is currently exploited in the development of chemotherapeutic agents. For instance, we may cite the renown case of the cisplatin (Floreà and Büsselberg, 2011), whose therapeutic effects are known since the 60’s. More recently the cytotoxic action of other organometallic compounds, in particular Ruthenium complexes (Rademaker-Lakhai et al., 2004; Yan et al., 2005; Antonarakis and Emadi, 2010; Suss-Fink, 2010), has gained great attention in the development of novel chemotherapeutic drugs that are presently in clinical trial phase. Furthermore, since DNA lesions can be induced not only by ground state chemistry, but may also results from photochemical or photophysical activated energy- or electron-transfer processes the use of light-triggered therapeutic strategies and drugs appears promising in enhancing specificity of action and hence reduce unwanted side-effects. A particular attention should be paid to the light-induced activation of singlet oxygen that is promoted by opportune sensitizers such as porphyrins and lies at the heart of photodynamic therapy strategies (Dougherty et al., 1998; Pandey, 2000; Agostinis et al., 2011; Ethirajan et al., 2011).

From an experimental point of view, one relies on the use of techniques able to identify and possibly quantify the different lesions formed in DNA. Combination of mass-spectrometry with gas-phase chromatography is usually used to provide such informations both for isolated and cellular DNA (Frelon et al., 2000). Indeed, the difficulty in studying the reactions taking place in the complex macromolecular environment usually pushes toward the use of simple models, such as (artificial) mono- or dinucleotide. Even if such simplified techniques allow to infer and extrapolate valuable information on the nucleobase reactivity, still it is important to recall that the role of the DNA environment is almost totally neglected, while it may have huge chemical consequences. On the other hand NMR and X-ray spectrometry may provide extremely useful structures of lesioned DNA double-strand (Gold et al., 2014; Jain et al., 2014; Zalesak et al., 2014; Mutter et al., 2015). Even if in this case most often DNA inherent flexibility and complex dynamic are not properly taken into account, and hence the structural evolution and the role played by different lesions in driving such reorganization may be totally neglected. Obviously, to assess the response of DNA to UV/vis radiation the use of spectroscopic techniques is crucial. This can go from the standard and bench-scale use of optical techniques such as absorption, luminescence and electronic dichroism (Vorlickova and Palecek, 1974; Brabec et al., 1992; Ding et al., 2009) to more sophisticated time-resolved technique (Gustavsson et al., 2010, 2013; Vaya et al., 2012), allowing to follow the time-evolution of the different excited states. Furthermore, dichroism and spectroscopic titration are commonly used to infer the different and possibly competitive interaction modes between DNA and sensitizers (Carvlin et al., 1982; Wang et al., 2012; Lauria et al., 2014).

As this brief survey evidences, the induction of DNA lesions triggered by different sources of stress involves a very complex
interplay between different phenomena taking place at molecular level, and having more general and systemic consequences. On the other hand, it is precisely that complexity that strongly suggests a multidisciplinary approach to tackle this non trivial problem, in particular combining adequate spectroscopic techniques with state-of-the-art molecular multiscale modeling. Molecular modeling is invaluable in providing atomistic or even electronic scale description of the involved phenomena. For instance, molecular dynamics techniques allow to assess for the structural deformation of lesioned DNA and its time evolution, as well as to infer the existence of different stable interaction modes with smaller sensitizers. The ground- and excited-state behavior and the reactivity, for instance in terms of free energy profiles along reaction coordinates, are described by using hybrid quantum mechanics/ molecular mechanics (QM/MM) methods allowing to take into account the role of the environment (Senn and Thiel, 2007, 2009; Meier et al., 2013; Monari et al., 2013). QM/MM also allows to validate the experimental use of simpler and more homogeneous model systems (Ding et al., 2009). The growing computational power experienced in the latter years, as well as the presence of more and more efficient algorithms and codes allows to tackle more and more complex problems and environments, providing an unprecedented comprehension of biologically relevant mechanisms.

In the present review we will present some examples of the molecular modeling of DNA under external stress, both concerning ground-state and photochemical pathways, underlying the fundamental questions that may be answered by a proper simulation and modeling.

2. Modeling of DNA Oxidative Stress

DNA oxidative stress is dominated by a combinatorial radical chemistry, in biological media hydroxyl (HO•) and peroxyl (HOO•) radical being by far the principal players. However, other non-radical reactive chemical moieties can play a role and have a strong biological influence: O2, H2O2, alkylating agents, low-energy electrons, and each of them has been tackled by simulation means.

2.1. An Ubiquitous First Step: Hydrogen Abstraction

We will hereby focus on radical induced ground-state reactivity, and mostly on the hydroxyl radical HO•, as its study exemplifies the methods employed in DNA damage molecular modeling. The hydroxyl radical is prone to abstract an hydrogen, with subsequent production of one water molecule and an highly reactive radical specie embedded in the biological macromolecule (see Figure 1); hence, this first step may also constitute the initiation step in radical chain reactions.

It has been shown that for a given double-strand DNA fragment (ds-DNA), an hydrogen uptake can occur at many positions, on the nucleobase (Cadet et al., 1999) but also on the sugar moiety (Pogozelski and Tullius, 1998), with a ratio of 9:1. Another reactive pathway activated by the hydroxyl radical is the direct addition of HO• onto the an ethylenic position of a given nucleobase B, thus forming in situ a radical entity [B–OH]•. Once again [B–OH]• may undergo further fragmentation or evolve toward the formation of a peroxyl nucleobase. Thus HO• has a deletious and versatile, chemical outcome with many possible subsequent fragmentation patterns. For instance, as schematized in Figure 2, a possible evolution may lead to the generation of abasic sites, or to the oxidative strand scission of nucleic acids (Cooke et al., 2003). Abasic lesions are characterized by oxidative pathways that lead to the disruption of the base pyrimidine or purine kernel, leaving only the sugar as part of the strand. On the other hand strand scission, mostly affect the sugar and the backbone (Figure 2) and ultimately results in a breaking of the strand continuity. If the previous lesions may easily be recognized and repaired by specific enzymes, also more dangerous lesions, particularly refractory to the repair may be produced (Bergeron et al., 2010). The former are produced by the attack of an oxidized nucleobase onto a vicinal one (either situated on the same strand or on the opposite strand), this gives rise to intra- or interstrand cross-links respectively. This particularly complex and significant case will be dealt to in Section 2.3.

