Basic Radiation Chemistry and Radiation Chemistry of DNA Damage



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Ernest Rutherford

Nobel for physics 1908, first to split the atom in 1917.

"All science is either physics or stampcollecting"







J.F.Ward, 2007

Time Scale (in seconds) of Radiation Action

- 10⁻¹⁵ Radiation energy is deposited Ionizations occur
- 10-12
- 10⁻⁹ Primary Radical reactions
- 10⁻⁶ Oxygen reacts with radicals
- 10⁻³ Permanent products are formed
- 10⁰ Cell begins to respond biochemically
- 10³ Enzymatic repair Mutation/transformation occur
- 10⁶ Cell death scored Cancer appearance

Mechanisms of Radiation Damage to Cells: Sources of Experimental Data

In vitro Model compounds irradiated in solution or alone

> In vivo Irradiation of living cells

In silico Computer Models of "biophysical" processes

Biophysical models

Important variables are often not accommodated, e.g.

Effects of oxygen

Presence of <u>sensitizers or protectors</u>

Identities of radiation products

Amounts of each damage

Response of cells to the damage

Effects of cell cycle

Genetic differences between cells

Conclusion: These models are limited in their ability to predict effects of variables.

Time Scale (in seconds) of Radiation Action

- 10⁻¹⁵ Radiation energy is deposited
 Ionizations occur
 10⁻²
- 10-9 Ra. Vical reactions
- 10-6 Oxygen Ar's with radicals
- **10-3 Permar.ent pro. 'ucts are formed**
- 10^o Cell begins to respond biochemically

10⁶

Enzymatic repair Mutation/transformation occur Cell death scored Cancer appearance

In silico - Biophysical Models supported by DOE and the European Union

See: Physical and Chemical Mechanisms in Molecular Radiation Biology" (W. Glass and M.N. Varma eds.) Plenum Press, (New York) (1991). Basic Life Sciences 58 (1991).

S. Curtis Mechanistic Models pp. 367-382

L.A. Braby Phenomenological Models pp. 339-361

Sidney Brenner, Science, July 17, 1992

Those who prefer the airy realm of theory to the area of decisive experiment aren't necessarily doing so by choice: I always say it's important to distinguish between chastity and impotence.

Contents of Typical Cell

From B. Alberts, Molecular Biology of a Cell, 4th Edition, Garland Press, 2002.

Cell Size 4 x 10⁻⁹ cm⁻³, Density 1.15 g/cm³ – Mass 4.6 x 10⁻⁹ g



Water Radiolysis

H₂O radiation (ionization) \rightarrow $H_2O^+ + e^-$ (unstable)

 $H_2O^+ + H_2O \longrightarrow H_3O^+ + OH$

e⁻ + bulk water \rightarrow e (H2O)₆⁻

(hydrated electron)

Radical Damage to Intracellular Molecule (RH)

 $-ionization \rightarrow RH^+$ RH (1) H_2O —ionization \rightarrow $H_2O^+ \rightarrow$ OH^* (2) $R^{\bullet} + O_2 \rightarrow RO_2^{\bullet}$ (5) $\mathsf{GSH} \rightarrow \mathsf{RH} + \mathsf{GS}^{\bullet}$ R' + (6) 1 and 2 occur within 10^{-12} s.

5 and 6 in 10⁻⁵ s.

Techniques used in Radiation Chemistry.

