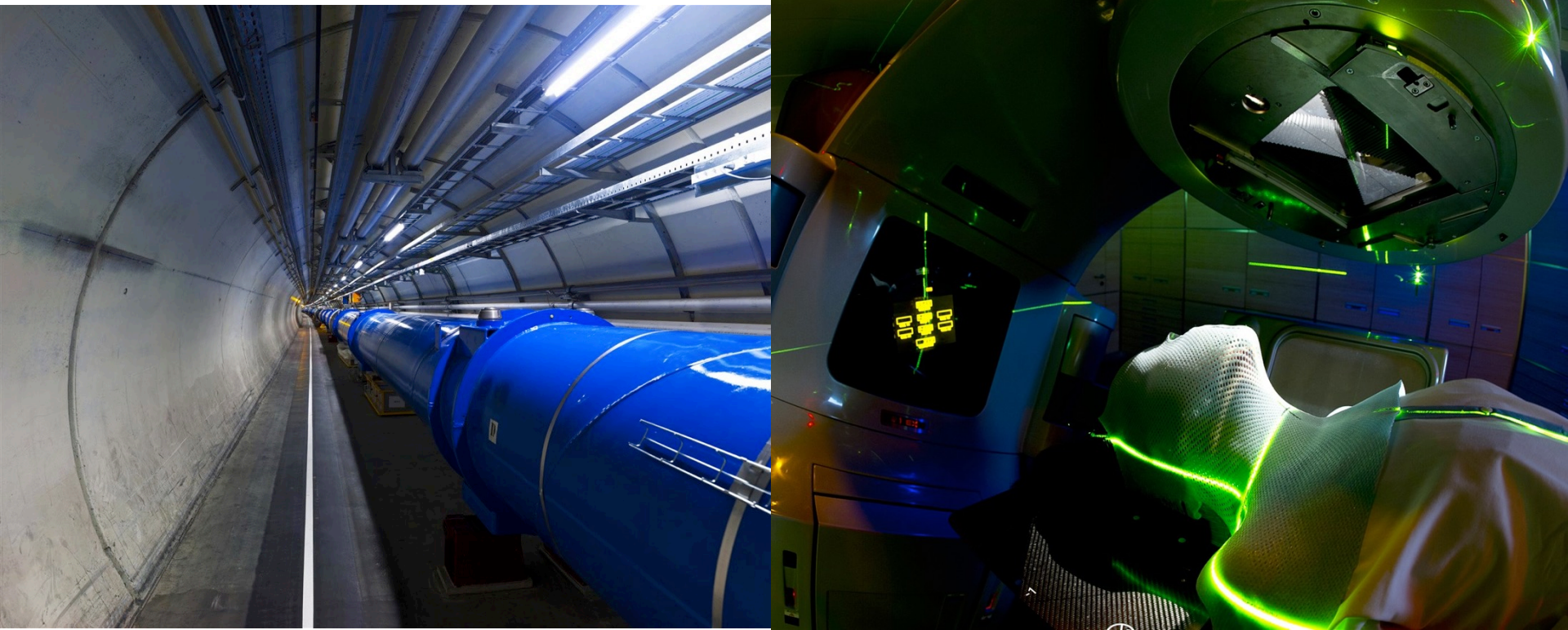


Introduction to Radiobiology



JAI Graduate Accelerator Physics Course, 2022

Manjit Dosanjh
manjit.dosanjh@cern.ch

Radiation Dose

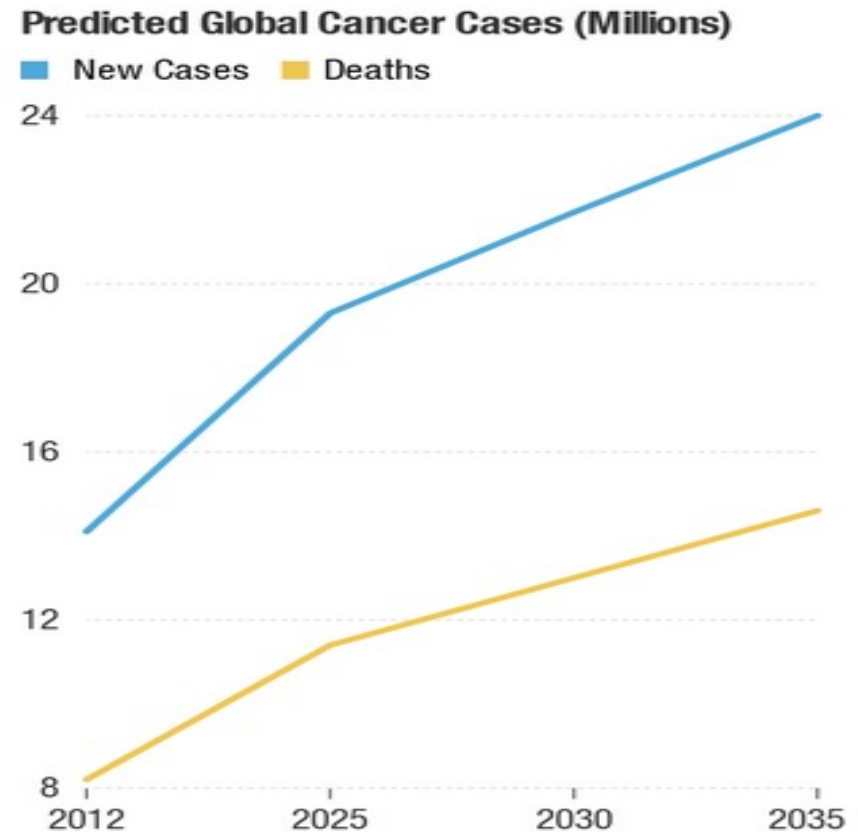
- Radiation effects depend on DOSE = Energy Deposited by Radiation per Unit Target Mass
- **Dose in biology/clinics is measured in Gray (Gy) (=1 joule / kg)**
- ...but different radiations have different effectiveness (Q)
- **Equivalent dose= $Q \times D$ is measured in Sievert (Sv)**
- For x-, γ -rays and electrons: 1 Gy = 1 Sv
- But not always equal, for example: 1 Gy α -particles= 20 Sv (Q=20)
- Average background radiation dose on Earth= 3 mSv / year
- CERN **1mSv** for non-professional and 20mSv/year professional exposure
- Occupational limit= 50 mSv/year
- Lethal dose= 4.5 Sv
- Radiotherapy= 60-70 Gy (to the tumour)
- Average background radiation dose in space = 1 mSv/day

Cancer is growing global challenge

- Globally **19.3** million new cases per year diagnosed and **910** million deaths in **2020**
- This will increase to **27.5** million new cases per year and **16.3** million deaths by **2040**

Radiation therapy is a key tool for cancer treatment

- 3 “Cs” of RT : cure, conservative, cheap
- About 50% patients are treated with RT
- No substitute for RT
- Number of patients is increasing

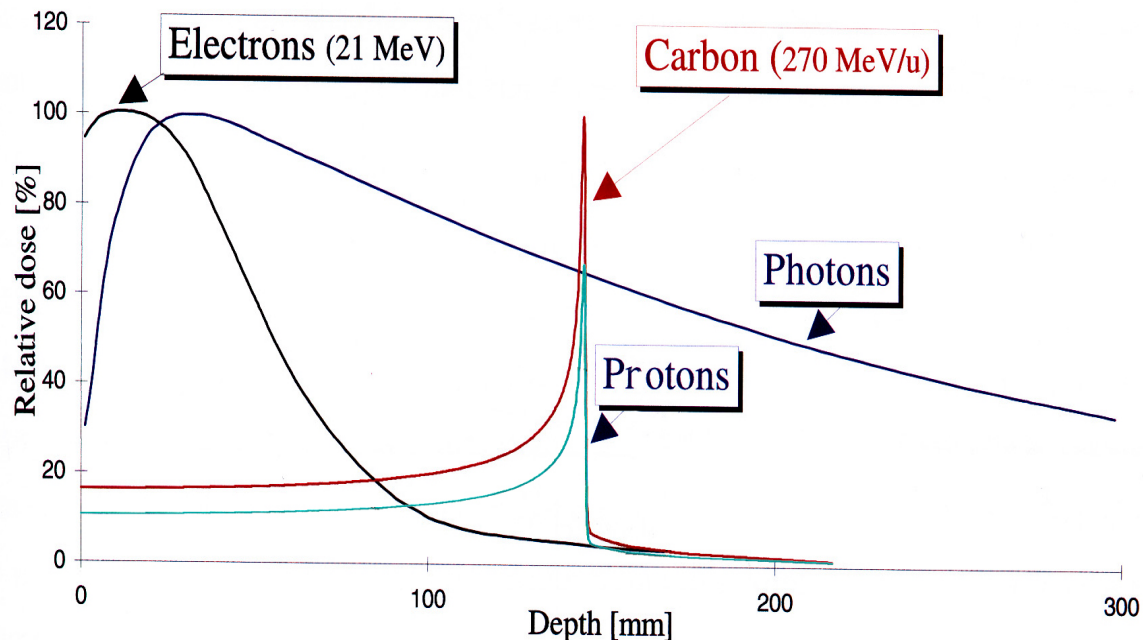


Aims of Radiotherapy:

- Irradiate tumour with sufficient dose to **stop cancer growth**
- **Avoid complications** and **minimise** damage to surrounding tissue

Current radiotherapy methods:

- MV photons
- 5 - 25 MeV electrons
- 50 - 300 MeV/u hadrons



Questions

- What is radiobiology?
- Why do we need biology for radiotherapy?
- What kinds of biology are important for radiotherapy?
- How do you investigate biological effects of particle beams?
- What do the data tell you?
- Do we know everything we need to know?

What is radiobiology?

