Biological Evaluation of DNA Biomarkers in a Chemically Defined and Site-Specific Manner

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Abstract: As described elsewhere in this Special Issue on biomarkers, much progress has been made in the detection of modified DNA within organisms at endogenous and exogenous levels of exposure to chemical species, including putative carcinogens and chemotherapeutic agents. Advances in the detection of damaged or unnatural bases have been able to provide correlations to support or refute hypotheses between the level of exposure to oxidative, alkylative, and other stresses, and the resulting DNA damage (lesion formation). However, such stresses can form a plethora of modified nucleobases, and it is therefore difficult to determine the individual contribution of a particular modification to alter a cell’s genetic fate, as measured in the form of toxicity by stalled replication past the damage, by subsequent mutation, and by lesion repair. Chemical incorporation of a modification at a specific site within a vector (site-specific mutagenesis) has been a useful tool to deconvolute what types of damage quantified in biologically relevant systems may lead to toxicity and/or mutagenicity, thereby allowing researchers to focus on the most relevant biomarkers that may impact human health. Here, we will review a sampling of the DNA modifications that have been studied by shuttle vector techniques.

Keywords: DNA lesion; DNA damage; shuttle vector technique; replication block; mutagenicity; mutational spectrum; mutational signature; DNA repair; DNA adduct bypass; site-specific mutagenesis

1. Introduction

The human genome is constantly exposed to and damaged by endogenous chemicals, such as reactive oxygen species, lipid peroxidation intermediates, and alkylating agents. These electrophilic reactive chemicals, as well as environmental carcinogens and administered drugs, are known to generate various DNA adducts [1–3]. Some of the adducts block DNA replication or cause mutations and have been used as biomarkers to monitor the level of DNA damage or of disease progression [4–6]. One of the major goals for researchers is to understand the deleterious consequences of those lesions within the cell or animal. Among the different methods for studying the biological effects of the adducts, use of shuttle vectors containing a chemically defined lesion at a specific site has provided information about the biological and toxicological properties of the adduct [4,7]. The shuttle vector-based methods normally involve the steps outlined in Figure 1. Oligonucleotide synthesis: An oligonucleotide (oligo) containing a structurally defined lesion at a specific site is made either through a biomimetic route (in situ formation by direct chemical reaction, followed by HPLC purification of site-specifically modified oligo), or purely synthetically using a normal or convertible nucleoside phosphoramidite, etc. Vector construction: An ss- or ds-DNA vector containing the modified oligo is built by cutting the parent vector with one or a pair of restriction endonuclease(s), followed by ligation of the 5′-phosphorylated modified oligo. Cellular processing: The vector is transfected into different types of cells (e.g., Escherichia. coli (E. coli) or mammalian), and cellular polymerases are allowed...
to replicate or transcriptionally bypass the lesion under different repair or bypass conditions. *Data analysis:* DNA is extracted, amplified using PCR, and the biological outcomes are analyzed, which include the ability of the lesion/adduct to block polymerases or cause a mutation when processed by a polymerase during cellular replication. This assessment could be done by plaque or colony counting and picking with Sanger sequencing, $^{32}$P-post labeling and thin-layer chromatography (TLC), liquid chromatography-mass spectrometry (LC-MS), next-generation sequencing (NGS), etc. [4,5,7–9]. The shuttle vector-based method was initially introduced by Essigmann [7,9–11], further developed and utilized by Wang [4,5], Moriya [12,13], Livneh [14,15], Greenberg [16,17], Basu [18,19], Lloyd [20,21], Loechler [22,23], Fuchs [24,25], Pages [26,27], and others. Several informative review articles have been written by these authors on designing and applying the methods.

**Figure 1.** Schematic overview of the shuttle vector-based methods for evaluating DNA biomarkers.

In this work, we will review a variety of DNA biomarkers or probes that have been studied using the shuttle vector techniques and briefly summarize their biological outcomes. In all cases, focus is placed on the effect of the lesion to block replication and to cause mutations. For the details regarding the formation of DNA damage and other properties of the lesions, please refer to the original literature or review articles. We apologize in advance to researchers whose work we could not include in this review. After detailed discussions on individual lesions, we will provide some perspectives on possible future directions.

2. Discussions on Individual Modifications

Below, we will cover modified DNA structures generated from oxidative stress, alkylation, and other processes (Figures 2–5). In the following sections, the biological effects of a certain lesion are briefly summarized. Please see Figures 2–5 for chemical structures and Table 1 for detailed information.
Table 1. Bypass efficiency and mutagenicity of DNA modifications.