Pulse radiolysis

Electron Spin Resonance

Product analysis

Pulse Radiolysis



Source of Rate Constants for Reaction of Free Radicals



Notre Dame Radiation Laboratory, Radiation Chemistry Data Center

http://www.rcdc.nd.edu/index.html

Typical Rate Constants – 'OH radicals

		Liters per mole per second
DNA	bases	5 x 10 ⁹
	deoxyribose	1.2 x 10 ⁹

Amino acids – glycine 1.7×10^7 tryptophan 1.3×10^{10}

Ethanol 1.3×10^9

Dimethyl sulfoxide

6.5 x 10⁹

Typical Rate Constants – e⁻(aq)

Liters per mole per second 1.3×10^{10}

DNA bases 5×10^9 deoxyribose 1×10^7

Oxygen

Amino acids - glycine $1 \ge 10^7$ tryptophan $3.2 \ge 10^8$ cystine $1.1 \ge 10^{10}$

Typical Rate Constants – (O_2^-)

Lite	ers per mole per second <10 ⁶	
Glycine Tryptophan Cystine	<0.42 <24 <0.4	
Superoxide Dismutase	3.2 x 10 ⁹	

•OH radical attack on DNA



Miral Dizdaroglu

Reactions of Hydroxyl Radicals

1. Hydrogen atom abstraction

RH + OH \rightarrow R' + H_2O

2. Addition to a double bond



DNA base alterations produced by 'OH

Fuciarelli et al., Int. J. Radiat. Biol <u>58</u>; 397.

Base Alteration	% of total base damage
8-hydroxyguanine	43
8-hydroxyadenine	7
Formamidopyrimidine-adenine	3
Formamidopyrimidine-guanine	6
Thymine glycol	27
Cytosine glycol	14

Also Double lesions Box et al., Free Radicals Biol. Med. 31; 856.

Characteristics of Single Strand Breaks (SSB)

- Are induced by both direct ionization (35%) and OH radicals (65%). *Roots and Okada Int. J. Radiat. Biol.* <u>21</u> 329.
- 15 % of OH radicals reacting with DNA cause SSB. Scholes et al. J. Molec. Biol. <u>2</u> 379.
- 30% of directly ionizing events cause SSB. Raskasovskiy et al Radiat Res. <u>153</u> 436.
- A base is released at the site of each SSB. *Ward and Kuo* <u>66</u> 485.
- The termini of SSB are 5' phosphates and 3' phosphoglycolates (35%) and 3' phosphates (65%). *Henner et al. J.Biol. Chem.* <u>258</u> 713.
- 70% of SSB are overt breaks, 30% are alkali labile sites. *LaFleur et al. Int. J. Radiat.Biol.* <u>30</u> 223.
- Alkali labile (abasic) sites are not the same as acid induced abasic sites. *LaFleur et al. Int. J. Radiat.Biol.* <u>35</u> 241.
- 1 Gray produces 1000 SSB per cell Elkind and Redpath, In Cancer, A Comprehensive Treatise <u>6</u> 51.
- Most SSBs are repaired with a half-life of 3-4 min.

Ratio of base damages to SSB

Milligan et al. Radiat. Res. 146 436.

Supercoiled plasmid DNA used as target.

SSB measured as relaxed open circle

DSB measured as linear

Base damage measured as increase in strand breaks after treatment with base damage specific enzymes.

Base damage produced by OH radicals are ≈ 2.6 times as frequently as SSB.

Direct Ionization of DNA

Initial events (observed at 4°K) *e.g. Debije and Bernhard, J. Phys. Chem.* <u>B 104</u>, 7845.

Electron loss leads to guanine cation radicals and deoxyribose radicals.

Electron gain leads to pyrimidine anion radicals.

Products, (after warming and dissolution):

Bases, same types of alterations as those from 'OH radical attack. *Swarts et al. Radiat. Res.* <u>145</u>, 304.

30% of directly ionizing events cause SSB. Raskasovskiy et al Radiat Res. <u>153</u>, 436.

What have we learned from *in vitro* radiation chemistry?

Structures of altered bases.

Strand break end-groups.

"Direct" and "Indirect" mechanisms cause the same types of damage.

Relative yields of these damages.

Reactions in which oxygen or radiation modifiers might act.