- Radiobiology is a branch of science which tackles the action of radiation on biological cells, tissues and living organisms.
- When we deliver a known physical dose with a high degree of accuracy to similar cells/tumours - the effect on cells/tissues varies widely
- Radiobiology plays an important role in safe and effective application of radiation in imaging and radiation therapy for cancer treatment as well other applications such as radiation-protection.
- We need to understand the biological factors that influence the sensitivity of cells to deliver better radiation therapy

Radiobiology

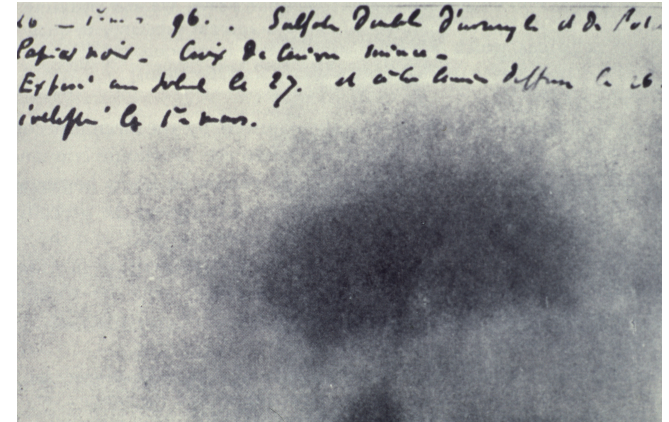
- The response to radiation is different in normal tissues and cancer tissues:
 - at the cellular level
 - at the tissue level
- These differences are due to the underlying biological properties of different tissues and cancers
 - What is the relevance of these differences?
 - How do they have to be taken into consideration?
 - How can we exploit them?
- Different types of ionizing radiation have different and differing effects on cells, tissues, tumours

.....of radiation biology

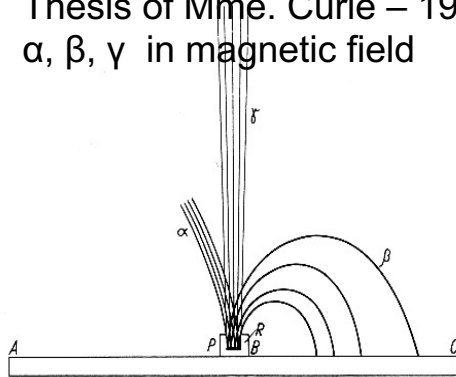


Henri Becquerel (1852-1908)

1896:
Discovery of natural
radioactivity

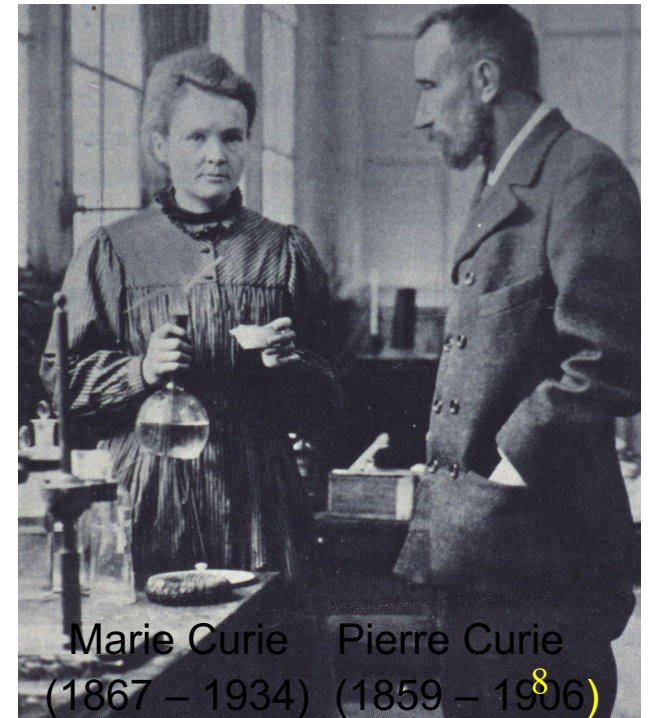


Thesis of Mme. Curie – 1904
 α , β , γ in magnetic field



**1898: Discovery of
radium**

**used immediately
for “Brachytherapy”**



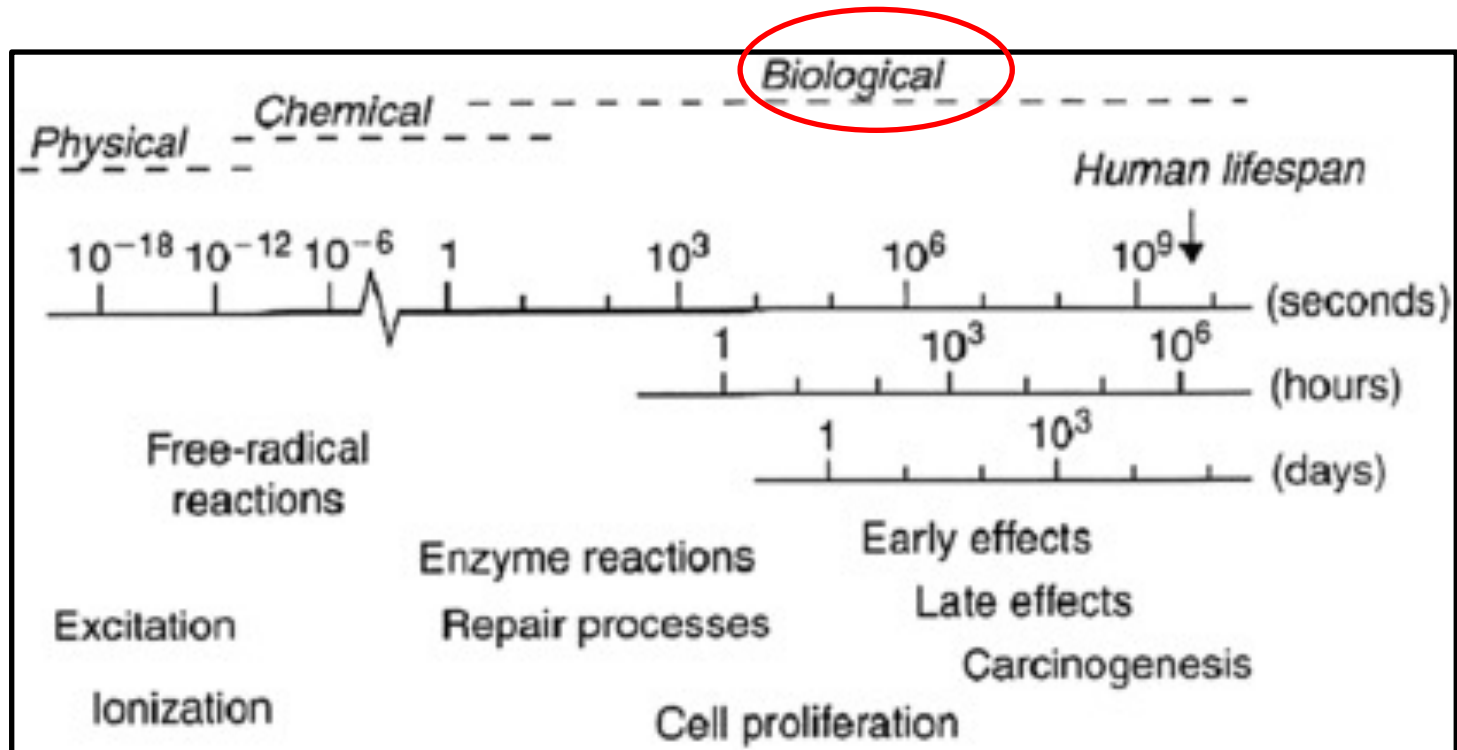
Marie Curie Pierre Curie
(1867 – 1934) (1859 – 1906)⁸