| Oxidative Lesion | Bypass Efficiency | Mutation | Cell |
|------------------|-------------------|----------|------|
| 8-oxo-G          | 88% [28,29]       | G>T 3% (44%, MutY-) [28,29] | E. coli |
|                  |                   | G>T 8% [30] | Human |
| Fapy-dG          | 31-43% (TXN sequences) [31] | G>T 1.2-1.9% (0.7-2.1%, MutM-MutY-) [31] | E. coli |
|                  |                   | G>T 10% [30] | Human |
| Nt                | 7% (57%, SOS) [32] | G>C 8.9%, G>A 19%, G>T 22% (G>C 2.5%, G>A 13%, G>T, 18%, SOS) [32] | E. coli |
| Oa                | 52% [28], 51% [33], 108% (118% MutY-) [29] | G>T 97% [28], 99% [33], 97% (no change, MutY-) [29] | E. coli |
| Oz                | 57% [28]          | G>T 86% [28] | E. coli |
| Ca                | 65% [28]          | G>T 95% [28] | E. coli |
| Gh                | 75% [34], 20% (30% MutY-) [29] | G>C 98%, G>T 2% [34], G>T 40%, G>C 57%, G>A 3% (no change, MutY-) [29] | E. coli |
| Sp1               | 9% [34], 19% (38%, MutY-) [29] | G>C 72%, G>T 27% [34], G>T 78%, G>C 19%, G>A 1% (no change, MutY-) [29] | E. coli |
| Sp2               | 9% [34], 17% (30%, MutY-) [29] | G>C 57%, G>T 41% [34], G>T 49%, G>C 48%, G>A 3% (no change, MutY-) [29] | E. coli |
| Ur                | 11% [35], 10% (10% MutY-) [29] | G>T 99% [35], G>T 54%, G>C 35%, G>A 9% (no change, MutY-) [29] | E. coli |
| Iz                | 60% (71%, SOS) [32] | G>C 88%, G>A 2%, G>T 1.1% (G>C 75%, G>A 3.4%, G>T 5.5%, SOS) [35] | E. coli |
| Cyclo-dG         | 11% (6% pol V-) [36] | G>A 20% [36] | E.coli |
| S-cdG            | 4% [37]          | G>T 35%, G>A 20% [37] | Human |
| Cyclo-dA         | 31% (15% pol V-) [36] | A>T 11% [36] | E. coli |
| S-cdA            | 6% [37]          | A>T 12% [37] | Human |
| Tg                | 96% [38]         | E. coli |
| 5ClC             | 75% (75% AlkB-) [39] | C>T 5% (same in AlkB-) [39] | E. coli |
| 5-OH-C           | C>T 0.05%, C>G 0.001% [40] | E. coli |
| 5-OH-U           | C>T 83% [40]     | E. coli |
| Ugi               | C>T 80% [40]     | E. coli |
| THF AP site      | 6% [28], 5.8% [32], 4% (4% MutY) [29] | AP>T 50%, AP>C 26%, AP>G 7%, -1 del 13% (no change, MutY-) [29] | E. coli |
| 2-deoxyribonolactone | 5% (3% pol II-), (1% pol V-) [41] | T 35%, C 42%, A 12%, G 8%, 5′ U (T 42%, C 38%, A 6%, G 14%, 5′ C) [41] | E. coli |
| Alkyl Modification | Bypass Efficiency | Mutation | Cell |
| m1G               | 15% (3%, AlkB-) [9], 20% (2%, AlkB-) [33] | G>T 3% (G>T 57%, G>A 17%, G>C 6%, AlkB-) [9], G>T 4%, G>A 2% (G>T 52%, G>A 20%, G>C 4%, AlkB-) [33] | E. coli |
| m2G               | 90% (84% AlkB-, 98% DinB-, 96% AlkB- and DinB-) [42] | G>A 3% (2.7%, AlkB-3%, DinB-3%, AlkB- and DinB-) [42] | E. coli |
| e2G               | 100% (98% AlkB-), (106% DinB-), (99% AlkB- and DinB-) [42] | G>A 2%, G>C 1% (G>A 2.3%, AlkB-), (G>A 2%, AlkB- and DinB-) [42] | E. coli |
| N2-CM-dG         | 100% [43]        | Not mutagenic [43] | Mouse |
| R-N2-CEdG        | 39% (13% pol V- [44] | Not mutagenic [44] | E. coli |
| S-N2-CEdG        | 75% (28% pol V- [44] | Not mutagenic [44] | E. coli |
| FF                | 101% (100% AlkB-), (28% DinB-), (36% AlkB- and DinB-) [42] | G>C 1%, (G>A 1%, G>T 1%, AlkB-DinB-) [42] | E. coli |
| Hf                | 92% (88% AlkB-), (28% DinB-), (40% AlkB- and DinB-) [42] | G>C 2% [42] | E. coli |
| O6-mG            | G>A 99% [45,46]  | E. coli |
| O6-POB-dG        | 70% [47]         | G>A 90%, G>T 2.5% [47] | E. coli |
| O6-PHB-dG        | 40% [47]         | G>A 95% [47] | E. coli |
| O6-CM-dG         | 10% [47]         | G>A 10% [47] | E. coli |
| O6-ACM-dG        | 40% [48]         | G>A 6% [48] | Human |
|                    | 2% [47]          | G>A 30% [47] | E. coli |
Table 1. Cont.