Modeling has been early invoked to provide a rationale, and ultimately predict, the preferential hydrogen abstraction sites: indeed, because of its general high reactivity, the hydroxyl radical selectivity is governed by the accessibility to the solvent. The latter is an information reliably estimated by bioinformatics approaches such as the computation of Solvent Accessible Surface Area (SASA). Tullius et al. have exploited this approach to rationalize the HO•-induced DNA strand breaking (Balasubramanian et al., 1998), hence providing a clear-cut structural basis to rationalize the abstraction site preference. Indeed the authors have clearly shown the existence of a very good correlation between the reactivity of the different sites and their solvent. The C5 carbon atom was indeed found to contribute to the total cleavage by 57%, the C4 position accounting for the 22% and the C3 for the 17%. The most reactive C5 position on the other hand shows a solvent accessible area of 46%, while the C4 and C3 are characterized by 28 and 14% accessible surface, respectively. Furthermore, this rather simple...
molecular dynamics (CPMD) to describe the hydrogen abstraction on a solvated guanine (Wu et al., 2004; Abolfath et al., 2012). This work nicely underlines the contrast between the static approaches where temperature effects and hence thermal fluctuations are neglected and the dynamic picture. Indeed while in vacuum the hydrogen abstraction occurs at around $t = 0.07$ ps, water solution slows down the reaction rate roughly by a factor of two ($t = 0.12$ ps). The hampering effect of the water environment is due to thermal fluctuation of the solvent molecules, that are also highly organized because of hydrogen-bond networks, ultimately resulting in a strong disturbance of the OH· motion. The extension of this protocol to a ds-DNA model would constitute an interesting perspective. However reaching such a goal will require describing most of the system at the molecular mechanics (MM) level, while the reactive center (i.e., the nucleobase) should be described quantum mechanically (QM). Furthermore, water molecules close to the nucleobase should be included in the QM partition, but due to water high mobility one has to take into account the fact that some solvent molecules may move out from the significant QM partition while others may enter. Such schemes, known as adaptative QM/MM, have been pushed forward over the last years (Nielsen et al., 2010) but are for now applied only to model reactions. An interesting alternative for large scale molecular modeling could rely on the use of reactive MD, which has been realized relying on the ReaxFF force field (Abolfath et al., 2011).

However, QM/MM-MD simulators, e.g., in a Car-Parrinello framework, have nowadays been successfully performed both for reactivity in DNA double strands and in solvated nucleoside (Abolfath et al., 2012; Garrec et al., 2012). Compared with reactive MD force-fields and previous calculations, these studies have the advantage to exploit a less-parameterized density functional. Furthermore, they also provide a more direct comparison of free-energy profiles between an isolated system and a DNA fragment. Indeed, it is worth emphasizing that DFT has been employed in a near-exclusive way not only for ground-state reactivity, but also for calculating eletronical vertical excitation (Durbeej and Eriksson, 2003). More recently, several works have also described photo-induced hydrogen abstraction (Szabla et al., 2015), resorting to a multi-reference description of the electronic density through complete active space self-consistent (CASSCF) or second order perturbation theory (CASPT2) approaches. Remarkably, in the previous contribution the authors identified a reactive conical intersection joining the $S_0/S_1$ of $\beta$-2′-deoxycytidine. Although, from the $S_1$ minimum the system should overcome a rather large barrier of 0.59 eV to reach the conical intersection, the excess vibrational energy at Franck-Condon geometry safely allows for its crossing and hence justify the active photochemistry. Even if multi-configurational methods can in principle be used within a QM/MM-MD framework, they are mostly restricted to the treatment of excited states due to their efficiency in the description of non-adiabatic potential energy points (conical intersection) and to the computational bottleneck that limits the time-scale of the MD trajectory to hundreds of fs (Szabla et al., 2015). However, unlike for other systems or reactions, the exchange-correlation density

![FIGURE 2](image.png)

FIGURE 2 | Evolution to abasic site and DNA strand scission following oxidative attack on the DNA sugar moiety.

approach may easily be generalized to preview the reactivity in presence of diffusion controlled reactants and may be extended to the investigations of RNA reactivity and repair processes. Monte-Carlo simulations of site-specific radical attack to DNA bases have also proved their usefulness (Nijkooy et al., 1999; Bulent et al., 2008), with the advantage to go beyond the single-structure and static description. On the other hand, the almost immediate estimate of SASA paves the way toward massive parallel-sequencing-based hydroxyl radical probing of RNA accessibility (Kielpinski and Vinther, 2014).

The hydrogen abstraction is the key limiting step of the whole damage process (Regulus et al., 2007; Nikitaki et al., 2015), and its in-depth understanding can take full profit from quantum mechanics calculations: the latter are most often rooted in density functional theory (DFT), which allows to discriminate the most stable radical centers. Most particularly, DFT can easily be used to calculate the carbon–hydrogen bond dissociation energy (BDE) on different nucleobase and sugar potential reactive sites. This description allows to go beyond the simple solvent accessibility, indeed a lower BDE will be recognized as a crucial factor favoring the reactivity of the specific site.

For instance, DFT confirms that hydrogen abstraction for pyrimidine nucleobases, i.e., cytosine and thymine, occurs preferentially on C5 positions and on the corresponding methyl group (C5m), not only because of solvent accessibility but also due to electronic favors (Ji et al., 2005; Frances-Monneris et al., 2013; Yadav and Mishra, 2013). Interestingly enough, while most natural occurring nucleobases lead to the formation of $\pi$-radicals upon H abstraction, the substitution of the carbonyl oxygen with sulfur induces the formation of the quite rare $\sigma$-radicals (Besci et al., 2001; Vianello and Maksic, 2003; Gomzi, 2011).

It is remarkable that a sound agreement can be achieved between experiment and theory even when calculations on model, isolated moieties are performed (Bera and Schaefer, 2005). More recently, Papiez et al. have employed Car-Parrinello
functional dependence is less pronounced for the study of DNA and nucleobase hydrogen abstraction reactivity, and different functional classes usually provide the same insights. In contrast, the inclusion of a dispersion is particularly important as it was evidenced for instance by the Bickelhaupt's group (Poater et al., 2011).