First Radiobiological Experiment

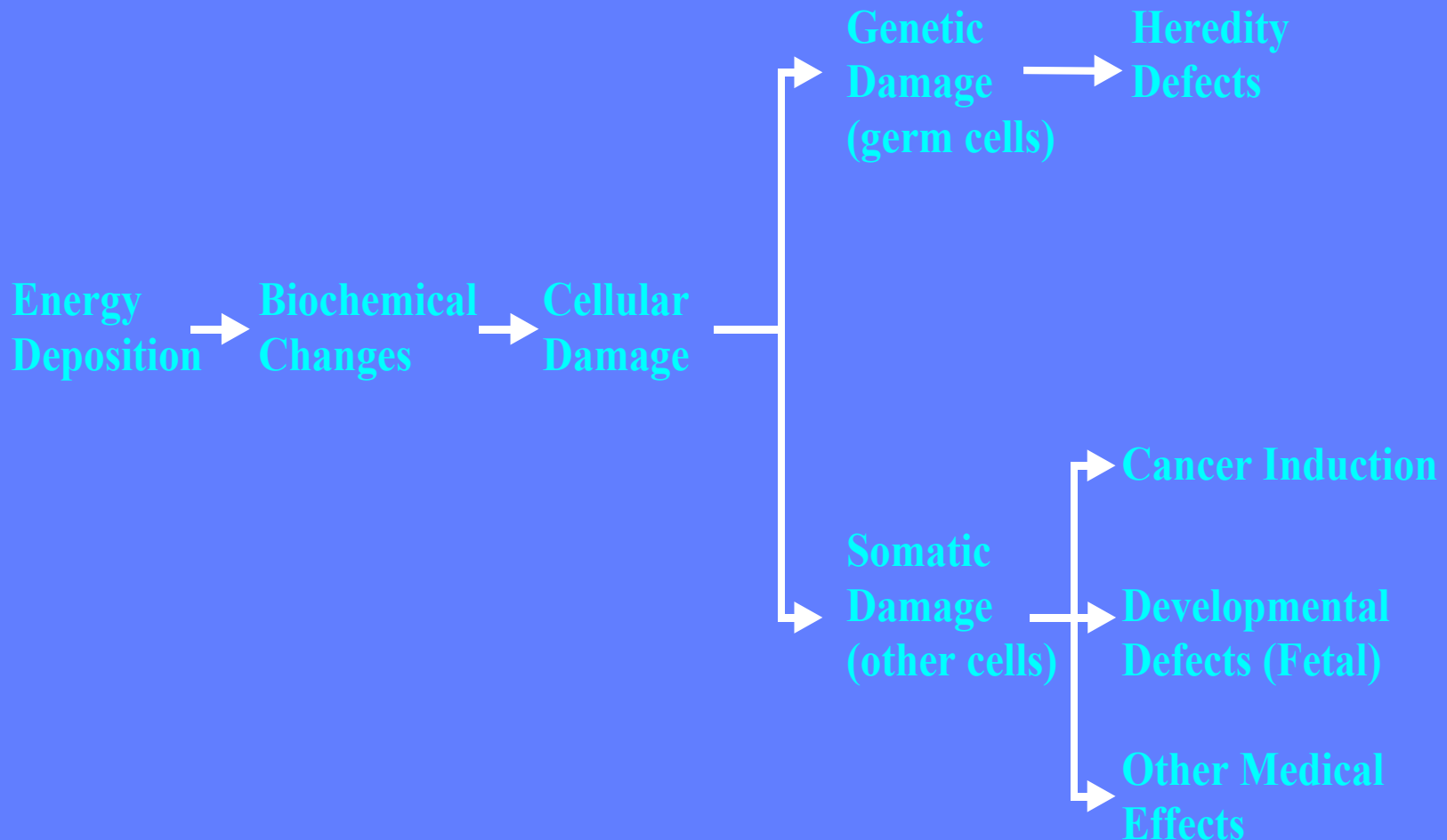


The first radiobiology experiment. Pierre Curie using a radium tube to produce radiation ulcer on his arm. Hall fig. 1-2

Time-scale effects in radiation biology



Major Events Which Follow Energy Absorption From Ionizing Radiation



Effects of radiation damage on cells

Radiation causes damage to all cellular molecules, but DNA damage is most critical as most cellular and molecular components can be replaced

- Cells stops growing - cell cycle arrest:

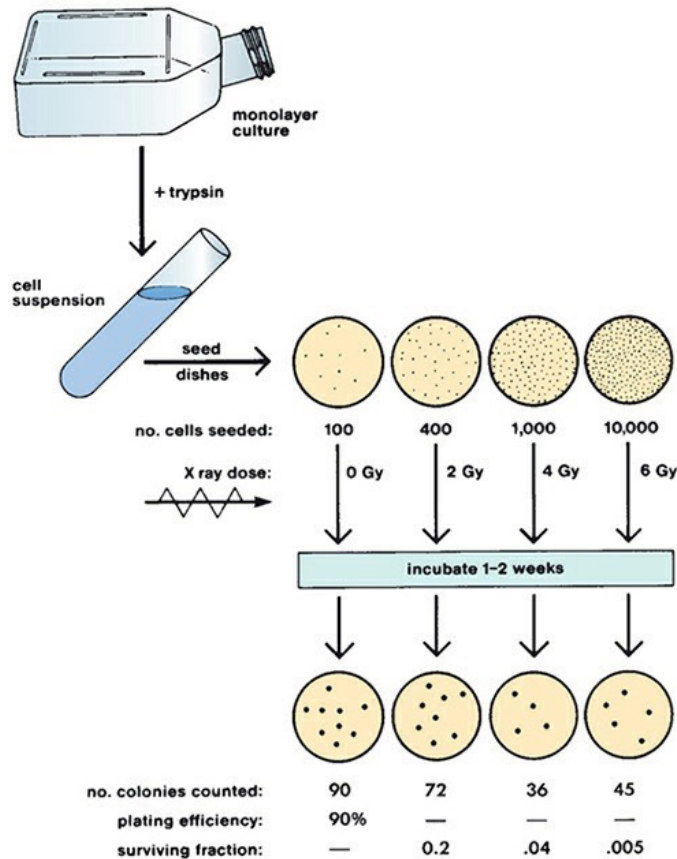
DNA damage or blockage of replication by triggering "checkpoint" responses, which delay cell cycle progression, promote and give time to repair and protect genome integrity.

- DNA repair:

DNA repair processes either remove the lesion(s)/damage or misrepairs the induced damage such that all surviving progeny of an irradiated cell carry the burden of radiation exposure, e.g. gene mutations

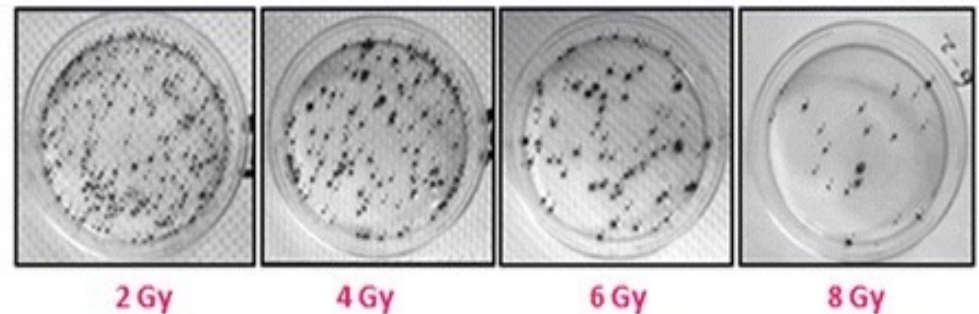
- Cell death

Cell culture techniques and cell survival curves



Puck and Marcus (1956) promoted the study of radiation on individual cells... using cell culture and cell survival

Radiosensitivity measured by Clonogenic Assay in 14 days

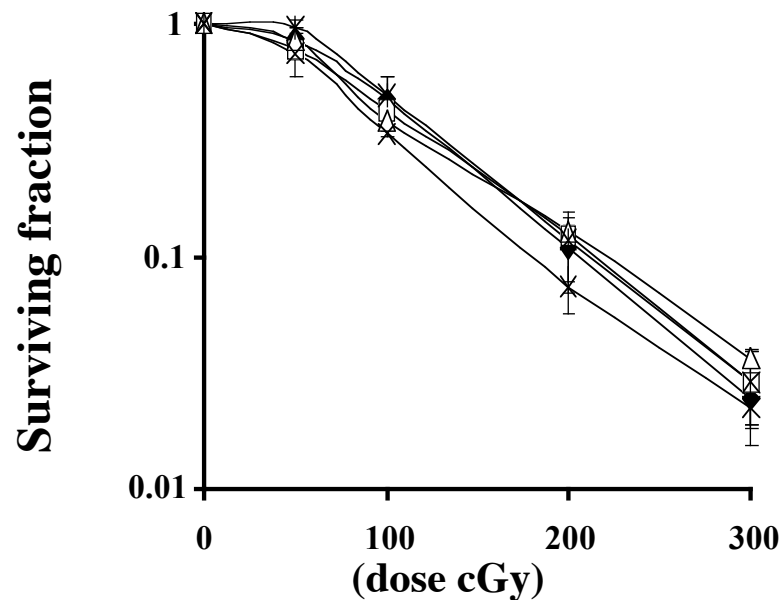


$S/S_0 = \text{colonies produced} / \text{cells plated} * \text{PE}$
 PE = plating efficiency (correction factor derived from control samples)

Cell survival curves

- Describe the relationship between the radiation dose and the proportion of cells that survive reproductive integrity
- Cell survival as a function of radiation dose is graphically represented by plotting the surviving fraction on a logarithmic scale on the Y axis against dose on a linear scale on the X-axis.
- Cell surviving fractions are determined with in vitro or in vivo techniques.

1956: The first in vitro radiation survival curves on mammalian cells carried out by Puck & Marcus



Cell killing by different radiation types

Cell survival after exposure can be expressed in terms of a logarithmic curve of survival versus dose.

For X- or γ -rays (said to be sparsely ionizing), the dose-response curve has an initial slope, followed by a shoulder; at higher dose, the curve tends to become straight again.

For α -particles or low energy neutrons (said to be densely ionizing), the dose-response curve is a straight line from the origin (i.e., survival is an exponential function of dose).