| Alkyl Modification | Bypass Efficiency | Mutation | Cell |
|--------------------|------------------|----------|------|
| O°-HOEt-dG         | 15% [47]         | G>A 40% [47] | Human |
| PdG                | 25% [12]         | G>T 6% [12] | Human |
| α-OH-PdG           | 17% [12]         | G>T 11% [12] | Human |
| γ-OH-PdG           | 73% [12]         | Not mutagenic [12] | Human |
| 1,6-ωG             | 4% (2% AlkB-) (18% AlkB-DinB-) [8] | G>A 6%, G>T 6%, G>C 2%, ~1/2 del 5% (G>A 13%, G>T 13%, G>C 1%, ~1/2 del 9%, AlkB-), (same in AlkB-DinB-) [8] | E. coli |
| 2°F-2°3-ωG         | 21% (26% AlkB-) (14% AlkB-DinB-) [8] | G>A 30% (30% AlkB-), (30% AlkB-DinB-) [8] | E. coli |
| mA                 | 100% (12%, AlkB-) [9] | A>T 0.06% (0.61%, AlkB-) [9] | Human |
| eA                 | 85% (5% AlkB) [49], 130% (9% AlkB-) [49] | <0.5% (A>T 25%, A>G 5%, A>C 5%, AlkB-) [33], A>C 1%, A>T 1% (A>T 22%, A>C 8%, A>G 7%, AlkB-) [49] | E. coli |
| EN°-CMdA           | 98% [36]         | Not mutagenic [36] | E. coli |
| 5°-HB-dA           | 6% (35% pol k-) [48] | Not mutagenic [48] | Human |
| R,R-N°,N°-DHB-dA   | 100% [51]        | <1% [51] | Human |
| 5,5°,N°-N°-DHB-dA  | 60% [51]         | A>G 1% [51] | E. coli |
| R,S,1,N°-γ-HMHP-dA | 10% [51]         | A>T 2% [51] | E. coli |
| O°-MedT            | 60% [52], 55% [53] | T>A 1%, T>G 1% [52], T>A 56% [53] | Human |
| O°-EtdT            | 21% (5% pol V-) [55] | T>C 35%, T>A 15%, T>G 5% (T>C 10%, pol V-) [55] | E. coli |
| O°-nPrdT           | 45% [52]         | T>A 5%, T>G 3% [52] | Human |
| O°-nPdT            | 35% [52]         | T>A 12%, T>G 5% [52] | Human |
| O°-nBudT           | 30% [52]         | T>A 13%, T>G 6% [52] | Human |
| O°-BudT            | 15% [52]         | T>A 4%, T>G 2% [52] | Human |
| O°-BudT            | 15% [52]         | T>A 4%, T>G 2% [52] | Human |
| O°-POB-dT          | 3% [54]          | 12% T>A, 38% T>G [54] | E. coli |
| m3T                | 6% (4% AlkB-) [9] | T>A 32%, T>C 6%, T>G 2% (T>A 47%, T>C 9%, T>G 2%, AlkB-) [9] | E. coli |
| N3-EtdT            | 17% (3% pol V-) [55] | T>C 15%, T>A 21%, T>G 3% (Not mutagenic, pol V-) [55] | E. coli |
| N3-CMdT            | 55% [56]         | T>A 66% [56] | E. coli |
| O°-CMdT            | 40% [48]         | T>A 81% [48] | Human |
| O°-EtdT            | 40% [48]         | T>C 68% (25% pol C) [48] | Human |
| O°-MedT            | 32% [56]         | T>C 58% [56] | Human |
| O°-nPrdT           | 76% [55]         | T>C 84% (Not mutagenic, pol V-) [55] | E. coli |
| O°-nPdT            | 33% [56]         | T>C 82% [56] | Human |
| O°-nBudT           | 35% [56]         | T>C 42% [56] | Human |
| O°-BudT            | 30% [56]         | T>C 44% [56] | Human |
| O°-BudT            | 32% [56]         | T>C 29% [56] | Human |
| O°-BudT            | 24% [56]         | T>C 42% [56] | Human |
| O°-BudT            | 20% [56]         | T>C 25% [56] | Human |
| O°-BudT            | 22% [56]         | T>C 25% [56] | Human |
| m3C                | 100% (10% AlkB-) [9], 113% (14% AlkB-) [39], 98% (5% AlkB-; 115% DinB-; 7.5% AlkB-DinB-) [42], 100% (15% AlkB-) [39] | C>T 1% (C>T 14%, C>A 14%, C>G 2%, AlkB-) [9], Not mutagenic (C>T 55%, C>A 30%, C>G 1%, AlkB-) [57], Not mutagenic (C>T 41%, C>A 41%, C>G 4%, AlkB-) [42], Not mutagenic (C>T 32%, C>A 30%, AlkB-) [39] | Human |
### Table 1. Cont.

| Alkyl Modification | Bypass Efficiency | Mutation | Cell |
|--------------------|-------------------|----------|------|
| eC                 | 24% (13% AlkB-)   | C>A 24%, C>T 11% (C>A 49%, C>T 31%, AlkB-) [33] | E. coli |
| H-edC              | 1% [50]           | C>G 40% [50] | E. coli |
| 5hmC               | 10% [50]          | C>A 60%, C>T 32% [50] | Human |
| 5fC                | 100% [59]         | Not mutagenic [59] | E. coli |
| 5caC               | 72% [58]          | Not mutagenic [58] | Human |
| eC                 | 110% (Ada-, decreases from 140% to 70%) [61] | TT>GT 50%, TT>GC 15% [61] | E. coli |
| Rp-Me-PTE          | 30% [61]          | Not mutagenic [61] | E. coli |
| Sp-Me-PTE          | 190% [61]         | Not mutagenic [61] | E. coli |
| Rp-Et-PTE          | 40% [61]          | Not mutagenic [61] | E. coli |
| Sp-Et-PTE          | 160% [61]         | Not mutagenic [61] | E. coli |
| Rp-pPr-PTE         | 79% [61]          | Not mutagenic [61] | E. coli |
| Sp-pBu-PTE         | 100% [61]         | Not mutagenic [61] | E. coli |