In vivo radiation chemistry



Ionizations in a Typical Cell

Cell Size 4 x 10⁻⁹ cm⁻³, Density 1.15 g/cm³ – Mass 4.64 x 10⁻⁹ g

Water	% 70	picograms 3250	Ionizations/Gray 1,000,000	
Inorganic Ions	1	46		
Small Metabolites	3	139		
Proteins	18	835	260,000	
RNA	1.1	51	16,000	
DNA	0.25	12	3,600	
Phospholipids	3	35	10,800	
Other lipids	2	23	7,200	
Polysaccharides	2	23	7,200	J.F.Ward, 2007

Assessment of Significance of Damage in a protein (as an example)

Protein of Mol. Wt. 100,000

20,000 molecules per cell.

Mass of protein

= $10^5 \times 2 \times 10^4$ Daltons = $[10^5 \times 2 \times 10^4 / 6 \times 10^{23}] \times 10^{12}$ picoograms = **3.3 x 10**⁻³ pgm

Energy deposited therein

- 1 Gray = I Joule per kgm. $\equiv 6.25 \times 10^{15} \text{ eV}$ per gm
- ~ 20 eV are needed to cause an ionization.

1 Gy causes 3.1×10^{14} ionizations per gram

Number of ionizations in 3.3 x 10^{-3} picograms would be 3.1 x 10^{14} x 3.3 x 10^{-3} x 10^{-12}

= 1

So from 1 Gray the damage produced by direct ionization is:

1 altered site in the 20,000 copies of this protein

<u>Assessment of relative contributions of direct</u> <u>Ionization and 'OH Radicals in Cells</u>

Cell targets react with 'OH;

RH + $OH \rightarrow$

(1)

If a compound is added which competes with the target for 'OH, the amount of damage to the target will be reduced:

DMSO + $OH \rightarrow$

(2)

Fraction of 'OHs reacting with RH:

<u>k₁[RH]['OH]</u>. k₁[RH]['OH] + k₂[DMSO]['OH]

.I F Ward 2007

Effect of scavengers on Cell Survival

J.D. Chapman et al. Radiat. Res. 56; 291-306.



FIG. 1. The effect of various concentrations of DMSO and cysteamine on the radiation inactivation rate of air-saturated and hypoxic Chinese hamster cells.

Effect of 'OH scavenger on chromosome aberration yield



Littlefield L.G. et al. Int. J. Radiat. Biol. 53; 875-890.

Figure 2. Dose-response relationships for dicentric induction in lymphocytes exposed to X-radiation in absence or presence of DMSO. Dicentric cell⁻¹ data fitted to linear-quadratic dose-response function.

OH Radical Scavengers Reduce Biologically Significant Damage

1964 Bacteria

Johansen, I. and Howard-Flanders, P., Radiat. Res. 24: 184-189.

1972 Mammalian Cells

Roots, R. and Okada, S., Int. J. Radiat. Biol., 21; 329-342. 1973 Chapman, J.D. et al., Radiat. Res. 56; 291-306.

1979 High LET

Chapman, J.D., Radiat. Environ. Biophys. 16; 29-41.

1984 Transformation

Yang T.C. and Tobias, C.A., Adv. Space Res. 4; 207-218.

1987 Mutations

Corn B.W. et al., Radiat. Res. 109; 100-108.

1988 Chromosome Aberrations

Littlefield, L.G., Int. J. Radiat. Biol. 53; 875-890.

1995 **DNA double strand breaks – α particles** *deLara, C.M. et.al., Radiat. Res. 144; 43-49*

- 2000 Cell killing from ¹²⁵I decays Walicka, M.A. Radiat. Res. 154; 326-30.
- 2001 Genomic Instability Limoli, C.L. et al. 31; 10-19

Conclusions of authors:

65 % of most biological effects are caused by 'OH radicals

1/2 maximum protection is provided by 0.13M DMSO

•OH radicals cause base damage and strand breaks in DNA. However:

Other evidence suggests the unimportance of OH

- Treatment with H₂O₂ does not kill cells (*Ward, Blakely and Joner, Radiat Res.103:383-92*).
- ➤ There are high endogenous levels of oxidized bases (≈2.4 -48 Gy). Collins et al. Arch Biochem Biophys. <u>423</u> 57-65
- α-particles produce a lower yield of OHs than γ-rays, but are more effective in killing cells (Objection raised by T. Alper).