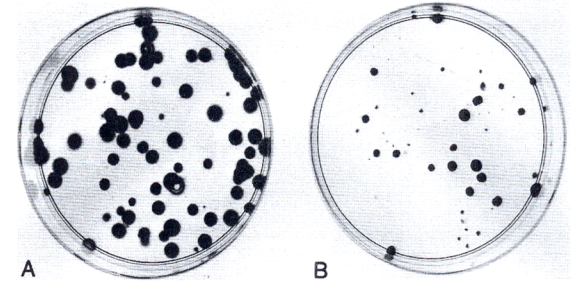
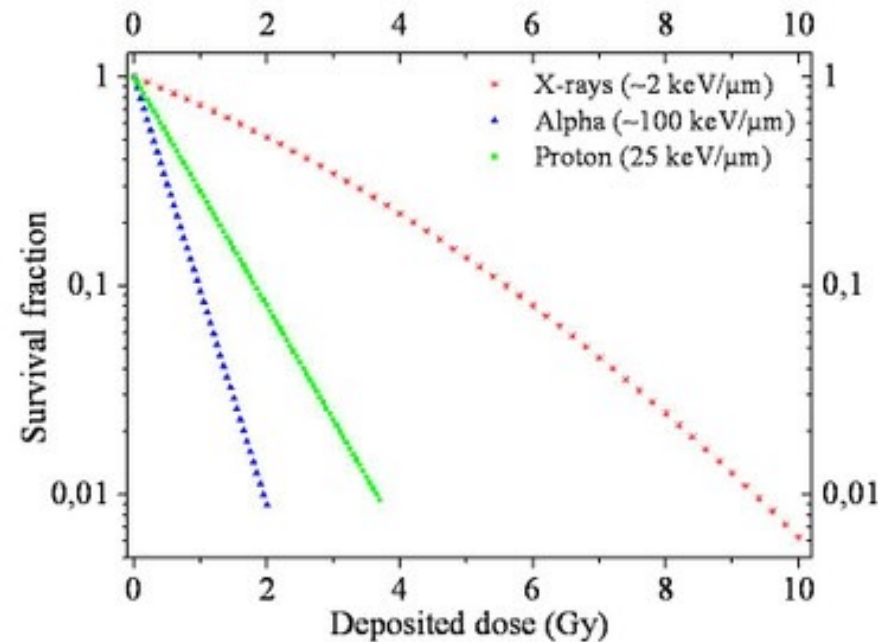
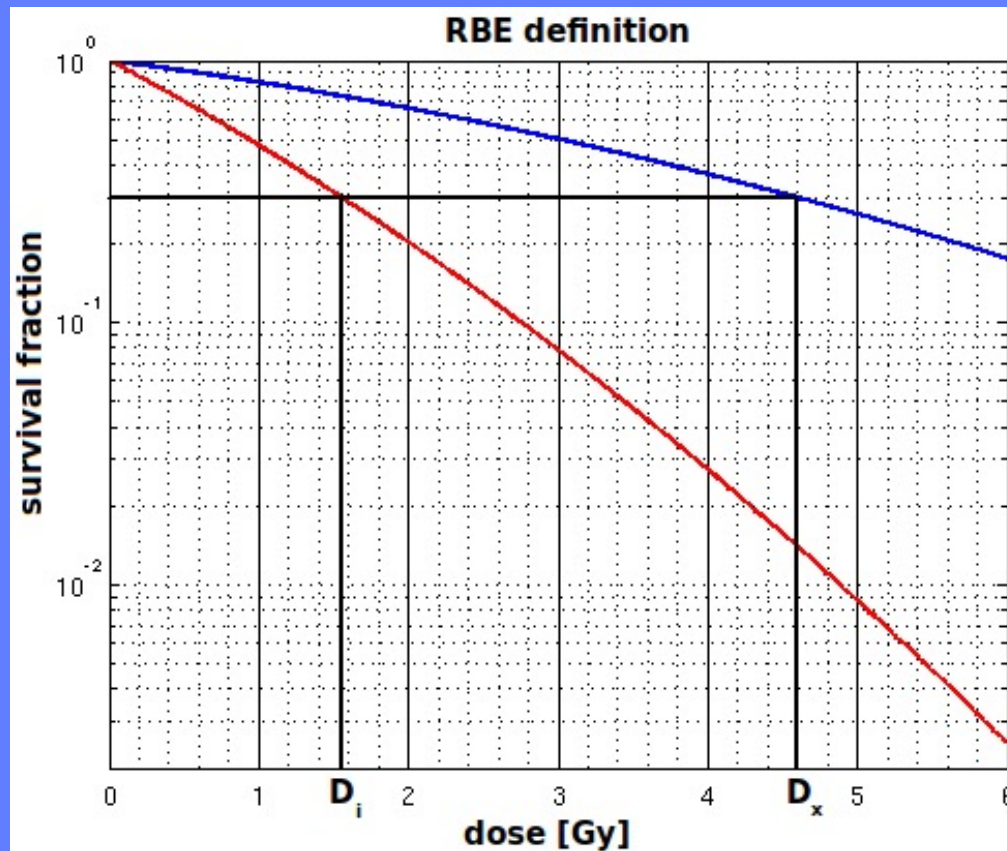


Figure 3.1. Colonies obtained with Chinese hamster cells cultured in vitro. **A:** In this unirradiated control dish 100 cells were seeded and allowed to grow for 7 days before being stained. There are 70 colonies; therefore the plating efficiency is 70/100, or 70%. **B:** Two thousand cells were seeded and then exposed to 800 rad (8 Gy) of x-rays. There are 32 colonies on the dish. Thus:
Surviving fraction = Colonies counted [colonies seeded \times (PE/100)]
= 32/2000 \times .7
= 0.023



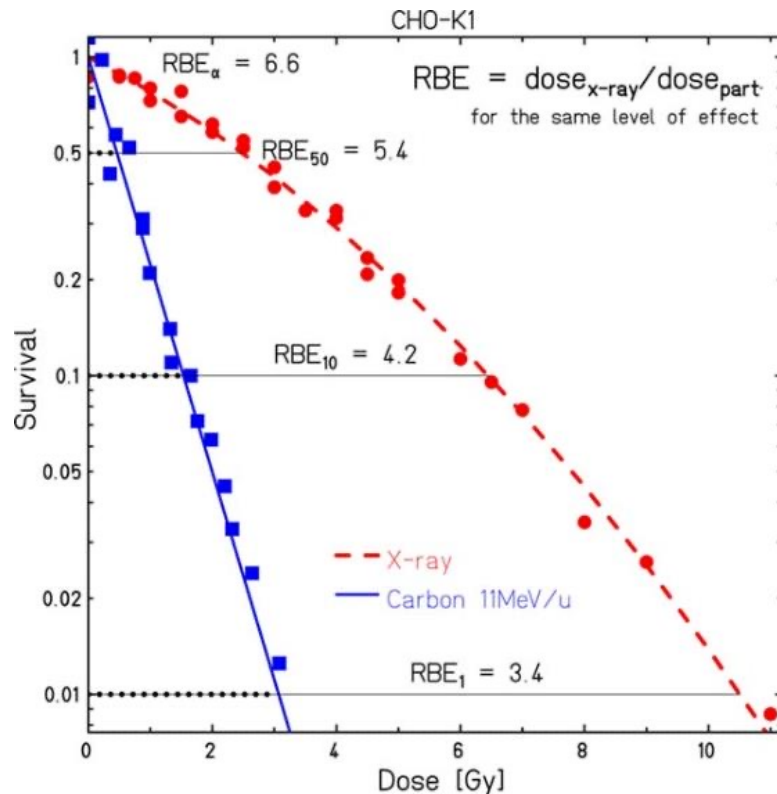
Cellular Survival Curves and Relative Biological Effectiveness

(reference vs test radiation)



$$\frac{D_x}{D_i} = \text{RBE}$$

RBE and how does it vary

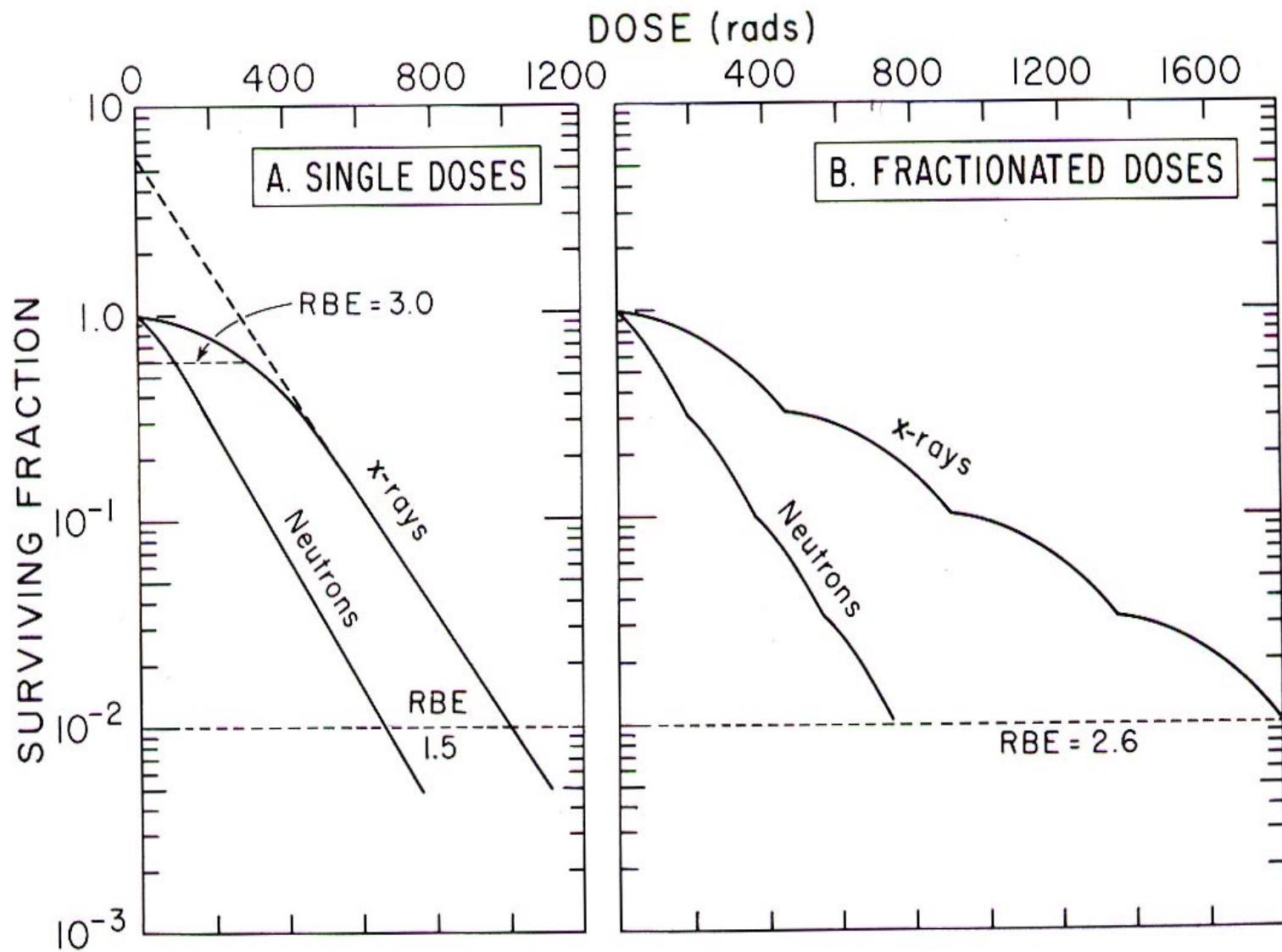


RBE critically depends on both physical and biological parameters:

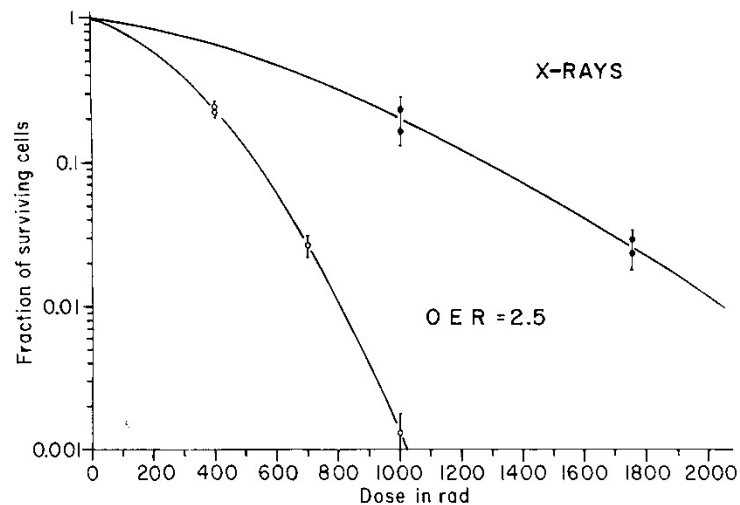
- Dose and Dose Rate
- Varies with types of radiation
- Varies with types of cell/tissue and radiosensitivity
- Varies with the biological effect under investigation
- Varies with dose rate and fractionation
- An increase in RBE in itself does not offer therapeutic advantage unless there is differential effect between normal and tumour tissues
- OER (oxygen enrichment ratio) effects RBE
- Effected by presence of other chemicals present

Fractionation

- Increased cell survival when a dose is split into two or more fractions separated by a time interval
- There is a point at which an increase in the number of fractions will no longer increase survival—plateau in the response



Oxygen for high and low LET radiations



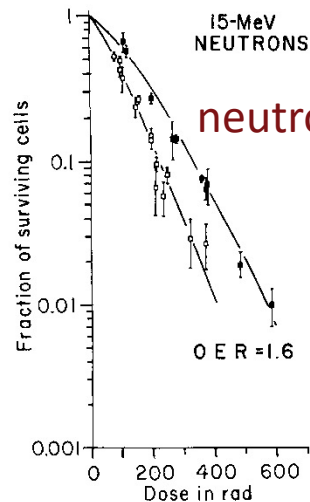
X-rays

OER varies with LET:

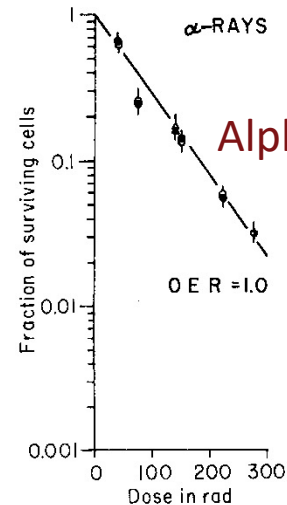
X-rays=2.5

Neutrons=1.6

Alpha particles=1.0



neutrons



Alpha particles

Figure 6.2. The oxygen enhancement ratio (OER) for various types of radiation. The OER for α particles is unity. X-rays exhibit a larger OER of 2.5. Neutrons (15-MeV $d^+ \rightarrow T$) are between these extremes, with an OER of 1.6. (Adapted from Barendsen GW, Koot CJ, van Kersen GR, Bewley DK, Field SW, Parnell CJ: Int J Radiat Biol 10:317, 1966; and Broerse JJ, Barendsen GW, van Kersen GR: Int J Radiat Biol 13:559, 1967, with permission.)

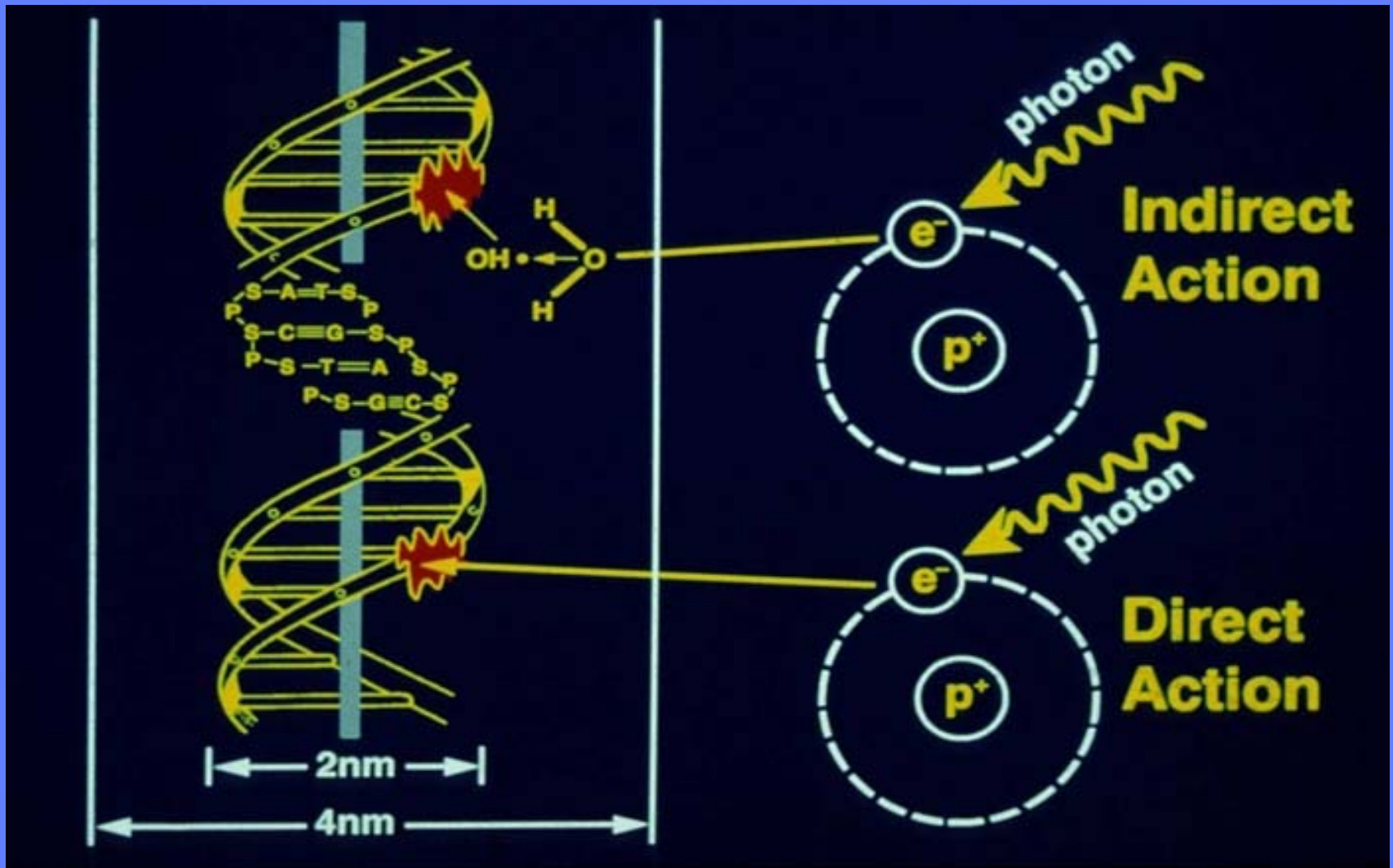
Re-oxygenation in Radiotherapy

- Hypoxia confers resistance to X-rays/gamma rays – also to chemotherapeutic drugs....tumours are normally hypoxic
- Human tumours that do not respond to radiotherapy may not re-oxygenate -radioresistance
- Optimal fractionation regimen depends on reoxygenation

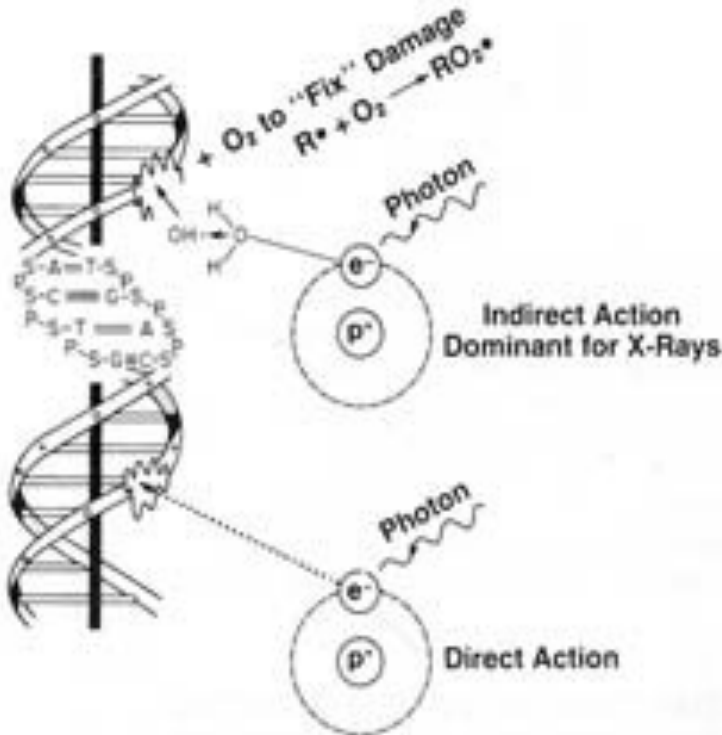
Is the measurement of cellular survival in vitro adequate?

- Which cell types are relevant?
- What about DNA damage and repair?
- What about molecular endpoints?
- What about acute effects in vivo?
- Are there late effects different from acute response?
- What about risk of secondary cancer?