#### Bulky Lesion Bypass Efficiency

| Alkyl Modification | Bypass Efficiency | Mutation | Cell |
|--------------------|-------------------|----------|------|
| N2-MC-dG           | 38% [62]          | G>T 18% [62] | Human |
| N2-2,7-DAM-dG      | 27% [62]          | G>T 10% [62] | Human |
| AL-II-dG           | 9% [63]           | G>T 9% [63] | E. coli |
| AFB1-N7-dG         | 51% [66]          | Not mutagenic [66] | Human |
| AFB1-FAPY          | 13% [66]          | Not mutagenic [66] | Human |
| C8-AF-dG           | 97% [66]          | Not mutagenic [66] | Human |
| AL-I-dA            | 100% (5% Rev3L-) [67] A>T 50% (Not mutagenic, Rev3L-) [67] | Mouse |
| AL-II-dA           | 5% [63]           | A>T 22% [63] | Mouse |

#### Crosslinked Lesion Bypass Efficiency

| Alkyl Modification (dG strand) | Bypass Efficiency | Mutation | Cell |
|-------------------------------|-------------------|----------|------|
| AP-dG (dG strand)             | 38% (43% Pol 63), (13% Pol 69), (2% Pol 70) [70] | G>A 2-5%, G>T 1-2%, G>C 1% [70] | Human |
| AP-dG (AP strand)             | 18% (25% Pol 71), (4% Pol 72), (1% Pol 73), (5 Pol 74) [70] | AP>T 74%, AP>C 10-20%, AP>G 4-6%, AP>A 1-2% [70] | Human |
| 1,2-GG-γ-ε-DDP*              | 11% [71]          | <0.2% (A>T 4.4%, SOS) [72] | E. coli |
| 1,2-AG-γ-ε-DDP*              | 22% [72]          | <0.2% (A>T 4.4%, SOS) [72] | E. coli |
| 1,2-GTG-ε-β-PP*              | 13% (14% SOS) [72] | <0.7% [72] | E. coli |

#### Crosslinked Lesion Bypass Efficiency

| Alkyl Modification | Bypass Efficiency | Mutation | Cell |
|--------------------|-------------------|----------|------|
| γ-HOPA mediated peptide crosslink | G>T 5%, G>C 3% [20] | Human |
| γ-HOPMA mediated peptide crosslink | Not mutagenic [20] | Human |
| 5fC mediated peptide crosslink | C>T 7%, C>G 1%, C del 2% [73] | Human |
**Table 1. Cont.**

| Other Nucleotide Analog | Bypass Efficiency | Mutation | Cell   |
|-------------------------|-------------------|----------|--------|
| H                       | 5% [74]           | T>A 41%, T>C 5%, T>G 4%, −1 del 13% [74] | E. coli |
| F                       | 13% [74]          | T>A 9%, T>C 1%, T>G 1% [74] | E. coli |
| L                       | 20% [74]          | T>A 5% [74] | E. coli |
| B                       | 12% [74]          | T>A 24% [74] | E. coli |
| I                       | 10% [74]          | T>A 46%, T>C 1%, T>G 1%, −1 del 6% [74] | E. coli |
| KP1212                  | 128% [57]         | C>T 10% [57] | E. coli |
| xG                      | 11% (45% SOS) [75] | G>A 95% [75] | E. coli |
| xA                      | 80% (108% SOS) [75] | <1% [75] | E. coli |
| xT                      | 73% (102% SOS) [75] | T>A 73% [75] | E. coli |
| xC                      | 29% (33% SOS) [75] | C>A 10% [75] | E. coli |
| α-dG                    | 3% [76]           | G>A 60%, G>C 6% [76] | E. coli |
| α-dA                    | 20% [76]          | Not mutagenic [76] | E. coli |
| α-dT                    | 1% [76]           | Not mutagenic [76] | E. coli |
| α-dC                    | 1% [76]           | C>A 72% [76] | E. coli |
| dxG                     | 25% [77]          | Not mutagenic [77] | E. coli |
| dxA                     | 75% [77]          | A>G 10% [77] | E. coli |
| dxT                     | 150% [77]         | Not mutagenic [77] | E. coli |
| dxC                     | 125% (CXT), 175%(GXG) [77] | Not mutagenic [77] | E. coli |
| sG                      | 98% [78]          | G>A 11% [78] | E. coli |
| sH                      | 98% [79]          | G>A 8% [79] | Human |
| s8mG                    | 91% [78]          | G>A 94% [78] | E. coli |
| sO2HG                   | 95% [79]          | G>A 40% [79] | Human |
| 2′-F-G                  | 87% [78]          | G>A 77% [78] | E. coli |
| J                       | 52% [60]          | Not mutagenic [60] | Human |

**Figure 2.** Structures of oxidative lesions.

**Figure 3.** Structures of alkyl modifications (continued on the next page).
Figure 2. Structures of oxidative lesions.

Figure 3. Structures of alkyl modifications (continued on the next page).

Figure 3. Structures of alkyl modifications.

Figure 4. Structures of bulky and crosslinked lesions.

Figure 3. Structures of alkyl modifications. Cont.
Toxics causing $35\%$ dihydro cyclo deoxyguanosine Cyanuric mutagenicity its mutagenic $2.1$. of mainly yielding in cells causes but $72\%$. 

Figure 3. Structures of alkyl modifications. Cont.

Figure 4. Structures of bulky and crosslinked lesions.

