Communication

Current and future methodology for quantitation and site-specific mapping the location of DNA adducts

Gunnar Boysen\textsuperscript{1,2},* Intawat Nookaew\textsuperscript{2,3}

\textsuperscript{1} Dept. Environmental and Occupational Health, Fay W. Boozman College of Public Health, University of Arkansas for Medical Sciences
\textsuperscript{2} The Winthrop P Rockefeller Cancer institute, University of Arkansas for Medical Sciences
\textsuperscript{3} Dept. Biomedical Informatics, College of Medicine, University of Arkansas for Medical Sciences
* Correspondence: egboysen@uams.edu

Abstract: Formation of DNA adducts is a key event for a genotoxic mode of action and their formation is often used as surrogate for mutation and cancer. Interest in DNA adducts are twofold, first, to demonstrate exposure, and second, to link DNA adduct location to subsequent mutations or altered gene regulation. Methods have been established to quantify DNA adducts with high chemical specificity, and to visualize the location of DNA adducts, elegant bio-analytical methods have been devised utilizing enzymes, various chemistries, and molecular biology methods. Traditionally, these highly specific methods cannot be combined, and the results are incomparable. Initially developed for single-molecule DNA sequencing, nanopore-type technologies are expected to enable simultaneous quantitation and location of DNA adducts across the genome. Herein, we will briefly summarize the current methodologies for state-of-the-art quantitation of DNA adduct levels and mapping of DNA adducts and describe novel single-molecule DNA sequencing technologies that are expected to achieve both measures simultaneously. Emerging technologies are expected to soon provide a comprehensive picture of the exposome and identify gene regions susceptible to DNA adduct formation.

Keywords: DNA adducts; nanopore; Oxford Nanopore Technology; mass spectrometry; adductomics; exposome

1. Introduction

Chemical carcinogenesis. Although DNA is a very stable molecule for storing biological information, it is under relentless attack by reactive compounds of endogenous and exogenous origin that covalently bind to DNA, forming so called DNA adducts \cite{1}. Since identification of the first DNA adduct by Reiner, B. and Zamenhof in 1957 \cite{2}, several thousand studies on DNA adducts have been reported and are reviewed from different perspectives \cite{3–10}. The ability of a compound to form DNA adducts, directly or after metabolic activation, is considered a critical event in chemical carcinogenesis \cite{11} and a key event for genotoxic mode of action of toxicants \cite{12,13}. The binding to DNA, has been widely used as biomarker of exposure in molecular epidemiology studies to link exposure to adverse health outcomes \cite{14–16}. Formation and stability of specific promutagenic DNA adducts has been established \textit{in vitro}, in cell cultures, animal experiments and molecular epidemiology studies \cite{5,6,17}. Recent advances in technology, especially in mass spectrometry and single molecule sequencing, allows for ‘omics’ type monitoring of all DNA adducts, DNA adductomics, and is expected to provide unprecedented insight into the total exposome \cite{18–21}.

This perspective summarizes the current methodologies for quantitation of DNA adduct levels and mapping the location of DNA adducts along the genome and highlight novel single-molecule analysis expected to be capable for achieving both measures.
2. Methodology

Historically studies on DNA adducts have mirrored advances in analytical and bio-analytical chemistry technologies, starting with paper chromatography, enzyme-linked immunosorbent assay, $^{32}$P-Post-labeling [22], followed by liquid [23] and gas chromatography separations with various detection systems, including UV, fluorescence and electrochemical detectors [24,25].

![Scheme 1](image.png)

**Scheme 1.** Overview of current approaches for measuring DNA adduct levels and mapping the location of DNA adducts. Left: DNA digestion or release of DNA adducts, prior to quantitation by LC-MS. Middle: Labeling of DNA adduct sites prior to localizing by amplification-base sequencing. Right: Single molecule sequencing using nanopore-based technologies.

2.1. Mass Spectrometry-based DNA Adduct quantitation.

The introduction of electrospray ionization by Dole and others, in 1968 [26] revolutionized mass-spectrometry and biochemistry [27]. Electrospray mass-spectrometry is now a commonly used technique for qualitative and quantitative analyses of any type of compound, including DNA adducts [24,25]. Improvements of mass analyzers allows monitoring of thousands of molecules simultaneously with ultra-high mass resolution and accuracy [28–31]. The basic and most frequently used approach for quantitation of DNA adducts is to isolate DNA, liberate the DNA adducts from DNA by chemical means or enzyme hydrolysis and quantify the released nucleotide- or base-adducts by LC-MS (Scheme 1 left) [32]. To improve sensitivity various sample enrichment procedures maybe included such as solid phase and liquid-liquid extraction or pre-separation by HPLC [3]. If needed, stable isotope standards are added at the beginning to account for any potential loss during sample workup and to improve measurement accuracy [33,34].