2.2. Probing the Role of the Environment for Single-nucleobase Reactions

When coming to DNA reactivity and damage generations, experiments are performed on isolated nucleobases, small DNA fragments (trinucleotides), longer oligonucleotides or even larger ds-DNA structures. It is now well recognized that reactivity in model systems can largely differ due to the geometrical constraint imposed within a ds-DNA and the strongly heterogeneous environment. Providing a most direct comparison between reaction profiles for isolated moieties vs. nucleobases embedded in a complex macromolecular environment allows to pinpoint the structural and electrostatic factors that can dramatically tune or reserve the intrinsic reactivity of nucleobases. Many elementary lesions in DNA subunits have been first targeted based on QM studies on isolated fragments (Duncan Lyngdoh and Schaefer, 2009).

Predictions of the reactivity order between different nucleobases can indeed be assessed on model π-stacked systems (thymine propensity to form lower triplet states or dimerize (Durbeej and Eriksson, 2003), lower ionization potential of guanine, ...). Both DFT and (multireference) post Hartree-Fock approaches can be employed, (e.g., on the guanine radical cation Hutter and Clark, 1996; Sponer et al., 2001; Jurecka et al., 2006), DNA reactivity has been a field of investigation for conceptual DFT, with derivation of DFT-based descriptors to rationalize intrinsic trends in DNA reactivity (Labet et al., 2014). Yet, this supposes that reactivity is governed only by electronic factors, neglecting the influence of the strongly heterogeneous environment, and the fact that the flexibility of short oligonucleotides can induce noticeable deviations from the canonical B-DNA parameters.

Cluster models where a reaction profile is obtained employing hybrid QM/MM (Barnett et al., 2006) or QM/MM’ approaches (Cauët et al., 2013; Ceron-Carrasco and Jacquemin, 2013; Jacquemin et al., 2014) provide a first step toward more realistic systems. However in these models dynamics is neglected and implicitly such schemes assume that reactivity occurs with no distortion of the B-helix (extremal nucleobases are kept frozen). Furthermore, even larger deviation of the ideal B-DNA structure can be induced by the lesions themselves, with consequences that can be crucial for further reactivity or for the repair ratio.

Structural and dynamic changes of an oxidized abasic site in ds-DNA have been studied by molecular dynamics to unravel representative conformation(s) of the double-helix and infer the role played by structural modifications on the DNA chemical reactivity (Chen et al., 2008; Patel et al., 2013). Indeed it has been shown that abasic sites show a great conformational flexibility (Patel et al., 2013), and most notably MD has unambiguously shown that abasic site is much better stabilized in the B-DNA double strand when coupled with a cytosine. Indeed in this case the lesion is accommodated in the duplex and stabilized by a pair of hydrogen bonds, in contrast purines nucleobases may at most form one hydrogen bond and as a consequence the lesions constantly flip in and out of the duplex. The blocking of the abasic site inside the B-DNA by cytosine has a strong consequence on reactivity since it can favor reactive conformations leading to interstrand cross-links.

Indeed, MD simulations provide a starting point for tackling the reactivity, within the framework of hybrid QM/MM–MD simulations. A decade ago Parrinello and coworkers rationalized the role of the B-DNA (Gervasio et al., 2004) on the reactivity of the guanine radical cation. This pioneer work constitutes a first probe of a reversal for the stability of the GC+:C vs. G(−H):C -C(H)+ couple in B-DNA, which is due to the electrostatic interaction with the negatively charged backbone as well as to the different geometrical structures adopted by the pair.

QM/MM–MD simulations of DNA reactivity are becoming more and more popular, with several choices concerning the bias techniques (metadynamics Gervasio et al., 2004, umbrella sampling, thermodynamic integration Garrec et al., 2012) to sample the free energy reaction profile along one or several reaction coordinate(s).

The next subsection is devoted to theoretical studies focusing on reactions implying two nucleobases, where the distortion of the DNA structure is even larger and hence a proper dynamic description of the reaction becomes crucial.

2.3. Bridging Two Nucleobases: Structure and Reactivity

After the initial radical hit, modified nucleobases (X), such as oxidized AP, see Figure 2, can be formed which are prone to form covalent intra- or interstrand cross-links (ICL) with vicinal nucleobases. One often lacks the knowledge of a proper experimental structure of the damage to ascertain the conformation of the reactant, or to probe the final structure of the product (Patel et al., 2013).

Formation of ICLs obeys a multistep pathway that can be studied based on model systems (Labet et al., 2008; Sviatenko et al., 2012). But the study of ICLs also differs from single-nucleobase lesions since the B-DNA structure is much strongly impacted by the formation of this kind of defect, which could be in line with the fact they are refractory to repair.

The formation of ICLs can be formally written as X → B, where X denotes the modified (oxidized) nucleobase and B one of the four canonical nucleobases (Figure 2). Their modeling covers two central aspects:

1. The reactivity to situate the barrier to covalently bind two nucleobases together. This can rely on DFT calculations performed on dinucleoside monophosphate, in gas-phase or with implicit solvation. Alternatively the reactivity can be treated within a B-DNA fragment, to account notably for the helical embedding. The barrier to overcome to bridge up two nucleobases together is strongly dependent on DNA conformation and on the nucleobases themselves. However
this dependence is non trivial, and attacks corresponding to strand offset have also been reported.

2. The structural consequences of the formation of such bonds, and in particular the free-energy necessary to induce the helical deformation. In this case molecular modeling can palliate the lack of experimental structures. In particular modeling provides starting conformations of the reactant, but perhaps more importantly, provides a computationally efficient way to build up in silico the structures of a given modified oligonucleotide.

The latter point is fundamental in the absence of NMR or X-ray structure even for the most commonly formed intra- and inter-strand oxidatively-generated lesions. Also, from a modeling perspective, the typical timescale for the reorganization of a 12 base-pair oligonucleotide bearing a central defect, poses a central challenge since it spans tens of nanoseconds, to be compared to the current limitations of 10–100 ps for QM/MM-MD schemes. The identification of suitable reaction coordinates and collective variables describing the process will allow the use of rare-event accelerating methods such as meta-dynamics, however the problem is still far from being trivial. Their inherent complexity also explains why the modeling of this class of lesions is much less developed, despite their strong biological significance.