Figure 5. Structures of other nucleotide analogs.
1-Methyldeoxyguanosine (m1G) is a strong replication block either with or without the repair enzyme AlkB (85% and 97%); it mainly causes ∼3% G>T mutation in WT E. coli, which increases to more than 50% in AlkB− E. coli (AlkB: alkyl DNA adduct direct reversal of damage repair protein) [9,33]. N2-methylguanine (m2G) weakly blocks replication by 10% in E. coli, there is no significant change when knocking out either AlkB or DinB (DinB: DNA polymerase IV), and a small amount of G>A mutation (3%) is seen [42]. N2-ethylguanine (e2G) does not block replication in E. coli and causes a low amount of G>A mutation (2%); eliminating AlkB and DinB does not change the replication bypass and mutagenicity significantly [42]. N2-carboxymethyl-2′-deoxyguanosine (N2-CdG) is a stable structural analog to the abasic site (AP site), which is not stable in E. coli. Tetrahydrofuran (THF) is a stable structural analog to the apurinic/apyrimidinic site (AP site), which is not stable in E. coli.
(N²-CMdG) and N²-(1-carboxyethyl)-2′-deoxyguanosine (N²-CEdG) do not block DNA replication and are not mutagenic in WT mammalian cells; however, each of them causes G>A (23%) and G>T (15%) mutations in mouse embryonic fibroblast (MEF) cells that are deficient in pol k [43]. N²-CEdG blocks replication in E. coli [44]. The R-N²-CEdG is a stronger replication block (61%) than S-N²-CEdG (25%); however, neither of them are mutagenic [44]. N²-furfurylguanine (N²-FF-dG) does not block replication in WT E. coli; however, it blocks replication about 72% in DinB cells [42]. It is not very mutagenic with or without DinB [42]. 2-Tetrahydrofuran-2-yl-methylguanine (N²-HF-dG) is similar in structure to N²-FF-dG and strongly blocks replication (72%) only when DinB is knocked out, and causes only 2% G>C mutation [42]. O²-methylguanine (O²mG) is very mutagenic and leads to almost 100% G>A mutation in Ada/Ogt/UvrB triple knockout E. coli (Ada/Ogt: alkyl DNA adduct direct reversal of damage repair protein; UvrB: nucleotide excision repair) [45,46]. N,Nitroso compounds induce DNA lesions: O⁶-pyridyloxobutyl-dG (O⁶-POB-dG), O⁶-pyridylhydroxybutyl-dG (O⁶-PHB-dG), O⁶-carboxymethyl-dG (O⁶-CMdG), which have two structural analogs: O⁶-aminocarboxymethyl-dG (O⁶-ACM-dG) and O⁶-carboxymethyl-dG (O⁶-CM-dG), which have two structural analogs: O⁶-a-aminocarboxymethyl-dG (O⁶-ACM-dG) and O⁶-HOEt-dG (O⁶-ACM-dG) and O⁶-CM-dG. Both O⁶-ACM-dG (2% bypass) and O⁶-HOEt-dG (15% bypass) strongly block DNA replication [47]. They also induce G>A mutation with 30% and 40% frequencies, respectively [47]. Major acrolein-dG adducts include 8α and 8β isomers of 3H-8-hydroxy-3-β-(D-2′-deoxyribofuranosyl)-5,6,7,8-tetrahydroxypro[3,2-a]purine-9-one (γ-OH-PdG), 6α and 6β isomers (α-OH-PdG), and 1,N²-(1,3-propano)-2′-deoxyguanosine (PdG) [12]. The bypass efficiency for γ-OH-PdG is 73% compared to dG control in human cells, and γ-OH-PdG is not very mutagenic (<1%) [12]. α-OH-PdG strongly blocks DNA replication with a bypass efficiency of 17% in human cells and it causes G>A mutation (11%) mutation [13]. PdG strongly blocks replication in human cells and mainly causes 6% G>T mutation [12]. Most of the derivatives of PdG moderately block DNA replication in human cells and cause mainly G>T mutation (2–8%) [81]. 1,N²-etheno(1,N²-eg) is a strong replication blocker (96%) in E. coli and causes G>A and G>T mutation by 6% for both, plus a small amount of G>C (2%) mutation; it also causes –1 and –2 frame shift mutations (5%), and knocking out AlkB leads to higher replication block and almost doubles the mutagenicity [8]. 2′-Fluoro-N²,3,3′-deoxyarabinoguanosine (2′-F-N²,3-eG), a stable analog of N²,3-etheno(1,N²-eg), blocks replication by 79%, and causes 30% G>A mutation in E. coli, with AlkB having no significant influence in its replication bypass and mutagenicity [8].

1-Methyledoxyadenosine (m1A) strongly blocks replication in AlkB- E. coli (88%), but it is not very mutagenic, causing <1% A>T mutation; m1A does not block replication in AlkB+E. coli cells [9]. 1,N⁶-ethenoadenine (eA) weakly blocks replication by 4% in WT E. coli, but significantly blocks replication (95%) when AlkB is knocked out; likewise, eA is not mutagenic in WT E. coli, but shows strong mutagenicity in AlkB- cells (25% A>T mutation) [33,49]. Bypass efficiency of eA in human cells is 17% [50]. 1,N⁶-ethenoadenine (eA) does not block replication in WT E. coli, but strongly blocks replication by 86% when AlkB is removed; it is not very mutagenic in either WT or AlkB- cells, causing only 2% A>C mutations [49]. N⁶-carboxymethyl-2′-deoxyadenosine (N⁶-CMdA) minimally blocks replication in E. coli and is not mutagenic [36]. S,N⁶-HB-dA (HB = 2-hydroxy-3-butene-1-yl) and R,R,N⁶,N⁶-DHB-dA (DHB = 2,3-dihydroxybutan-1,4-diy) do not block DNA replication and are not mutagenic in E. coli [51]. S,S,N⁶,N⁶-DHB-dA moderately inhibits replication with a 60% bypass efficiency, and causes minimal 1% A>G mutation [51]. R,S,1,N⁶-NHMP-dA (NHMP = 2-hydroxy-3-hydroxymethylpropan-1,3-diy) strongly inhibits DNA replication but causes only 2% A>T mutation [51].