FIG. 1. The effect of various concentrations of DMSO and cysteamine on the radiation inactivation rate of air-saturated and hypoxic Chinese hamster cells.

J.D. Chapman et al. Radiat. Res. 56; 291-306. J.F.Ward, 2007

Life Time of 'OH in mammalian cells

Fraction of •OHs reacting with target =

<u>k₁[RH]['OH]</u> k₁[RH]['OH] + k₂[DMSO]['OH]

When the amount of scavengeable damage is reduced by a factor of 2,

 $k_1[RH][OH] = k_2[DMSO][OH],$

Thus, the half life of the 'OH radical in its reactions with DNA intracellularly can be determined:

 $t_{1/2} = \ln 2 / k_2 [DMSO]$

So that if $[DMSO]_{1/2} = 0.13$ M and $k_2 = 6.5 \times 10^9$ LM⁻¹s⁻¹

 $t_{1/2} = 8.2 \times 10^{-10} s$

Evolution of an electron track

J.E. Turner et al. Radiat. Res. <u>96;</u> 437.



Radiation Chemical Concepts of early distributions of radicals

Entity	Energy	Size	Energy (%)	Events (%)
Spur	<100eV	4nm (diam.)	80	95
Blob	<500eV	7nm (diam.)	20	5
Short tracks	500-5,000eV			
DNA		2mm (diam)		
nucleosome		5.7nm thick		

Energy Deposition events

From Table II of Pimblott and Mozumder J. Phys. Chem. <u>95</u> 7928.

Numbers of •OH per event	% of total events	
1	54.4	
2	24.8	
3	8.0	
4	2.3	
5	1.3	
6	0.4	
		J.F.Ward, 200

Distance 'OH moves before reacting in mammalian cells

The diffusion of a particle in 3 dimensions is 2.45 [Dt]^{0.5} (*ref 1*)

For OH, D, the diffusion constant = $2 \times 10^{-5} \text{ cm}^2 \text{ s}^{-1}$ (ref 2)

Mean distance 'OH moves in mammalian cell before reacting with DNA is:

2.45 [2x10⁻⁵ x 8.2 x 10⁻¹⁰]^{0.5} cm

|= ∼3 nm.

 A. Einstein, Investigations on the Theory of Brownian Movement, ed. R. Fürth, translated by A.D. Cowper (1926, reprinted 1956); Einstein, Collected Papers, vol. 2, 170-82, 206-22
 J.E. Turner Atoms, Radiation, and Radiation Protection, 2nd ed. New York: Wiley-Interscience, 1995. Table 13.2)

Energy Deposition Event on top of DNA





Locally Multiply Damage Sites Combination Lesions Underlying Single Track Event Radiation **Signatures**

Relative LMDS Frequencies in DNA in Solution

J.R. Milligan et al. Int. J. Radiat. Biol. <u>76</u> 1475-1483

Locally Multiply Damaged Site	Percent total LMDS
DSB	20
Oxy-purine complex	33
Oxy-pyrimidine complex	46

Enzymes used to cut at base damaged sites, formamidopyrimidine glycosylase (oxy-purine) and endonuclease III (oxypyrimidine) can be inhibited by neighboring damage. Therefore the yields of base damaged sites will be higher.