One important breakthrough in the study of intrastrand-cross links has been recently performed using Car-Parrinello QM/MM molecular dynamics (Garrec et al., 2012). In this paper the authors have performed thermodynamic integration to unravel the free-energy profile of the reaction between cytosine and a nearby C5m-thymine radical leading to the so called G[8,5-Me]T lesion. The geometrical constrains imposed by the B-DNA environment helps in keeping the reactant in an optimal π-stacked conformations, and as a results the process is characterized by a relatively low activation energy (about 10.0 Kcal/mol) on the other hand isolated nucleotides shows a rather larger deviation from the ideal conformations and as a result the activation free energy goes up to about 70.0 Kcal/mol. Furthermore, the relatively small disruption of the ideal B-helix structure induced by G[8,5-Me]T could also explain the observed low repARATION rate.

All in all it clearly appears that the modeling of ground-state DNA reactivity is an extremely complicated problem requiring the correct equilibrium between a proper description of the electronic factors and a suitable sampling of the conformational space of a very flexible macromolecule. To tackle the latter problems some techniques developed in particular in the field of the enzymatic catalysis (Garcia-Mesenguier et al., 2013; Zinovjev et al., 2013; Zinovjev and Tunon, 2014) should be considered both to define proper reaction pathways and to perform a good sampling compatible with the reaction time-scale. On the other hand the problem of an accurate sampling of the conformational space or of the definition of suitable collective variables is less dramatic when dealing with excited-state phenomena, i.e., photochemistry and photophysics, even if still the coupling of electronic factors with vibrational and dynamical ones cannot be underestimated. These aspects will be the focus of the next section.

3. Interaction between DNA and UV Radiation

DNA is constituted by π-stacked nucleobases that can be regarded as chromophores and whose absorbance is mostly confined in the UVB regions of the spectrum. The absorption region capable of producing DNA lesions can be enlarged for instance by photosensitization, i.e., by the interaction with endogenous or exogenous compounds (Epe, 2012; Cadet and Wagner, 2013).

3.1. Direct UV Absorption and Decay

Even if UVB wavelengths are normally filtered by the ozone layer, and efficient deactivation pathways of the nucleobases excited states exist, DNA lesions resulting from direct photoactivation are usually reported and are considered as extremely dangerous for their potential carcinogenicity.

The photochemical pathways leading to the formation of the cyclobutane thymine dimer \((T \leftrightarrow T)\) and of the 6-4 photodadducts \([64]-PP\) in the case of stacked nucleobases has been the subject of interesting combined theoretical and spectroscopic study (Banyasz et al., 2012). After a thoughtful validation of TDDFT techniques by comparison with high-level CASPT2 techniques the authors efficiently analyze the differences between the paths leading to the two main photoproducts \([T \leftrightarrow T]\ or \([64]-PP\). In particular in the case of \(T \leftrightarrow T\) dimerization, after absorption at 5.36 eV, an efficient and barrierless path in the first \(\pi\pi^*\) excited state leads to a short lived minimum at 4.45 eV connected to the conical intersection (CI) region. Furthermore, the CI is characterized by a very short intermonomer distance, most notably the C5-C5′ and the C6-C6′ distances have been found to be of 2.5 and 2.06 Å, respectively, hence the topology of the CI clearly favors dimerization. Notably, no sterical hindrance or electrostatic effect due to the backbone are observed. On the contrary the formation of the \([64]-PP\) adducts, and in particular of the key oxetane intermediate, involves the population of a charge transfer (CT) state. The CT state involves both thymine and is found at 5.50 eV. From Franck-Condon region the excited states may relax toward a very short living minimum that efficiently evolves toward the oxetane conical intersection region. Once again, the geometrical features of this conical intersection favor dimerization, in particular very short intermonomer distances are found (C6-C4′ 2.46 Å and C5-O8′ 1.65 Å). Note that environment effects are crucial since the backbone charges efficiently stabilize the charge-separated state compared to the gas-phase situation, and hence strongly favor its population. Furthermore, a barrier in the CT state potential energy surface, and probably due to dynamical solvent effects, needs to be overcome. This fact supports the evidence that although the CT state is reachable upon UVA irradiation, shorter wavelengths excitation are needed to allow the system the excess kinetic energy.

Recently, more complex photochemical mechanisms have been taken into account (Esposito et al., 2014). Once again combining spectroscopy and modeling, the study of a trinucleotide of sequence TCG allowed to gain insight into the competition between the formation of cyclobutane pyrimidine
dimer (CPD) and (64)-PP adducts. Interestingly the role of
the methylation of the 5′ position of guanine has been shown
to be far from innocent in dictating the relative ratio between
the CPD and (64)-PP. Indeed, the authors found that upon
methylation the ratio between the quantum efficiency for the
CPD production versus the (6-4)PP one ($\psi_{CPD}/\psi_{6-4PP}$) goes
from 1.52 to 3.68. The important increase of the selectivity
among CPD has been rationalized by combining QM and
MD calculations and ascribed to the sugar puckering which
modulates the stacking arrangement of the reactants. Indeed,
methylation strengthens interaction with flanking T and globally
stabilizes conformers that are more reactive toward CPD
formation. Since C5 position methylation of guanine is known
to play important role in epigenetic gene regulations and is
found as hot-spot in skin cancer the biological relevance of
these findings are evident. The influence of the double helical
pairing has also been taken into account, globally confirming the
previous results.

The same group (Vayá et al., 2012) has also revealed the role
of the DNA structural organization in dictating its spectroscopic
and photochemical properties. Indeed the exciton coupling of
the nucleobases is reflected in an energy- and charge-transfer
along the double helix that is reflected in the fact that DNA
lesions hot-spot depend on the nearby base sequence and are
not randomly distributed. From a photophysical point of view
an interplay between charge and energy transfer is observed. Still
it will be important to determine the sequence effects favoring
charge-separation and charge-recombination.