O²-Methylthymidine (O²-Me-dT) can be bypassed by 55% in human cells and mainly causes T>A mutation (56%) [53]. O²-[4-(3-pyridyl-4-oxobut-1-yl)thymidine (O²-POB-dT) exhibits genotoxicity showing 26% bypass efficiency and is mutagenic with 47% T>A transversion [53]. Both O²-Me-dT
and $O^2$-POB-dT strongly block DNA replication in *E. coli* (95% and 97%) [54]. $O^2$-Me-dT induces 10% T$\rightarrow$A and 10% T$\rightarrow$G mutations [54]. $O^2$-POB-dT induces 38% T$\rightarrow$G and 12% T$\rightarrow$A mutations [54]. $O^2$-Ethylthymidine ($O^2$-EtTdT) is a strong replication block (79%) in *E. coli*, and knocking out pol IV increases the blocking activity, while knocking out pol V increases the replication block even more [55]. It is very mutagenic and forms T$\rightarrow$C (35%), T$\rightarrow$A (15%), and T$\rightarrow$G (5%) mutations, and mutation frequency drops when pol V is knocked out [55]. The bypass efficiency of $O^2$-dT alkyl adducts in *E. coli* depends on the size of the alkyl lesion [82]. More than 20% of adducts can be bypassed during replication for ethyl and methyl substitutions, but less than 10% can be bypassed for propyl, and less than 5% for butyl adducts, with the major mutation type being T$\rightarrow$C point mutation [82]. $O^2$-alkyldT lesions strongly inhibit DNA replication (40–85%) in mammalian cells [52]. The blockage effect increases with the size and branching of the alkyl groups [52]. These lesions cause T$\rightarrow$A and T$\rightarrow$G mutations [52].

3-Methyldeoxycytidine (m3C) strongly blocks replication in *E. coli* by 94% and is very mutagenic, generating mainly T$\rightarrow$A (32%) transversion mutation; eliminating AlkB slightly increases its replication blocking power and mutagenicity [9]. N3-Ethylthymidine (N3-EtdT) strongly blocks replication by 83% in *E. coli*, and knocking out pol V or pol IV increases its blocking activity; it is very mutagenic causing T$\rightarrow$A (21%), T$\rightarrow$C (15%) and T$\rightarrow$G (3%) mutations, and removing pol V eliminates the mutagenicity of this adduct [55]. N3-carboxymethylthymidine (N3-CMdT) strongly blocks replication by 45% in *E. coli*, with the major mutation being T$\rightarrow$A (66%); and knocking out pol V slightly increases the mutation rate; however, knocking out pol IV decreases the mutation rate [36]. $O^4$-carboxyethylthymidine ($O^4$-CMeTdT) is a strong replication block (51%) and very mutagenic, causing 86% T$\rightarrow$C mutation [36]. N3-CMdT, $O^4$-CMeTdT and $O^6$-carboxymethyl-dG ($O^6$-CMdG) moderately block DNA replication in human cells [48]. $N^3$-CMeTd causes T$\rightarrow$A (81%) mutation; $O^4$-CMeTd causes T$\rightarrow$C (68%) mutation; $O^6$-CMdG causes G$\rightarrow$A (6.4%) mutation; neither $N^6$-CMdA nor $N^4$-CMeDc block replication or induce mutation [48]. $O^4$-Ethylthymidine ($O^4$-EtTdT) does not strongly block replication (24%) in WT *E. coli*, but it cannot be efficiently bypassed in pol II/IV/V triple knock out cells [55]. The major mutation of $O^4$-EtTdT is T$\rightarrow$C (84%) transition; however, it does not cause mutations in *E. coli* lacking pol V [55]. $O^4$-Alkylthymidine ($O^4$-alkyldT) lesions moderately block DNA replication in human cells; pol $\zeta$ and pol $\iota$ promote the bypass of all $O^4$-alkyldT lesions except $O^4$-MedT [56]. The $O^4$-alkyldT lesions induce only T$\rightarrow$C transition mutations in cells [56].

3-Methyldeoxyctydine (m3C) has been demonstrated to strongly block replication (>90%) and generate mainly C to T (50%) and C to A mutations (30%) in the AlkB- *E. coli* cell [9]. However, the lesion is not mutagenic and not blocked by the replicative polymerases in the WT (*AlkB*+ cell) [9]. 3-Ethyldeoxyctydine (e3C) does not block replication in *E. coli*; however, it dramatically blocks replication when knocking out AlkB (91%) [9]. e3C causes 17% C$\rightarrow$T, 11% C$\rightarrow$A, and 2% C$\rightarrow$G mutations in AlkB- *E. coli*, but is not mutagenic in WT cells [9]. The m3C, e3C, and m1A lesions presumably have their methyl or ethyl groups removed by AlkB’s direct reversal of DNA alkyl damage mechanism prior to encountering the DNA polymerase [9]. $N^4$-carboxymethyl-2'-deoxyctydine ($N^4$-CMeCyd) weakly blocks replication (17%) and is not mutagenic in *E. coli* [36]. 5-Methylcytosine (5mC) and its derivatives 5-hydroxymethylcytosine (5hmC), 5-formylcytosine (5fC) and 5-carboxylcytosine (5caC) neither block replication nor cause mutation in *E. coli* [39,59]. 5mC also does not block replication in human cells, but there are some blockades of 5hmC (5%), 5fC (25%), and 5caC (28%) towards DNA replication in human cells [58]. 3,$N^4$-ethenocytosine (eC) is a toxic adduct, which strongly blocks replication (76%) and leads to mutation with a pattern of dominant C$\rightarrow$A (24%) and less C$\rightarrow$T (11%) mutations in WT *E. coli*; in AlkB- cells, the blockage of replication increases to 87% and mutagenicity rises up to 49% C$\rightarrow$A and 31% C$\rightarrow$T mutations [33]. Lipid peroxidation-derived product 4-oxo-2(E)-nonenal reacts with dG, dA, and dC in DNA to form heptanone (H)-etheno (e) adducts [50]. H-edC shows strong DNA replication blocking in both *E. coli* (99%) and human cells (90%) [50]. It causes mainly C$\rightarrow$G (40%) mutation in *E. coli*; however, mostly C$\rightarrow$A (60%) and C$\rightarrow$T (32%) mutations are seen in human cells [50].