Direct and Indirect Interactions

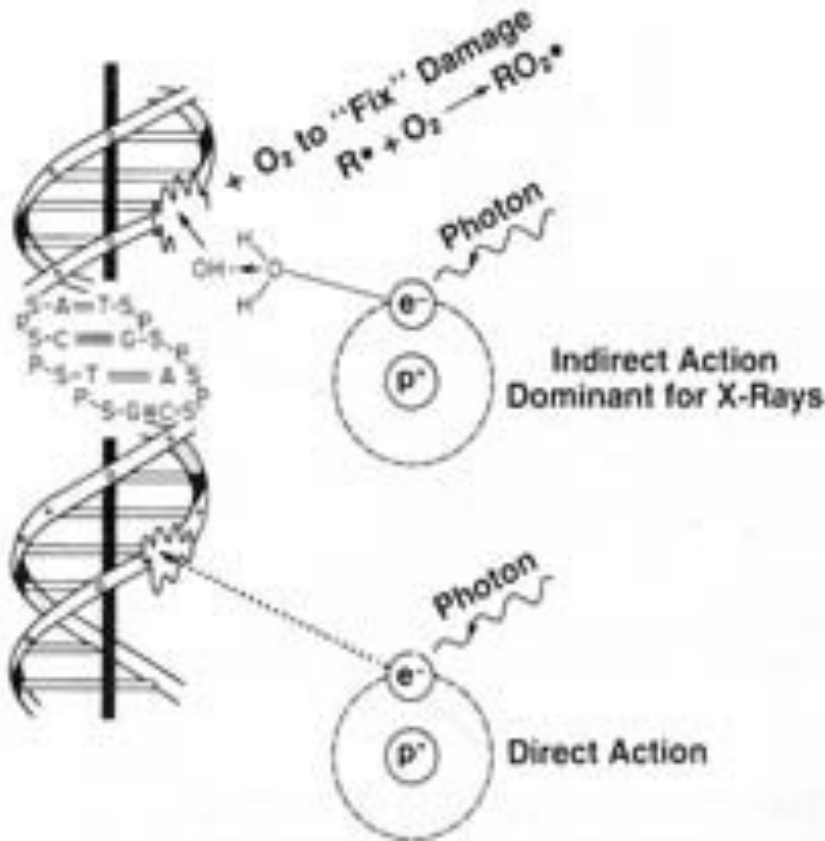


Indirect action in cell damage by radiation



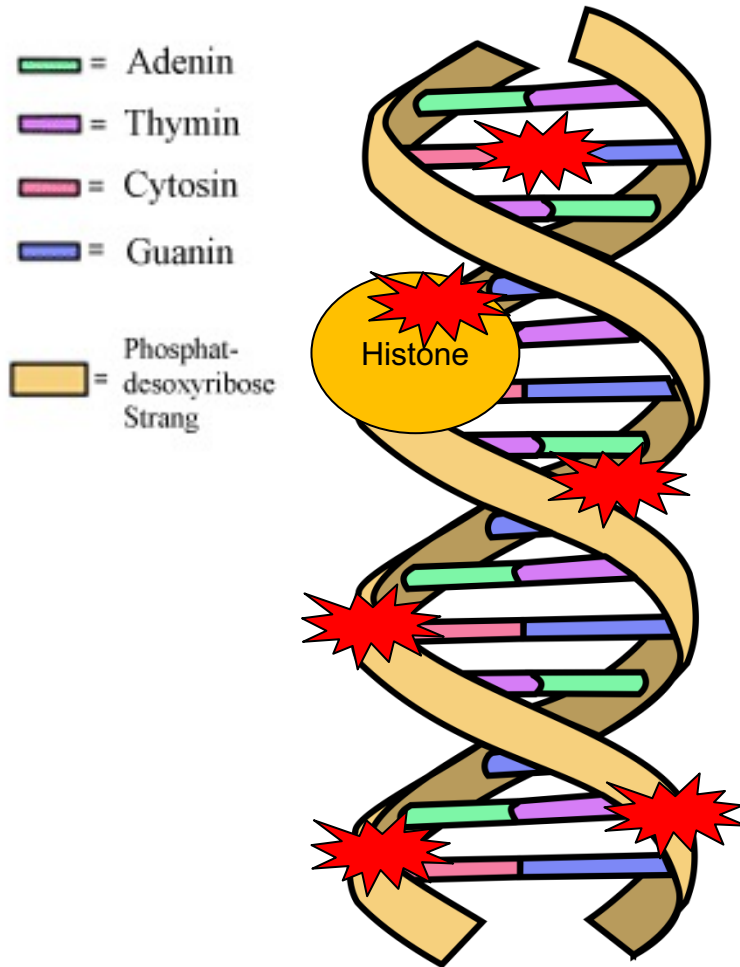
- In indirect action the radiation interacts with other molecules and atoms (mainly water, since about 80% of a cell is composed of water) within the cell to produce free radicals, which can, through diffusion in the cell, damage the critical target within the cell.
- In interactions of radiation with water, short lived yet extremely reactive free radicals such as H₂O⁺ (water ion) and OH• (hydroxyl radical) are produced. The free radicals in turn can cause damage to the target within the cell.
- The free radicals that break the chemical bonds and produce chemical changes that lead to biological damage are highly reactive molecules because they have an unpaired valence electron.
- About 2/3 of the biological damage by low LET radiations is due to indirect action.

Direct action in cell damage by radiation



- In direct action the radiation interacts directly with the critical target (DNA) in the cell. The atoms of the target itself may be ionized or excited through Coulomb interactions, leading to the chain of physical and chemical events that eventually produce the biological damage.
- Direct action is the dominant process in the interaction of high LET particles with biological material

DNA damage – 1 Gy



base damage

4000-5000 incidents

DNA-protein crosslinks

150 crosslinks

sugar changes

800 – 1500 changes

single strand breaks

1000 SSB

double strand breaks

30- 60 DSB

If cells are irradiated with x-rays, many breaks of a single strand occur. In intact DNA however single strand breaks are of little biological consequence because they are repaired readily using the opposite strand as template.

If the repair is incorrect (misrepair), it may result in a mutation.

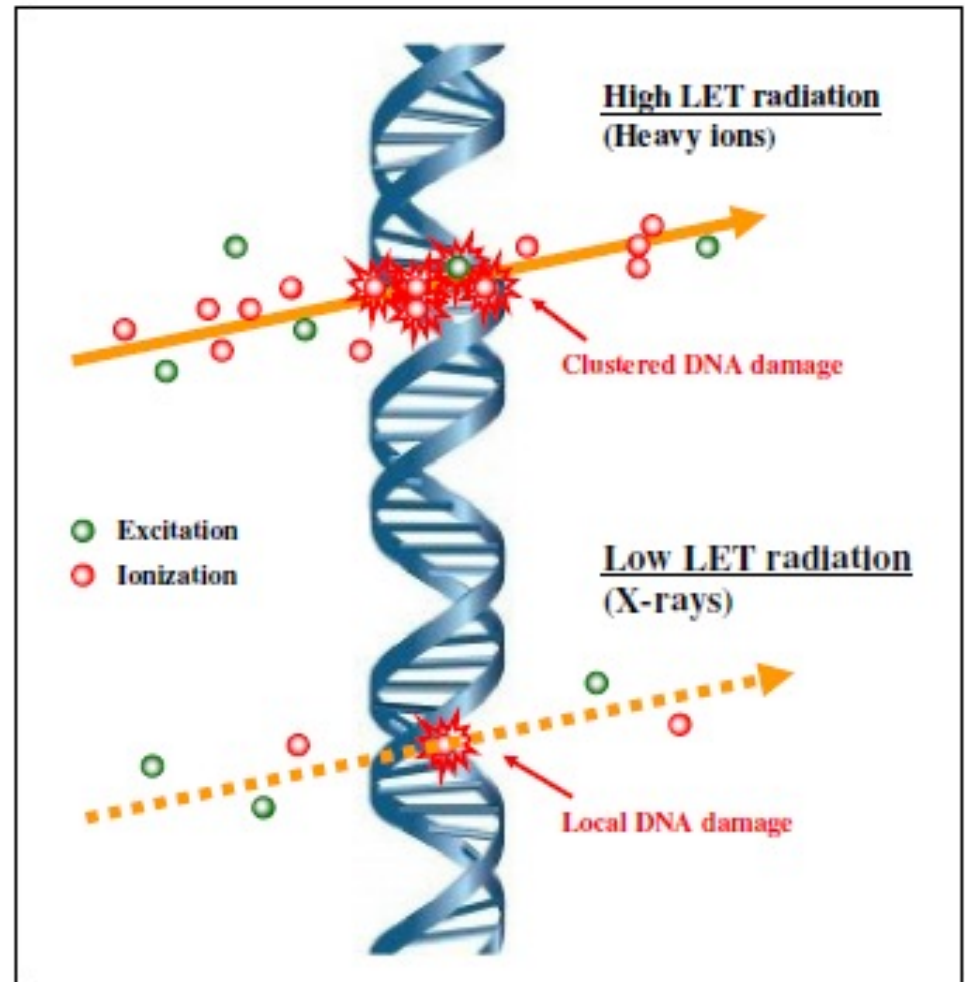
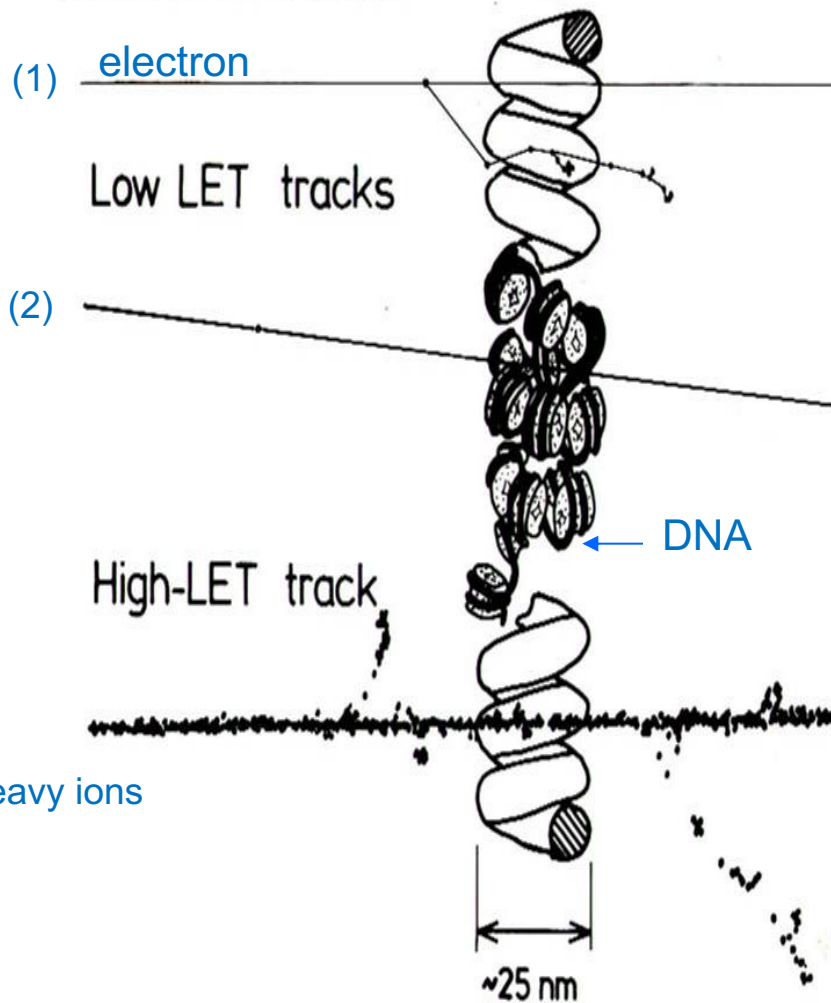
If both strands of the DNA are broken, and the breaks are well separated, repair again occurs readily because the two breaks are handled separately.