Barbatti and Lischka have reported a systematic study of the
deactivation pathways for the four DNA nucleobases (adenine,
thymine, guanine and cytosine) as well as for uracil (Barbatti
et al., 2010). The use of non-adiabatic state-hopping molecular
dynamics (Barbatti, 2011) has allowed the authors to give clear
insights on the deactivation pathways and to reveal two distinct
behaviors between purines and pyrimidines. Indeed, adenine and
guanine have a relatively simple photodynamic, characterized by
a rapid evolution from the $\pi\pi^*$ up to the conical intersection,
where they are funneled back to the S0 ground state. This simple
mechanism is also reflected in the extremely fast deactivation
with an estimated excited state life time of 0.28 and 0.77 ps for
guanine and adenine, respectively. In contrast, for pyrimidines a
much more complicated situation is found with the competition
of two channels: one evolving to the $\pi\pi^*$ conical intersection and
the other one toward the nπ*$ crossing seam. The ratio between
the two channels depending on the bases, for instance in cytosine
the direct $\pi\pi^*$ path is not activated at all. The more complex
photochemistry of pyrimidine bases, also due to the presence of
low lying $\pi\pi^*$ states with flatter potential energy surfaces, is
reflected in the longer life-time of excited states spanning values
from 0.53 to several ps.

Robb group (Groenhof et al., 2007) also reported a
computational study of the deactivation pathways of a cytosine-
guanine base-pair both in gas phase and embedded in a DNA
environment. Their QM and QM/MM multiconfigurational
calculations revealed that the deactivation pathway starts with a
proton transfer, that is followed by a radiationless decay leading
to an extended conical intersection seams. Indeed, the particular
topology of the conical intersection systems may lead to a number
of crossings between the first excited and the ground state, that
may ultimately lead to a diabatic blockage. After the crossing to
the ground state the proton is transferred back and in a majority
of situation the initial conformation is restored. Interestingly,
comparing QM and QM/MM calculations, the authors conclude
that the Watson-Crick pairing keeping the base pair closer
together enhanced photostability minimizing the fraction of
trajectory ending up in a different tautomeric state compared to
the initial configuration.

Novel strategies to prove the photostability of DNA
nucleobases start emerging, notably very recently Garaveli’s
group reported the modeling of near UV 2D-electronic
spectroscopy as a tool to characterize the double excited, multi
excimer and charge-transfer states that may be responsible of the
ultrafast deactivation dynamics. These advancements could pave
the way toward a better interplay between theory and modeling
to give an atomistic level description of the different dynamic
phenomena (Nenov et al., 2015).

Although, extremely efficient DNA relaxation pathways may
lead to a drawback evidenced by the works of Gonzalez’s group
that have underlined a femtosecond intersystem crossing taking
place in the case of cytosine (Richter et al., 2012). The explicit
inclusion of spin-orbit coupling in the calculation of the hopping
probability has evidenced a non-negligible population of the
triplet state manifold, indeed after 140 fs the T1 state has
an important population of 10%. This striking result, even
if it neglects the environment, is extremely important from
a photophysical point of view, since it illustrates a situation
in which the quasi-degeneracy between triplets and singlets
compensates the moderate, but non-negligible, value for the spin-
orbit coupling. Moreover, from a more biological point of view,
if confirmed in the DNA embedding it may open the way to the
population of triplet states, that could be responsible of indirect
production of DNA lesions.

3.2. DNA Photosensitization

In DNA photosensitization the absorption of light and
the subsequent photochemical pathways are mediated by
an interacting chromophore. By definition in the case of
sensitization the chromophore needs to be recovered inaltered
at the end of the process, acting in an almost catalytical way.
This excludes covalent interactions with DNA that characterize
the action of important antitumoral drugs such as the cisplatin.

The study of DNA photosensitization is complicated by the
fact that one should be able to achieve a good knowledge of
different phenomena. First of all, a clear picture of the structure
of the DNA/sensitizers aggregate is compulsory. Usually three
main modes of interaction have been characterized for canonical
B-DNA (Zeglis et al., 2007), see Figure 3:

- Groove Binding. In which the sensitizer lies close to the DNA
grooves, and stabilization is mostly driven by electrostatic
interactions with the negatively charged backbone. Note that
usually a rather important selectivity is evidenced between the
minor or major groove, and that the double-helical structure
is only scarcely distorted.
• Intercalation. In which the sensitizer slips between two base pairs. The stabilizing interactions are mostly due to dispersion and π-stacking with the DNA nucleobases, and hence this mode is predominant for rather large, planar, conjugated structures. The DNA helix is slightly more perturbed since an intercalation pocket needs to be created.

• Insertion. In which the sensitizer ejects one of the bases from its Watson-Crick pairing, taking its place in the helix. Conjugated systems with non-collinear fused rings, i.e., mimicking the pyrimidine geometries, are the most likely to prefer insertion over intercalation.

Moreover the three interaction modes can most often coexist, with a population ratio strongly dependent on the environmental factors such as the DNA sequence or the salt concentration. As a consequence, X-Ray resolved structures are quite rare, and structural determination becomes crucial, requiring a proper modeling that may confirm more indirect experimental observation such as the ones based on circular dichroism.

On that context we should cite the extensive molecular simulations performed by the Lavery group using molecular dynamics and different sampling techniques such as metadynamics and umbrella sampling (Wilhelm et al., 2012). The authors were able to unambiguously determine two stable states for the binding of Daunomycin drug to DNA: minor groove binding and intercalation. Furthermore, they were able to recover the free energy profile along the pathway connecting the two modes. A thermally accessible low barrier of about 6.5 kcal/mol was indeed found to separate the two stable modes, with intercalation being about 2.0 kcal/mol more stable. Interestingly, also an intermediate state was found with the drug only partially intercalated while DNA is strongly bended away from the drug. This intermediate state, sometimes called semi-intercalation, is also found in a number of other drugs or even in the case of minor-groove binding protein. In the case of daunomycin, however, the particularity lies in the fact that it is only connected to a very shallow minimum in the free energy profile, and hence does not constitute a kinetic blockade for the full intercalation.