5-Hydroxyethyluracil (5hmU) blocks replication by 20%, but it is not mutagenic in human cells [60]. The *S*$_p$ alkyl phosphotriester (*S*$_p$-alkyl-PTE) lesions display comparable replication
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bypass efficiency to unmodified DNA in E. coli; Sp-Me-PTE is mutagenic causing TT>GT (50%) and TT>GC (15%) mutations [61]. In contrast, Rp-alkyl-PTEs block DNA replication (30–70%) but are not mutagenic [61]. Interestingly, nPr- and nBu-PTEs exhibit higher bypass efficiencies than Me- and Et-PTEs [61].

2.3. Bulky Lesions

All the structures of modifications covered in this section are displayed in Figure 4. N-(deoxyguanosin-8-yl)-1-aminopyrene (C8-AP-dG) moderately blocks DNA replication in human cells [66]. N-acetyl-2-aminofluorene (C8-AAF-dG) strongly blocks replication [66]. 2-Aminofluorene (C8-AF-dG) slightly blocks replication [66]. All three adducts can be nearly bypassed in error free manner [66]. Aristolochic acids I and II (AA-I, AA-II) are found in all Aristolochia species and generate the aristolactam (AL) metabolite for forming DNA adducts with dA and dG. Both AL-II-dA and AL-II-dG strongly block DNA replication in MEF cells [63]. AL-II-dA causes 22% A(BP)-7,8-diol-9,10-epoxide-(C8-AF-dG) slightly blocks replication [66]. All the three adducts can be nearly bypassed in error free manner [66]. Aristolochic acids I and II (AA-I, AA-II) are found in all Aristolochia species and generate the aristolactam (AL) metabolite for forming DNA adducts with dA and dG. Both AL-II-dA and AL-II-dG strongly block DNA replication in MEF cells [63]. AL-II-dA causes 22% A>T mutation and AL-II-dG causes 9% G>T transversion [63]. Knocking out the rev3L gene dramatically suppresses bypass of AL-I-dA in MEF cells and abolishes A>T transversion [67]. Benzo[a]pyrene (BP)-7,8-diol-9,10-epoxide-N2-deoxyguanosine (BPDE-dG) is an adduct formed by benzo[a]pyrene; it predominantly miscodes with G>T (73%) and G>A (12%) mutations in WT MEF cells [68]. Knocking out rev1 gene decreases the bypass efficiency of BPDE-dG to 40% and changes the mutation frequency to 32% G>T and 18% G>A [68]. Knocking out the rev3L gene significantly decreases the bypass efficiency to 13% and decreases the mutation to 6% G>T [68]. Mitomycin C (MC) generates dG-N2-MC and dG-N2-2,7-Diaminomitosene (DAM) adducts, which can be bypassed 38% and 27% in human cells, respectively [62]. The major type of mutation is G>T mutation (18% for dG-N2-MC and 10% for dG-N2-2,7-DAM) [62]. Aflatoxin B1-N7-dG adduct (AFB1-N7-dG) is weakly mutagenic in E. coli, causing 1.5% G>T mutation [64]; and its FAPY adduct causes 14% G>T mutation [65].

2.4. Crosslinked Lesions

All the structures of modifications covered in this section are displayed in Figure 4. N2-guanine-N2-guanine interstrand crosslinks (ICLs), 3-(2-deoxyribos-1-yl)-5,6,7,8-(N2-deoxyguanosyl)-6(both R or S)-methylpyrimido[1,2-R]purine-10(3H)-one is a product induced by acetaldehyde/crotonaldehyde [69]. ICL-S and ICL-R moderately inhibit DNA replication in WT E. coli; however, their replication blocking effects increase in uvr- E. coli cells [69]. ICL-Rd is a moderate block in WT E. coli, but it almost completely blocks replication in uvr- cells [69]. All three lesions are weakly mutagenic in E. coli causing exclusively 5'-G>T (3%) transversions; no mutation is observed at the 3'-G site [69]. Similar mutations generated by these lesions are seen in human cells, except ICL-S has a slightly higher mutation frequency (6%) [69]. The crosslinks formed by cis-diaminedichloroplatinum (II) (cis-DDP, cisplatin) between two guanines or adenine-guanine strongly block DNA replication in E. coli, but they are not very mutagenic [72]. 5-Formylcytosine mediated peptide crosslink causes 7% C>T and 1% C>G mutation and 2% C deletion [73]. y-Hydroxypropanodeoxyguanosine (y-HOPdG) mediated crosslink between peptide and guanine is mutagenic, causing 5% G>T and 3% G>C mutations; however, the crosslink between peptide and y-hydroxypropanodeoxyadenine (y-HOPdA) is not mutagenic [20].