2.2. Adductomics, nontargeted screening of DNA adducts.

While most studies apply targeted mass spectrometry using authentic DNA adduct standards, efforts are underway to move towards nontargeted ‘omics’ type screening of DNA adducts to obtain a complete measure of the exosome [35]. Therefore, DNA adducts are monitored in various modes of data dependent (MS/MS) or multistage (MS$^n$) scanning modes [19,20,36].

Independent of targeted or nontargeted approach, results are reported as number or concentration of DNA adduct per DNA, or the corresponding unmodified nucleotides (e.g. fmol/mg DNA, fmol / µmol dG). The mass spectrometry-base detection provides a high level of chemical specificity but no site-specificity.
2.3. Amplification-based Mapping of DNA Adducts.

Various methods have been developed for genome-wide and site-specific mapping of DNA damage (Scheme 1 middle). The general approach is based on mapping sequencing stop sites to localize the adducts [37–39]. Therefore, DNA adducts are first recognized or modified by enzymatic or chemical means, taking advantage of DNA repair enzymes to mark and excise DNA adducts.

The marked or cleaved DNA is then amplified, sequenced and the location of DNA adducts are obtained from strand ends or mismatched base pairs [37–39]. For example, Denissenko et al. used UvrABC excision nuclease, in combination with ligation-mediated polymerase chain reaction (PCR), to map the sites of bulky DNA adducts [40–42]. More recently, cyclobutene pyrimidine dimers (CPDs) were mapped with CPD-seq based on DNA cleavage mediated by T4 endonuclease V and APE1 [43,44]. XR-seq [45] and tXR-seq rely on TFIIH-mediated enrichment of damage-containing fragments cleaved by mammalian nucleotide excision repair enzymes [46–48]. Cisplatin-seq takes advantage of the HMG box A of HMGB1 protein’s preferentially binding to distorted DNA structures for selective enrichment of cisplatin-modified DNA [49]. Further, click chemistry, Click-Code-Seq, has been successfully applied to label 8-oxo-dG or 5-hm-dC sites in DNA prior to next generation sequencing [50,51]. DNA adduct mapping methods have been reviewed extensively [52–54].

Results are given as modified versus unmodified sites or motifs with high level of site-specificity. Unfortunately, these elegant methods are limited by (i) the breadth of enzyme specificity, which may identify a mixture of DNA adducts; (ii) excision of short DNA fragments that sometimes cannot be aligned with absolute certainty; (iii) reliance on completion of chemical reactions; (iv) restricted applicability to one type or class of DNA adducts; (v) the inability to distinguish different DNA adducts; and (v) limited chemical specificity.

2.3. Single Molecule DNA sequencing.

A relatively new method for analysis of DNA adducts is based on single molecule sequencing. This novel technology sequences and counts single DNA molecules, whether they are whole genomes or DNA fragments.

2.3.1. Nanopore Technology.

Nanopore-type technology utilizes electrochemical forces to pull single-stranded DNA in native form through tiny pores (Figure 1 right). The accompanying changes in electric current indicate the physicochemical properties of the DNA bases transiting through the pore, revealing the DNA sequence and potential DNA adducts [55]. A DNA adduct modulates the nanopore ion current signal while entering, passing through, and exiting the nanopore. This results in an electric current signature characteristic for the DNA adduct within a given 7-base sequence that includes the DNA adduct and the three adjacent 3’ and 5’ nucleotides that reside in the nanopore [55].

Burrows and colleagues pioneered the application of nanopore-type technology for sequencing DNA adducts in single-stranded DNA. With custom-made solid-state or protein-based nanopores, they showed the proof-of-principle for detecting N²-BPDE-dG-induced adducts [56], abasic sites [57–59], and other DNA adducts [60], including 8-oxo-dG [61–63].

2.3.1.1 Oxford Nanopore Technologies (ONT)

Using a similar principle, ONT developed and commercialized a nucleic acid sequencing technology that has the capability to sequence long to ultra-long molecules of DNA (> 2 Mb) in the native form, preserving the sequence position of the DNA adducts [64–66].

2.3.1.2 PacBio DNA sequencer.
While technical not a nanopore system, the PacBio RSII DNA sequencer also employs the single-molecule real-time (SMRT) sequencing principle. The DNA strands are converted into loops and amplified by polymers. The unique time needed by the polymerase for elongation of the DNA is indicative of the base added and DNA sequence.