Also the interaction between simple organic molecules and DNA should be described most carefully, which necessitates the resort to molecular dynamic simulations to sample the relevant possible conformations. A particularly striking case is the one of benzophenone, a paradigmatic sensitizers (Cuquerella et al., 2012), for which no experimental structure has ever been reported for the interaction with DNA. Indeed, by analyzing the molecular dynamics trajectory only 2 stable interaction modes were evidenced (Dumont and Monari, 2013), one of them was the usual minor-groove binding, but the second one was a novel mode called “double insertion” (Figure 3). In this mode the sensitizer, benzophenone, ejects both the bases constituting the Watson-Crick pair and take the place of both in the stacked structure. The use of the so-called non covalent interactions (NCI) technique to visualize non-covalent interaction (Johnson et al., 2010) allowed to pinpoint the emergence of a very strong hydrogen-bond between the ejected base and the phosphate of the backbone as one of the factors explaining the stabilization of the double-insertion mode, even in presence of a strong helical deformation. Furthermore, the calculation of induced circular dichroism, using QM/MM methods coupled to the MD simulation, evidenced a clear spectroscopic signature for the double-insertion in the UVA region that could allow for an easy experimental confirmation (Dumont and Monari, 2013). Indeed, while the minor groove-bound benzophenone does not show any
induced circular dichroism spectra, the double-inserted mode presents a well resolved negative band at 350 nm, corresponding to the $n^r\pi^*$ excitation. Since this band belongs to the UVA and is far from the native DNA absorbing region it is supposed that it could allow for a quite straightforward discrimination of the interaction modes. Recently and independently the double-inserted mode has been also evidenced in the case of copper complexes interacting with DNA (Galindo-Murillo et al., 2015) using very long scale molecular dynamics up to the $\mu$s scale.

Photosensitization involving the excitation of the chromophore, and the subsequent excited-state interactions with the nearby nucleobases, the influence of the environment on excited state properties is crucial and should be thoughtfully taken into account. Indeed the optical properties of many chromophores can be modulated by the environment, one of the most well known cases being the light-switch effects (Friedman et al., 1990), i.e., the activation of luminescence induced by DNA for the Ruthenium complex $\text{[Ru(dpzp)(bipy)]}^{2+}$ (bpy = 2,2'-bipyridine; dppz = dipyridophenazine). Indeed the latter complex happens to be dark in water, weakly luminescent in non protic solvent like acetonitrile and strongly luminescent in water. One of the first rationalization of this phenomenon was proposed by Batista, using TDDFT calculations coupled with a continuum description of the environment and a micro-hydration scheme (Batista and Martin, 2005). Batista suggested the presence of two quasi-degenerate triplet states: one metal to ligand charge-transfer (MLCT) centered on the dpzp ($^3\text{MLCT}_{dpzp}$) and one intraligand (IL) $\pi\pi^*$ triplet still centered on the dpzp ($^3\text{IL}_{dpzp}$). The authors attribute the quenching of luminescence by water to the stabilization of the dark $^3\text{MLCT}_{dpzp}$ that becomes the lowest triplet state and hence easily populated. The stabilization would be mostly due to hydrogen bonds between solvent molecules and the nitrogen dpzp atoms.

More recently a refined explanation for the light-switch effect was offered coupling a multiscale modeling to describe the interactions with the surroundings together with TDDFT and high level ab-initio calculations. Firstly the intercalation interaction mode and the main features of the absorption spectrum of $[\text{Ru(dpzp)(bipy)}]_2^{2+}$ interacting with DNA were modeled, also taking into account polarizable embedding and validating the computational protocol (Very et al., 2012). To this end the presence of the charge-transfer band due to the Ruthenium complex was evidenced at around 450 nm. The band was assigned to a complex manifold of different metal-to-ligand (MLCT) transitions. Subsequently (Very et al., 2014), the triplet excited state manifold of $[\text{Ru(dpzp)(bipy)}]_2^{1+}$ in different environments, namely water, acetonitrile and DNA was computed, and the distribution of electronic density analyzed. The authors concluded that in water the dark $^3\text{MLCT}_{dpzp}$ state was populated being the lowest one upon geometry relaxation, in acetonitrile on the other hand the lowest triplet excited state was still of MLCT character but was centered both on the dpzp and bipy ligands ($^3\text{MLCT}_{dpzp/bipy}$) since in contrast to the dpzp centered one the $^3\text{MLCT}_{bipy}$ are brights the mixed $^3\text{MLCT}_{dpzp/bipy}$ can be considered as weakly emissive, coherently with the experiment. Finally in the case of intercalation in DNA the triplet state becomes now entirely centered on the bipy ancillary ligand ($^3\text{MLCT}_{bipy}$) giving rise to a bright state and hence to the triggering of luminescence. Therefore, the light-switch effect does not happen because of a competition between MLCT and IL triplet states but instead by a change in the electronic density reorganization of the MLCT states operated by the environment.

Usually photosensitization can be classified considering the physical phenomenon producing the lesion with DNA. Type I photosensitization is due to a charge transfer usually toward guanine, type II sensitization on the other hand is based on an energy-transfer between the sensitizer (triplet) excited state and molecular oxygen giving raise to $^1\text{O}_2$ and the subsequent oxidative events. Finally, one should also take into account triplet photosensitization characterized by an energy-transfer between the photosensitizers and DNA bases, usually thymine; thymine dimers are the most common final products of triplet sensitization. Since the physical processes underlining these photosensitization pathways are different, modeling strategies should be tailored to provide a satisfactory description.

Concerning type I photosensitization some evidences of important electron transfer between nearby guanine and different ruthenium complexes have been reported already at Franck-Condon level using QM/MM partition and TDDFT calculations (Chantzis et al., 2013). Indeed, in this contribution the authors have calculated the absorption spectrum of ruthenium complexes intercalated in DNA extending the QM partition to accommodate the nearby guanine and adenine base. By analyzing the electron density reorganization, it has been shown that some vertical transitions exist around 300 nm characterized by a very strong charge-transfer, the calculation of natural population analysis (NPA) charges have allowed to estimate such charge-transfer to be close to 0.7 electron.