2.5. Other Nucleotide Analogs

All the structures of modifications covered in this section are displayed in Figure 5. A series of unnatural analogs of thymine (T) was developed by the Kool group to probe the biological requirements for DNA polymerases [74]. 3-Toluene-1-β-D-deoxyribose (H) strongly blocks replication (95%) and is very mutagenic causing T>A (41%), T>C (5%), and T>G (4%) point mutations and -1 frame shift mutation (13%). 2,4-Difluoro-5-toluene-1-β-D-deoxyribose (F) strongly blocks replication (87%) and is mutagenic causing T>A (9%), T>C (1%), and T>G (1%) mutations. 2,4-Dichloro-5-toluene-1-β-D-deoxyribose (L) strongly blocks replication (80%) and is slightly mutagenic causing T>A (5%) mutation.
(B) strongly blocks replication (88%) and is mutagenic, causing T>A (24%) mutation. 2,4-Diiodo-5-toluene-1-β-D-deoxyribose (I) strongly blocks replication (90%) and is very mutagenic causing T>A (46%), T>C (1%), and T>G (1%) point mutations and −1 frame shift mutation (6%) [74]. xG is an ‘expanded base’ of dG (retaining the hydrogen-bonding face), which strongly blocks replication (89%) and is very mutagenic, causing G>A (95%) mutation [75]. xA (expanded A) weakly blocks replication (20%) and is not mutagenic; xT (expanded T) weakly blocks replication (27%), but is very mutagenic, causing T>A (73%) mutation; xC (expanded dC) strongly blocks replication (71%) and is mutagenic, causing C>A (10%) mutation [75].

The α-anomer of deoxynucleosides (α-dN) can be generated as a result of hydroxyl radical attack on deoxyribose [76]. All α-dNs except α-dA strongly block replication in E. coli [76]. α-dC blocks almost 99% replication and causes 72% C>A mutation [76]. α-dG also strongly blocks replication and causes 60% G>A mutation [76]. α-dT blocks almost 99% replication but it is not mutagenic in WT E. coli [76]. α-dA is not mutagenic [76]. The anticancer agent 6-thioguanine (sG) and its derivative Sα-methylthioguanine (SαmG) do not block replication strongly in both E. coli and human cells [78]. sG causes 11% G>A mutation and SαmG causes 94% G>A mutation in E. coli [78]. sG is less mutagenic (8%) than SαmG (40%) in human cells as well [78]. Guanine-Sα-sulfonic acid (SO3HsG) is another derivative of sG [78]. It is not a strong replication block in E. coli, but it is very mutagenic, causing 77% G>A mutation [78]. The anti-HIV drug KP1212 is an analog of deoxycytidine [57]. It does not block replication in E. coli, but is mutagenic causing 10% C>T mutation [57]. Among the four 2′-deoxyxylonucleosides (xN), only xA and xG exhibit a replication block in E. coli [77]. xA is the only mutagenic lesion among the four and causes 10% A>G mutation [77]. Base J strongly blocks replication by 48%, but is not mutagenic in human cells [60].

3. Perspectives

In this review, we survey the biological effects of various DNA lesions or biomarkers studied by the shuttle vector techniques, allowing one to gain insight into how DNA damage or other chemically defined nucleobases are processed by polymerases and repair machinery in a natural cellular environment under physiological conditions. Among the new methods that have been developed or applied in the last decade, MS-based strategies and NGS methods have been demonstrated to be efficient for analyzing the lesion’s biological outcomes. LC-MS-based methods are sensitive and accurate for quantifying the degree of lesion bypass and point mutations [4,5]. NGS techniques allow for a large-scale population analysis on many samples at the same time and provide information on a genomic perspective [4,8]. Another possible direction for using vectors as probes to analyze biomarkers is to study the mutational spectrum or mutational signature of a certain chemical or damaging agent [83–86]. LC-MS- and NGS-based analyses not only consider the biological consequences at the lesion site, but also incorporate information from the neighboring bases, such as one or two nucleotides next to the lesion site from both the 5’ and 3’ direction. An oligonucleotide containing the modified base can be made surrounded by nearest (and next-to nearest) randomized bases and ligated into a shuttle vector. While cellular analysis may pull out a hotspot consensus sequence for poor repair and/or mutagenic replication, this will not answer the primary question of contextual bias in adduct formation. Shuttle vector systems whereby the vector is treated with the chemical to be assessed, followed by quantification of adduct type and amount, and transfection into isogenic cells of varying repair and/or replication backgrounds may tease apart the contribution of local sequence environment to adduct formation, repair, and replication. Such vectors were used over a decade ago [87], and coupled with NGS throughput and bioinformatics, may provide enough reads to make statistically significant claims. Shuttle vectors are currently, to our knowledge, mainly DNA-based; however, one can envision use of RNA-based vectors to study the effect of modified RNA bases on cellular processes such as viral replication, translation, reverse transcription, and possibly even repair. While the role of DNA damage in toxicology focuses mainly on the direct addition of chemical damage to DNA, pool mutagenesis has often been overlooked, and it would be interesting to leverage shuttle vector techniques to study
the incorporation of modified bases from the nucleotide pool in the form of damaged DNA or from DNA-based therapeutics.

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