LMDS containing Base Damage assessed in human cells

B. Sutherland et al. Proc. Natl. Acad. Sci. USA <u>97</u>, 103-108



PNAS

Relative LMDS Frequencies in Human Cells

B. Sutherland et al. Radiat. Res. <u>157</u>, 611–616

Locally Multiply Damaged Site	% Total
DSB	27.5
Oxy-purine complex	27.8
Oxy-pyrimidine complex	24.7
Abasic site complex	20

Note: Values are approximate since there are cross-sensitivities to the enzymes, and, cutting by the enzymes is inhibited by proximal damage (e.g. Weinfeld M. et al. Radiat Res. <u>156</u> 584-9)

Examples of Multiply damaged sites



Variables of Multiply Damaged Sites

Size - distance over which damage spread

Complexity – numbers of damages per site

Composition — variety of base damages and SSB

Spacing of constituent SSBs: DSB production by alternate method

Limoli and Ward Radiat. Res. <u>134</u> 160-9.

DNA labeled with 5-bromouracil.

DNA loaded with Hoechst dye 33258.

Exposed to UVA light; 360 nm (3.4 eV).

Measure DSBs and cell killing.

Hoechst 33258 binding to DNA

Kakkar et al. J. Biomolec. Struct. & Dynam. 2327

The Hoechst 33258-DNA complex. The drug molecule is shown highlighted in green.



DSB produced by photolysis(Dye, BrdU, UVA)

Limoli and Ward Radiat Res. 138 312.



J.F.Ward, 2007

Conclusions from photolysis experiment

DSBs can be produced by agents other than ionizing radiation.

Not all DSBs are equally effective in killing cells.

DSBs which are more closely opposed are more lethal.

Some ionizing radiation induced DSBs are nonlethal.

Potential Consequences of DSBs



DSB and mutations

HPRT gene in hamsters is 36 kbp in length

648 bp are in exons

Double strand break in an exon can lead to a point mutation.

Relationship between yields of point mutations and DSBs

Yield of point mutations = $8 \text{ per } 10^6 \text{ cells per } 2 \text{ Gy.}$

(T. Morgan et al. Mutat Res. <u>232</u> 171)

Size of exon 648 base pairs

Total DNA in exons in 10^6 cells = 6.48 x 10^8 base pairs

Yield of DSB is 5.8 x 10⁻³ per megabase pairs per Gy (*M. Löbrich et al. P.N.A.S.* <u>92</u> 12050)

2 Gy causes 5.8 x 10^{-3} x 6.48 x 10^{2} DSB in target exons

= <u>8 DSB in target exons</u>

Do all DSBs in exons yield point mutations?

The yields are equal. But DSBs with distant SSBs would be expected to be accurately repaired.

Other damage, i.e., LMDS in which base damage is present, are present in several fold higher in yield.

Repair and rejoining of these kinds of damage could also lead to a point mutations.

Are all DSB the same?

- a. Measurements of DSBs are carried out after stripping away all other cellular material.
 Such material (e.g. nucleosomes) may act to hold the ends of DSBs in register enabling their rejoining.
- Removing DNA from cell may also break hydrogen bonded base pairs in between SSBs on opposite strands.
 The hydrogen bonding could serve to hold the ends together favoring fast rejoining.
- c. DSBs measured by these means are greater than the yields existing in cells.
- d. Yields measured by biochemical methods (P.F.G.E., Elution, Centrifugation, etc.) are 37 per cell, but *in situ* by premature chromosome condensation (PCC) are 4-6 per cell. *Cornforth M. p. 563 in "DNA Damage and Repair" (ed. Nickoloff and Hoekstra) Humana Press.*

DNA and nucleosomes

Friedland, W. et al.Rad. Res 150 170-182



Chromatin fiber with zigzag structure. Blue: phosphate groups; white: sugar group atoms; green, yellow, red, violet; base atoms of adenine, guanine, cytosine and thymine, respectively; turqoise; histones

The attraction between DNA (polyanion) and the multiple positive charges on the histones can serve to keep the two ends of a DSB in register, aiding correct rejoining.

Breaks occurring in the linker region may not be so stabilized and may be more prone to separate from their partner end.

Nucleohistone packaging From: K.E. van Holde, Chromatin, Springer Verlag.