Also in the case of type II sensitization the role of the environment may be far from innocent. For instance in the case of a DNA interacting drug known as palmatine the production of $^1\text{O}_2$ as well as fluorescence is triggered only by interaction with DNA (Hirakawa et al., 2012). Once again only QM/MM methods coupled with MD allows for a rationalization of the photochemical pathways (Dumont and Monari, 2015). Indeed, it was shown that in water solution upon excitation to a valence $\pi\pi^*$ ($3.0 \text{eV}$), state palmatine relaxation crosses a CT state that happens to be, at his relaxed geometry, lower in energy than the valence state ($3.8$ against $4.0 \text{eV}$, respectively). This fact opens the way to efficient non-radiative pathways and hence induces quenching of luminescence and the inhibition of singlet oxygen production. In the case of the interactions with DNA (happening through minor-groove or insertion) already at Franck-Condon geometry the CT state is strongly destabilized by more than $1 \text{eV}$ comparing to the water-solution case. As a consequence the crossing with the $\pi\pi^*$ state is impossible even after geometry relaxation. Therefore, when interacting with DNA, palmatine evolves toward the $S_1$ ($\pi\pi^*$) state that can subsequently either relax radiatively (fluorescence) or give intersystem crossing to populate the triplet state and produce $^1\text{O}_2$. Intersystem crossing is also supported by the high spin-orbit
coupling calculated to be of the order of 40 cm$^{-1}$. We note that in this contribution (Dumont and Monari, 2015), it was evidenced that in the case of planar conjugated systems, the calculation of the absorption spectrum necessitates to take into account the effect of the low-frequency vibration modes, for instance through the coupling with MD. Indeed the calculation of absorption spectrum with static procedures gives totally unreliable results. Indeed, static absorption spectrum has a shift of about 100 nm compared to experimental results, while the inclusion of dynamic effects allows to perfectly match the observed absorption maximum (420 nm) providing one uses a long-range corrected functional. This situation was also observed for other DNA interacting systems such as the $\beta$-carboline harmane both for absorption and fluorescence spectra (Etienne et al., 2013, 2014).

The role of the environment in tuning type II photosensitization, and in particularly in driving the population of the triplet manifold was recently pointed out by Nogueira et al. in the case of the interaction between DNA and phenothiazinium dyes (Nogueira et al., 2013, 2014). The author evidenced that while in water solution intersystem-crossing is only driven by relatively inefficient vibronic spin-orbit coupling, the interaction with DNA modifies the energy order of the excited states and hence opens the way to a second symmetry allowed crossing. Indeed, in water solution the dye has to overcome significant barriers both from Franck-Condon region and $S_1$ minimum (0.77 and 0.51 eV, respectively) to reach the symmetry allowed intersystem crossing between $S_1$ and $T_3$. On the other hand in DNA, mostly because of the hydrogen-bonding networks, the modified potential energy landscape for singlet and triplet results in a significant lowering of the barriers that assume values of 0.15 and 0.30 eV, for Franck-Condon or $S_1$ minimum, respectively hence the intersystem crossing region can now be safely reached thanks to thermal energy and $^1O_2$ ne produced. The observed production of singlet oxygen in water is ascribed to the coupling between electronic and vibrational movements, i.e., to the so called vibronic spin-orbit coupling, happening between $S_1$ and $T_2$ state, however this channel being less effective that the fully allowed electronic spin-orbit coupling with $T_3$ the enhancement in the production of $^1O_2$ by the DNA environment is confirmed.

Triplet photosensitization is also an experimentally-known component of DNA damage that can be timely tackled by modeling. Since it involves the preliminary population of the photosensitizer triplet state manifold, one may ask if the macromolecular environment keeps the road to photosensitization open, or if it hinders its development. Acetophenone is a known triplet photosensitizer (Epe, 2012) and its efficiency is due to a very efficient population of the triplet state in gas phase (Huix-Rotllant et al., 2013). Molecular dynamics simulations have allowed to identify acetophenone stable interaction mode with DNA, namely intercalation (Huix-Rotllant et al., 2015). Furthermore, QM/MM calculations, performed both at CAS-PT2 and TD-DFT level of theory, have shown that the gas-phase quasi-degeneracy of the first singlet excited state $S_1$ with the two $\pi\pi^* \& \;\pi\pi^*$ triplet state is not significantly altered by the environment (Huix-Rotllant et al., 2015). The authors have also evidenced that all along the MD trajectory the spin-orbit coupling between the $S_1$ and at least one of the triplet is always quite high (about 20 cm$^{-1}$), i.e., a value that allows efficient intersystem crossing if the energy difference between the states is small (Richter et al., 2012). One of the most striking features of gas phase acetophenone photochemistry, explaining the high-efficiency of the intersystem crossing is indeed the presence of a three-point crossing between the singlet and the two triplets. In presence of DNA although the perfect crossing is lifted the two intersections still happens in a very close-by region of the configuration space (Huix-Rotllant et al., 2015). All these results point toward the fact that the DNA environment does not alter the efficient photophysical pathways leading to acetophenone triplet population during the very first vibrational motions. This in turn points toward the rationalization of the efficient DNA photosensitization experienced by acetophenone.

But in order to be efficient in terms of triplet-photosensitization, the population of the triplet manifold should be rapidly followed by energy transfer to the nearby thymine. This aspect has been recently deeply analyzed for the case of benzophenone (Dumont et al., 2015). The energy profile for the benzophenone-DNA triplet transfer has been obtained following an approximate reaction coordinate both for a double-inserted and a minor-groove bound benzophenone. Calculations have been performed by QM/MM to take into account the environment effects, while triplet energy has been obtained both at CASPT2 and TD-DFT level. In particular it has been evidenced that for double-insertion two very efficient and barrierless sensitization channels exist one proceeding from the $T_1$ state ($n\pi^*$) and the other from the $T_2$ ($\pi\pi^*$) triplet. On the contrary in the case of minor-groove binding the $T_1$ sensitization channel is hampered by a significant barrier (0.30 eV), while the $T_2$ channel is still active and virtually barrierless. Furthermore, it has been evidenced that due to the close-by distance of the interacting chromophores the transfer will proceed through the Dexter mechanism mainly driven by the electronic densities overlap. Estimation of the overlap has shown a slightly pronounced kinetic preference for the double-insertion mode, but still shows that the transfer from minor-groove is possible and efficient although with a 100 times lower efficiency compared to the double-inserted situation.