While ONT uses the disturbance in the ion signals, the PacBio system makes use of the time delay of the polymerase to identify DNA sequence. The standard base calling algorithms for both systems are optimized for the four main bases (A, T, G and C), and report errors when encountering unknown bases, potential DNA adducts. Therefore, efforts are under way to expand base calling algorithms to enable detection of DNA adducts and epigenetic modifications. Multiple groups successfully applied ONT for genome wide detection of epigenetic modifications, such as 5-methyl-2'-deoxycytidine (5mdC) and N6-methyl-2'-deoxyadenosine (N6mdA) [67–70]. Similarly, the PacBio system has been shown to be suitable for simultaneous detection of N6mdA, 5mdC and 5-hydroxymethylcytosine (5hmdC) [71]. The latter, is also capable for detection of the unique phosphothioate modifications of the phospho-ribose backbone, found in some bacteria [72,73]. Expansion of these commercial platforms to high abundant epigenetic marks is the first step to enable them to detect any DNA modifications, including DNA adducts derived from endogenous or exogenous sources.

2.3.2. Data analyses
2.3.2.1 ONT/ELIGOS.

Our team developed the Epitranscriptional/ Epigenomical Landscape Inferring from Glitches of ONT Signals (ELIGOS) software that uses ONT data to simultaneously detect RNA and DNA modifications, including DNA adducts [55,74,75]. The ONT/ELIGOS platform is a powerful tool for (i) detecting DNA adducts and (ii) discriminating DNA adducts of different sizes, regiochemistry, and functional groups [55]. ELIGOS takes this error information from the standard base-calling algorithm and calculates the odds ratios at each site as an indicator for a potential DNA adduct (Figure 1)[55].

![Figure 1](https://example.com/figure1.png)

**Figure 1.** Representative Odds ratios plot of a plasmid carrying a DNA adduct at position G640.

Next, ELIGOS generates a radar plot displaying the multiplex disturbances of the raw ONT signal from the DNA adduct in relationship to the dG-containing control plasmid (Figure 2a)[55]. The radar plot shows an 11-base sequence that covers the DNA adduct along with 5 preceding and 5 trailing deoxynucleotides that dwell in the nanopore during sequencing [55,76]. These radar plots are characteristic for each DNA adduct at each given position. The radar plots are used to identify the type of DNA adduct by comparing them to plots of standard DNA that contains the DNA adduct at the same sequence position and within the same sequence context (Figure 2a). Disturbances in the raw ONT signal are used for multivariant statistical analyses to obtain a measure of separation, as shown in Figure 2b.
2.3.3. CRISPR/cas-9 targeted sequencing.

DNA adduct measurement by single molecule sequencing technologies will generate a huge amount of data because the DNA adduct levels are extremely low. In principle one would need to sequence the whole genome 10^8 times to get a DNA adduct level at each site of 1 adduct/10^8 nnt as commonly reported. However, drawing from previous mapping approaches using CRISPR/cas-9 targeted sequencing will increase the number of reads and thereby sensitivity at the sites of interest such as cancer driver genes or mutation hotspots [77–79]. The CRISPR/cas-9 targeted sequencing may target DNA segments of 1,000 to 10,000 base pairs in length that can be read by ONT as a single molecule.

2.3.4. Limit of Detection.

The results of these single molecule analyses are number of DNA adduct X at Site Y per total number of DNA molecules / DNA molecules analyzed containing the site of interest, including unmodified DNA and DNA molecules carrying mutations or modifications at different sites. In theory a DNA segment of 2000 base pairs length that has been read 100,000 times has a limit of detection of;

- 1 adduct at a given site/ per 10^5 unmodified bases at position Y (e.g alkyl-dG/10^5 dG at position Y),
- 5 adducts per 10^9 unmodified nucleotides (e.g alkyl-dG/10^9 nnt),
- or approximately 1 adducts per 10^9 corresponding nucleotides (e.g. alkyl-dG/10^9 dG).

This theoretical limits of detection of ONT/ELIGIOS are in the range of the levels reported for endogenous and exogenous DNA adducts.

3. Discussion

Below we highlight some selected studies showing the application and utility of DNA adduct research.

3.1. DNA adduct levels.
The first question in the DNA adduct field was, and still is, to demonstrate the formation of exposure induced DNA adducts. For example, Benzo(a)pyrene (BaP) a ubiquitous environmental and occupational carcinogen, and tobacco smoke constituent has been studied extensively. BaP requires metabolic activation to the reactive 7,8,9,10-diepoxy-9,10-dihydroxy-benzo[a]pyrene (BPDE) that ultimately forms the promutagenic N2-BPDE-dG adduct. Applying an ultra-sensitive LC-MS method, with a limit of detection of one N2-BPDE-dG adduct per 10^11 nucleotides (1 adduct per 10 human cells) levels in lung of smoker and nonsmoker DNA were 3.1 and 1.3 N2-BPDE-dG adducts per 10^11 nucleotides, respectively [80].