Three types of DSB

1. In the linker region – the ends readily separate.

- 2. Held together by holding the ends in register on histones.
- 3. Held together by hydrogen bonding between complementary bases.
- Biophysical DSB measurement techniques detect all three types.

Within the cell they can have different outcomes.

DSB rejoining



FIG. 4. Total rejoining of radiation-induced DSBs after 80 Gy X irradiation. DSB rejoining in G_1 -phase A_L cells and G_0 GM38 cells (panel A) was determined by employing a standard pulsed-field gel electrophoresis assay (*38*) in which the fraction of DNA released from the well is used as an indicator of the relative number of DSBs present in the sample.

Fouladi et al. Radiat. Res. 150, 619-26 (1998).

Effect of dose on correct rejoining.

M. Kühne et al. Radiat. Protec. Dosim. 99, 129–132



Filled circles: Frequencies for correct rejoining as a function of X-ray dose per fraction (total dose 80Gy). 24 h incubation between doses. Open circles: Total rejoining. Closed square: correct rejoining after low dose rate γ -radiation (80Gyover 14 days – 0.24 Gy/h).

What have we learned from in vivo chemistry

- a. OH Radicals have short lives and travel short distances.
- b. OH Radicals react close to where they are produced.
- c. The clusters of ionization from radiation give rise to multiply damaged sites.
- d. There is a variety of MDS.
- e. All DSBs are not alike.

Conclusions about DSBs

- Many techniques for measuring DSBs recruit lesions whose ends do not separate within the cell.
- From the chemistry by which they are produced three classes of DSBs are predicted.
- The DSBs whose ends become available for misrejoining may be only 20% of the measured yields (5-6 cell⁻¹Gy⁻¹).
- DSBs whose ends do not separate may still be detrimental; as a source of point mutations.
- LMDS containing base damage do not give rise to DSBs but can be a source of point mutations.

Damage produced by 1 Gray

1,000 single strand breaks
3,000 damaged bases
37 double strand breaks (measured by harsh techniques)
5 actual double strand breaks (mild techniques)
190 multiply damaged sites

In contrast, UV damage

Sun exposure at 600m. produces thymine dimers in yield equivalent to 14 Jm⁻²s⁻¹ of 253.7 nm light

Klocker et al., Eur. J. Biochem <u>142</u>, 313.

This corresponds to a thymine dimer yield of 1.2 x 10⁵ per cell per hr.

Ward, Radiat. Res. 152 104.

Some Numbers

In a cell, a dose rate of 1mGray per year

produces 1 Actual DSB (PCC) every 185 years

- 1 DSB every 25 years
- 1 SSB per year
- 1 base damage every 4 months
- 1 LMDS every 2.5 years

Human body has 10¹⁴ cells, a dose rate of 1mGray per year produces in 1 second

2.5 x 10⁶ Actual DSB per second

1.9 x 10⁷ DSB every second

- 4.8 x 10⁸ SSB
- 1.4 x 10⁹ base damage
- 1.9 x 10⁸ LMDS

[Abkowitz et al. Blood 100, 2665

Number of pluripotent hematopoietic stem cells per human \sim 1.12 x 10⁴ - 2.24 x 10⁴] ? Calculate # of DSBs per dose.

Comparison of radical distribution from alphas and protons with that of electrons



J.E. Turner et al. Radiat. Res. 96, 437

Hamm et al. Radiat. Res. 104, Suppl. 8, s20.

J.F.Ward, 2007

"New" Paradigms of Radiation Action

Apoptosis **Bystander Effect Chromosome Instability Death Inducing Effect Gene Induction** Low Dose Hypersensitivity **Protein Mobilization**

From: The Breakdown of Desoxyribonucleic acid under Deuteron and Electron Bombardment.

C.L. Smith Arch Biochem. Biophys. 46. 12-17

"The assumption made is plausible and perhaps not too improbable

and is possibly true in essence if not in detail. It has not, however, yet

been confirmed by experiment."