Obviously a deeper characterization of energy- and electron-transfer processes will necessitate to take into account non-adiabatic and state-hopping dynamics. Furthermore, since the triplet manifolds are involved in most of the cases the spin-orbit coupling should not be neglected. This aspect coupled with the necessity to take into account the environment and many interacting chromophores and the relative long time scale of the full sensitization process will necessitate a considerable computational effort that can be achieved only through a proper methodological development. Still significant progresses have been made allowing to get a clearer vision of a fundamental and rather complicated process whose biological relevance cannot be underestimated.
4. Sensitizing Non-canonical DNA Structures

Even if the canonical B-DNA form is by far the most common one, particular DNA sequences may adopt the so-called non-canonical conformations. The biological relevance of such structures is now clearly evidenced. On that context particular significance has to be attributed to the guanine-quadruplex (G4), that are present in guanine-rich sequences such as the telomeres (Neidle and Parkinson, 2003). Telomeres protecting activity is a key factor to regulate cellular apoptosis, indeed the activation of telomerase, which is overexpressed in a vast majority of tumor cell lines, is strongly related to cancer cells uncontrolled replication (Neidle and Parkinson, 2002). The stabilization of G4 structures by DNA sensitization may result in an inhibition of the telomerase (Neidle, 2010), hence opening the way to novel anticancer therapies. In particular the selectivity of the sensitizers toward G4, instead of canonical DNA structure, will be a key factor governing the overall selectivity of the drug toward cancer cells.

The relative peculiar structure of G4 has necessitated an important computational effort to rationalize their structure and their stability (Ilchenko and Dubey, 2014), as well as the sequence effects governing their formation. Furthermore, since G4 are stabilized by interactions with cations to compensate the negative charges the pH and salt concentrations are key factors determining their appearance. In a vast majority of studies (Ilchenko and Dubey, 2014) QM methods were used to analyze the stabilizing factors of G4. Systematic DFT studies were performed (Jissy et al., 2011), in addition the role of different cations stabilizing the central channel was investigated in particularly connected to the emergence of polarizability anisotropy that may induce birifrangence potentially exploitable in biomolecular imaging.

A rather serious complication in the study of G4 is connected to the polymorphism observed for such structures (Ilchenko and Dubey, 2014). This experimentally observed evidence, has also been rationalized by other DFT calculations (Fonseca Guerra et al., 2010), in particular the role of the hydrogen-bond networks has been evidenced. Some of the same authors have also proposed new possible monomers leading to the formation of quadruplexes structure such as substituted xanthines (Szolomajer et al., 2011).

In parallel, the necessity to correctly represent the polymorphism of the G4 non-canonical structures has pushed toward the validation of force-fields, which are usually parameterized to reproduce the behavior of canonical B-DNA (Grunenberg et al., 2014). The comparison of force field descriptions with DFT calculations shows that extreme care should be taken to model the competitive rotamers. In particular while the OPLS and the MMFF force fields correctly reproduce experimental and DFT results, the DNA-popular Amber force field gives the opposite energy-order.

However, the combination of well parameterized force fields and of QM and QM/MM techniques allows to correctly reproduce the interaction of sensitizers with G4. The most common G4 sensitizers are rather extended and planar \( \pi \)-conjugated cations (Ilchenko and Dubey, 2014). Recently, Barone group (Terenzi et al., 2014) has used molecular modeling to predict the relative stability of G4 vs. B-DNA aggregates with square-planar organometallic Schiff-base complexes containing copper and zinc. MD simulations have shown that the Schiff-base strongly interact with G4 both via electrostatic interactions and through efficient \( \pi \)-stacking. The estimation of the G4/sensitizer free energy of formation gives values ranging from \(-34.6\) to \(-14.4\) kcal/mol therefore speaking in favor of a stable complex. It is noteworthy that these results have also been supported by biological tests confirming a potential anti-cancer activity.

It is evident that despite some difficulty the sensitization of non-canonical DNA structure, both at the ground and for the excited-state, is no more considered as an exotic novelty. On the contrary it is displaying all its potentiality in the rational design of novel therapeutic strategies. Molecular modeling is playing an active role on the field and will certainly drive not only the comprehension of the mechanisms leading to a selective sensitization, but also the prediction of entirely novel sensitizers.

5. Conclusions

The impressive progresses of modeling and simulation techniques have now made possible tackling problems as complex as the generation of oxidatively-induced DNA lesions and the interaction of DNA with potential drugs. However, this domain should still be considered very challenging also from a computational chemistry point of view. Indeed, many different factors should be taken into account and treated on the same footing to have a proper, and biologically relevant, description of the various phenomena comming into play.

Particular attention should be taken for instance to model the conformational degrees of freedom of the very flexible DNA macromolecule, this of course calls for adequate and long-scale molecular dynamics trajectory. Techniques allowing a good estimate of the free-energy, especially but not exclusively, in the case of sensitization binding energies are also of paramount importance. This aspect will of course require the development of efficient force fields as well as of proper sampling techniques.

The study of the electron density reorganization taking place on the ground- or on the excited states is also fundamental to go beyond the simple structural description and propose hints on the reactivity or the photochemistry of DNA alone or in presence of sensitizers. The envisaged QM or QM/MM techniques should conjugate the precision of the description with the efficiency necessary to make possible the sampling of important region of the configuration space, and hence having access to good statistical properties. This is especially true since in many case energy- or electron-transfer phenomena have to be taken into due accounts, i.e., one has to deal with some of the most challenging situations for QM methods.

We believe that from a methodological point of view the development of novel multiscale techniques in particular for what concerns excited state non-adiabatic dynamics as well as ab-initio QM/MM dynamics will allow the treatment of more and more complex scenarios, coupling a good accuracy with good statistics. On that context the development of proper sampling techniques allowing the definition of good and physical relevant
reaction coordinates taking into account all the relevant collective variables will certainly be one of the most important challenges of the next years.

Nowadays, the atomistic scale description of DNA related phenomena offered by multiscale modeling allows to rationalize and properly describe phenomena as diverse as the behavior of DNA under oxidative stress or the environment controlled behavior of phototherapy drugs. The selectivity of potential drug candidates toward different DNA configurations as well as the differential dynamical behavior of different DNA sequences are now accessible at least at a semi-quantitative level.

Modern day molecular modeling is therefore able to provide a molecular and electronic description of the factor governing DNA chemistry under external perturbation, therefore molecular modeling can nowadays be considered as a fundamental tool to provide unprecedented insights in chemical biology. Conversely, the knowledge of the mechanism of DNA lesions induction are also paving the way to the rational design of novel therapeutic agents. Therefore we may safely say that the communication between the in silico world of molecular modeling and the in vitro world of biological chemistry and cellular biology is now a sounding and promising reality.

**Author Contributions**

Both authors contributed equally to the present review.

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