With increases in sensitivity of the methodologies, it became apparent that DNA adducts are also formed by reactive compounds of endogenous origin, such as the 8-oxo-7,8-dihydroguanine (8-oxo-dG). Oxidative modifications of DNA were first believed to be an artifact during sample handling and processing, which is in part still true, but their presence in all cellular DNA in nowadays well established [81,82]. Endogenous 8-oxo-dG in human peripheral blood lymphocytes measured by LC-MS are about one 8-oxo-dG per 10^6 dG [83,84]. Consequently, endogenous 8-oxo-dG levels are five orders of magnitude higher than the exposure derived N2-BPDE-dG.

Most studies are on base adducts, potentially disrupting the DNA base paring. However, adduct formation at the phosphor-ribose backbone has been proposed long ago and recently been shown for DNA-phosphate adducts formed by the tobacco-specific nitrosamine 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanone [85].

3.2. DNA adductomics.

The emergence of nontargeted adductomics will generate a huge amount of DNA adduct data presumably representing the exposome. Studies of true DNA adductomics are limited, but first reports are very promising. For example, applying a nontargeted ‘omics’ approach, thousands of potential DNA adduct m/z features were observed in human tissues such as lung, bronchia or saliva [86–88]. A challenging task will be to accurately identify the low abundant DNA adducts resulting from exogenous sources that are dwarfed by epigenic marks and endogenous DNA adducts, that are present at levels several orders of magnitude higher [89,90].

3.3. DNA adduct mapping.

The second question in the DNA adduct field was, and still is, to determine the location of exposure-induced DNA adducts. Therefore, the mapping of BaP derived DNA adducts has been reported genome wide [56] and for specific genes such as P53 [91], kRAS and hRAS [92], suggesting DNA adduct formation at mutation hotspots in these genes. BaP treatment however has been shown to also induce oxidative stress which is known to cause multiple DNA lesions recognized by the UvrABC Nuclease, potentially confounding the results [93]. Elegant stable isotope labeled experiments confirmed the preferred binding of BPDE to the mutation hotspot sequence motif in TP53 [94] settling this discussion.

Further, several approaches have been developed for genome wide mapping of 8-oxo-dG with various degrees of resolution (reviewed by Poetsch [54]). The mapping revealed accumulation of 8-oxoG at sites of high nucleosome occupancy in yeast and different types of GC repeats accumulate large amounts of 8-oxoG, particularly telomeres and microsatellites [50], suggesting that DNA adduct formation is not random across the genome. These DNA adduct mapping studies highlight the importance of the DNA adduct location to elucidate subsequent biological outcomes.

The current nanopore-type Single Molecule DNA sequencing technologies do not reach the chemical specificity obtained by mass spectrometry, but efforts are underway to combine nanopore with mass spectrometry [95–97]. Further, since the ion signal disturbances are cause by physio-chemical properties of the DNA molecule transitioning the nanopore,
refined artificial intelligence (AI) assisted data analyses in future may allow chemical specific identifications of the DNA adducts at any location in the genome. This novel technology is still in the implementation phase and future studies are needed to evaluate and establishing chemical specificity, sensitivity and accuracy for measuring DNA adducts.

4. Conclusions

With the current technologies on hand, it’s easy to determine whether DNA adducts form in the target tissue to establish the internal dose derived from external exposure. When the goal is to understand the exposome, mass spectrometry-based adductomics is a method of choice. Detection and quantitation of DNA adducts derived from a mixture of pollutants in the target tissue will unambiguously demonstrate, with high chemical specificity, that the subject has been exposed and the toxicant has reached the tissue of concern.

If the goal is to understand how exposure leads to changes in cell homeostasis and promotes or prevents disease development many DNA adduct mapping approaches are available. These mapping methodologies will enable the investigator to demonstrate, with biological specificity, that the exposure or treatment induces modifications in the DNA at the promoter or gene region of interest.

The new Single Molecule DNA sequencing technologies are expected to be suitable for addressing both questions mentioned above. They will provide a comprehensive picture of the DNA adducts level and their locations across the genome.

Lastly, a novel approach is needed for meaningful interpretation of the DNA adductome-based exposomes and future DNA adduct maps to improve our understand in cancer etiology and explain the origin of mutational signatures established for various tumor types [98].

5. Patents

“Not applicable.”

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Conflicts of Interest:

The authors declare no conflict of interest.
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