By contrast, if the breaks in the two strands are opposite one another, or separated by only a few base pairs, this may lead to a double strand break (DSB).

A DSB is believed to be the most important lesion produced in chromosomes by radiation.

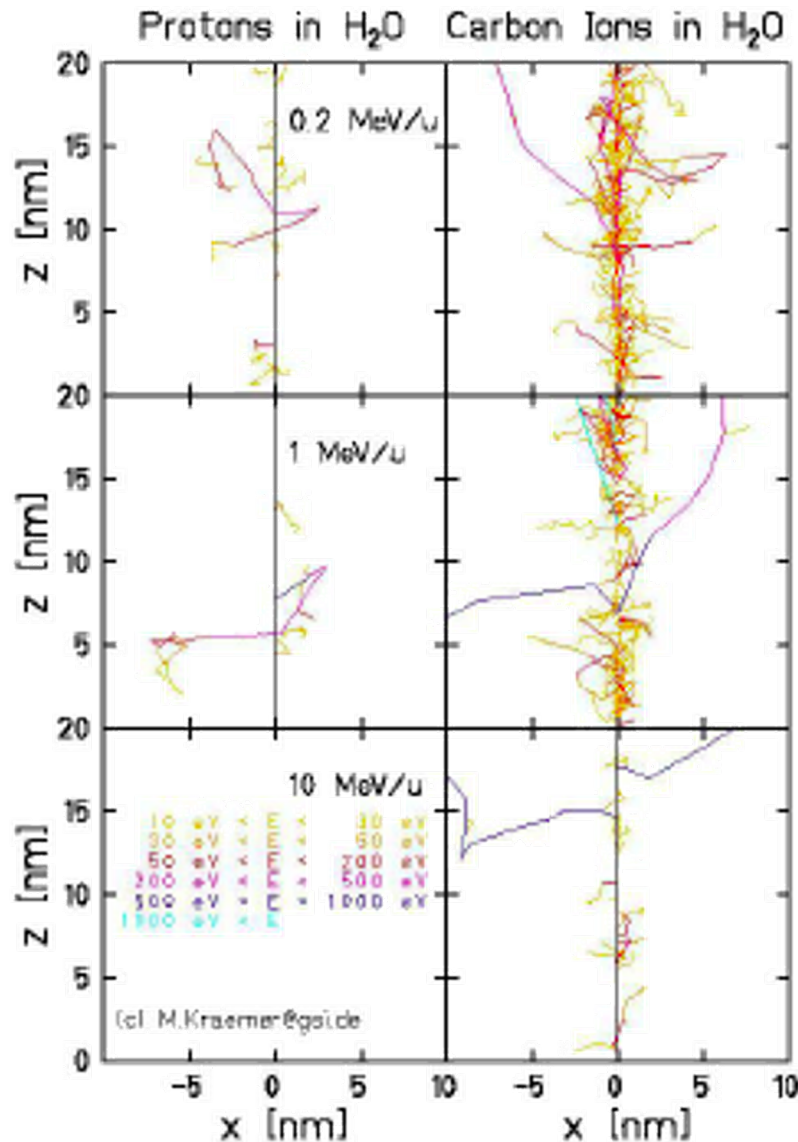
All radiation tracks are highly structured on the scale of DNA

Tracks in chromatin fibre



Delta-ray electron

Track Structures of Proton vs. Carbon Ions



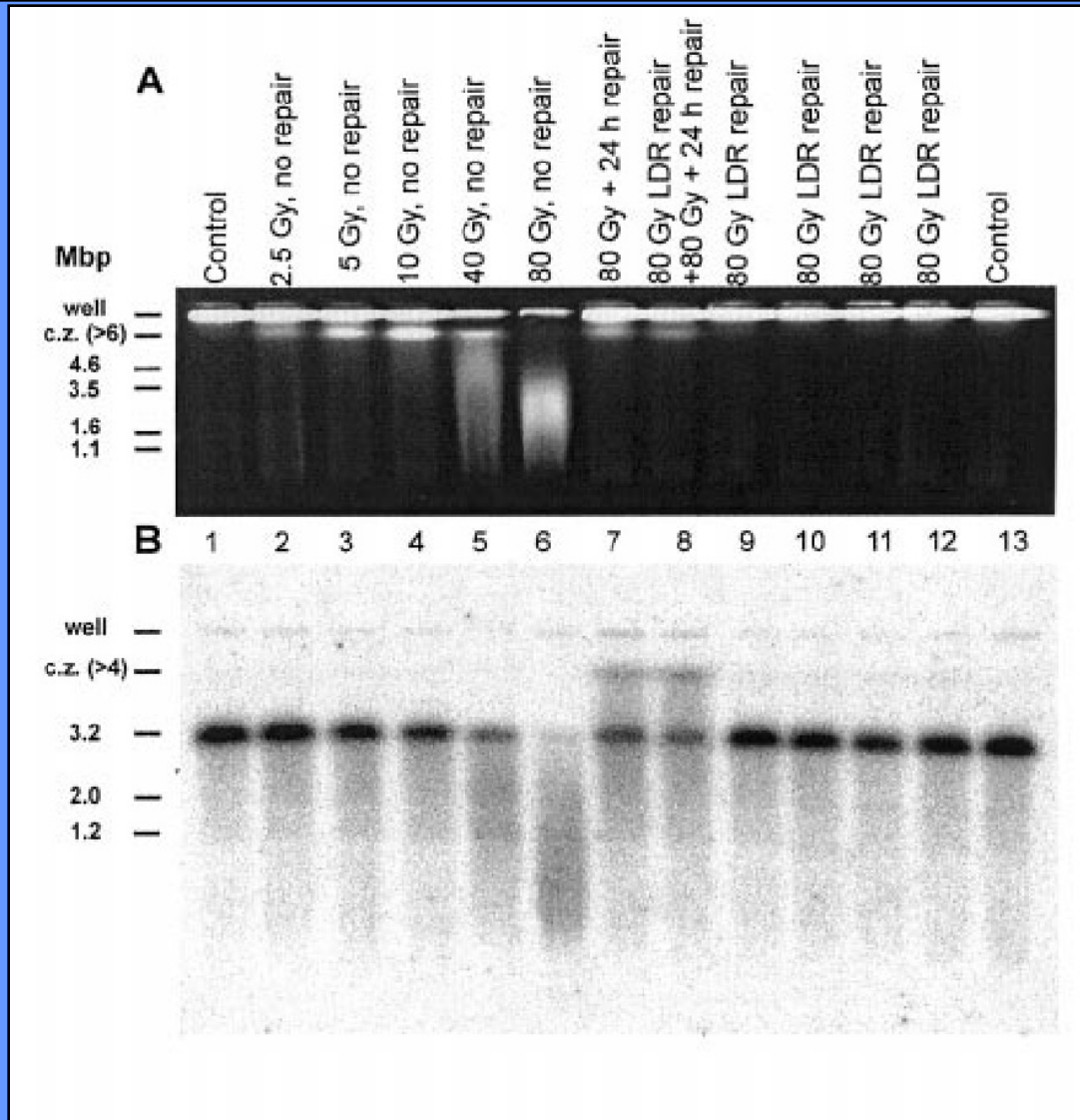
Linear Energy Transfer (LET) stands for the radiation energy deposited per unit length in tissue.

- X-rays and proton beams are low-LET radiations
- Heavy ion beams are high-LET radiation in Bragg peaks

Biological advantages:

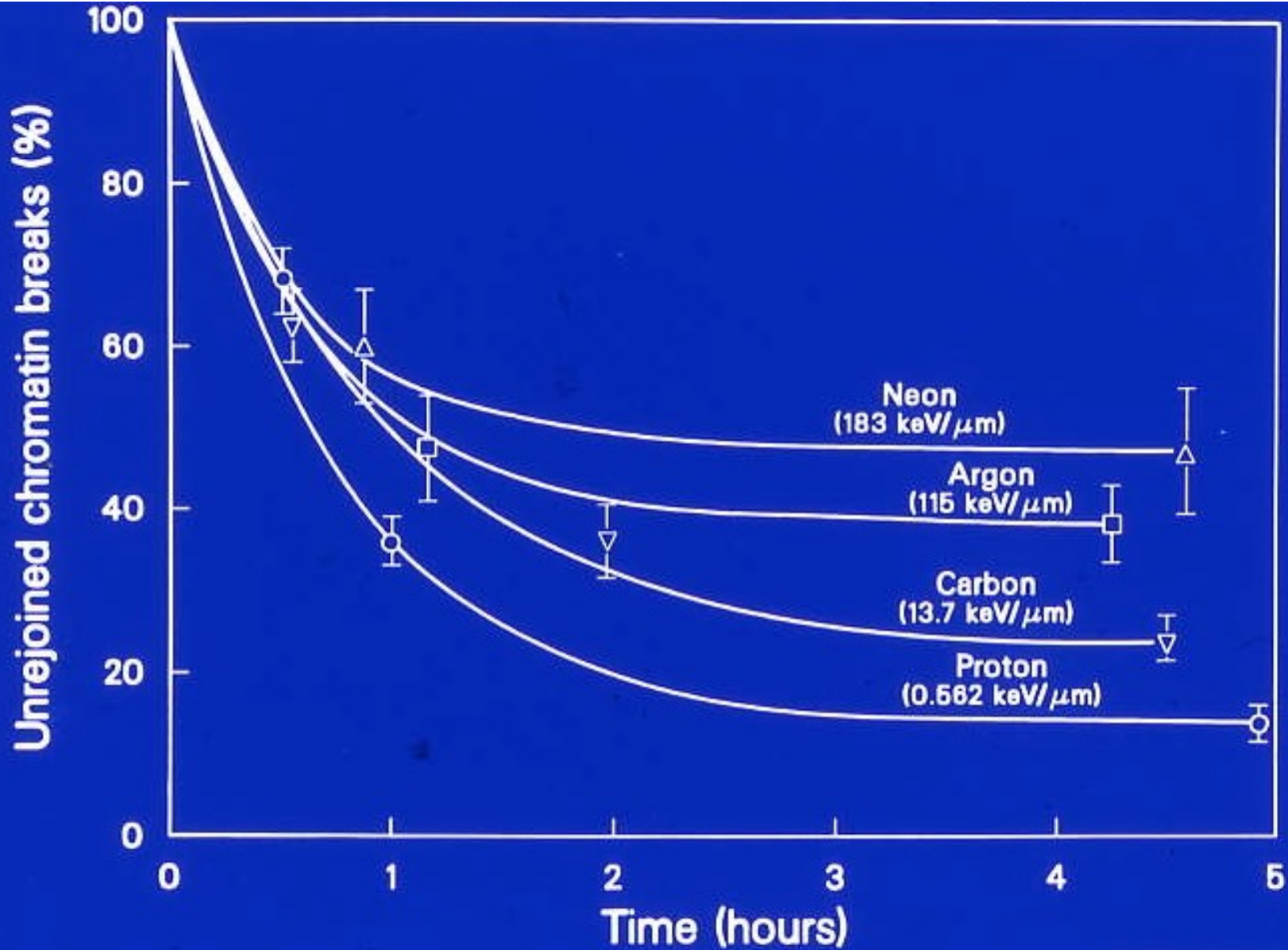
- High LET to provide significant differences in DNA damages
- Suppression of radiation repair
- Yet avoids some complications with higher-Z ions

DSB induction and rejoining in normal human fibroblasts following X-irradiation



*Rothkamm K. et al.,
Cancer Res., 2001.*

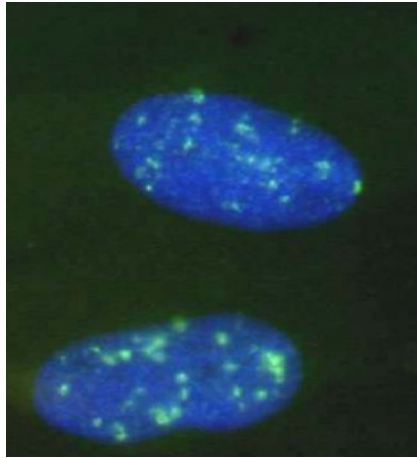
Chromatin Rejoining From Heavier Ion Damage is Slower



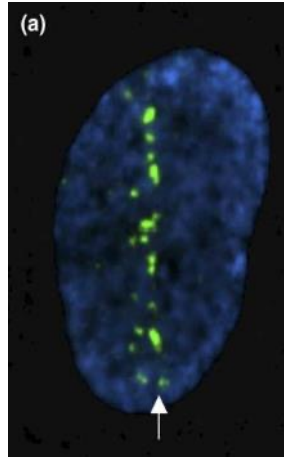
XCG 884-6604 A

Dose, LET and RBE

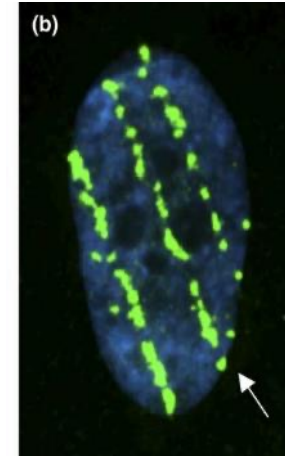
- Cellular response is determined by the level and **quality of DNA damage**, which reflects the energy deposition pattern.



X-rays

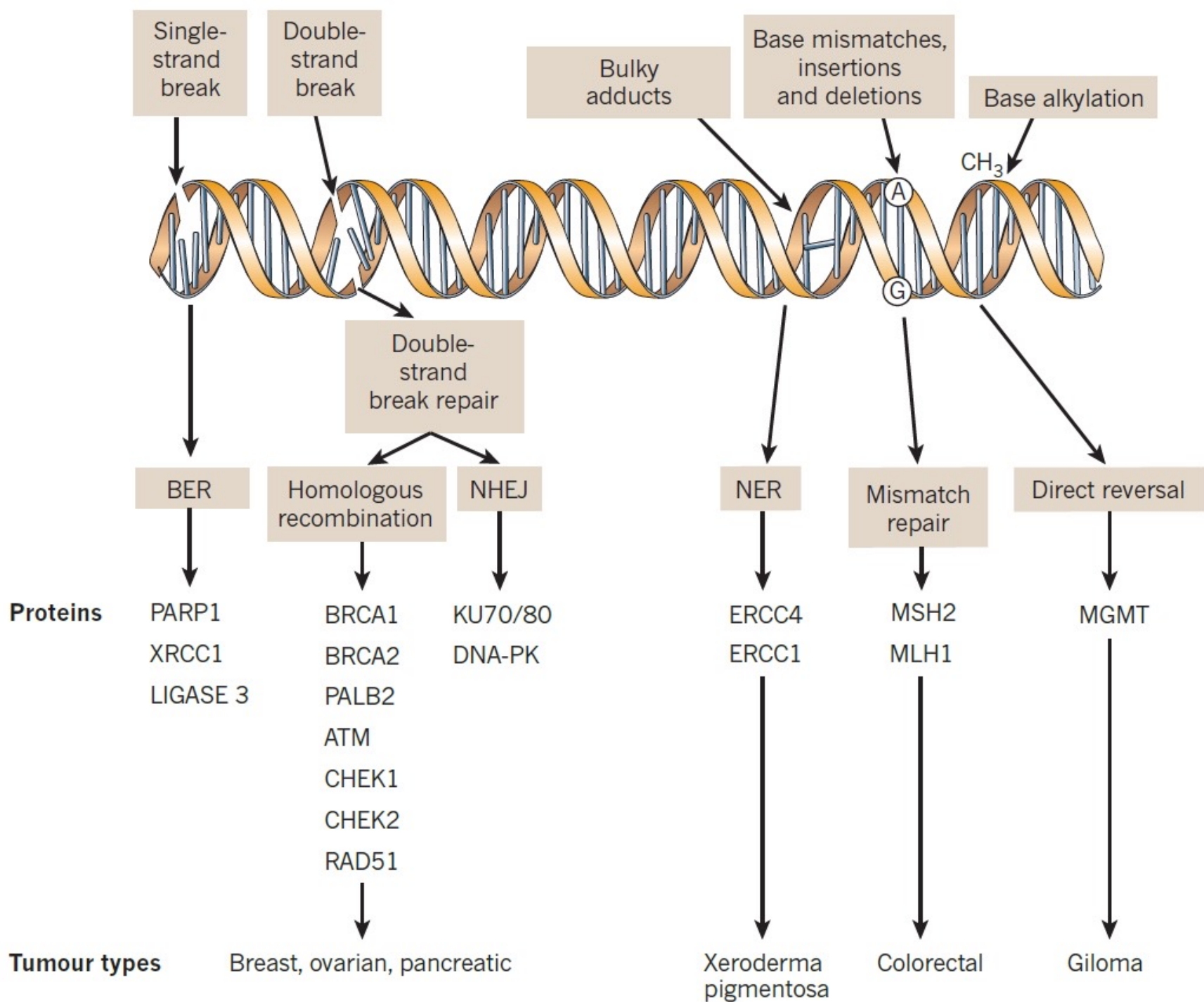


54 keV/μm Si ions



174 keV/μm Fe ions

- Severity of DNA damage** depends on lesion proximity and repairability, hence **it is not a constant value** but depends on physical (particle type, LET, dose) and biological (cell type, oxygenation status, repair capacity) parameters.
- RBE varies with the particle energy and the change of the beam composition (SOBP and nuclear fragmentations): its distribution is **not homogenous** across a treatment field.



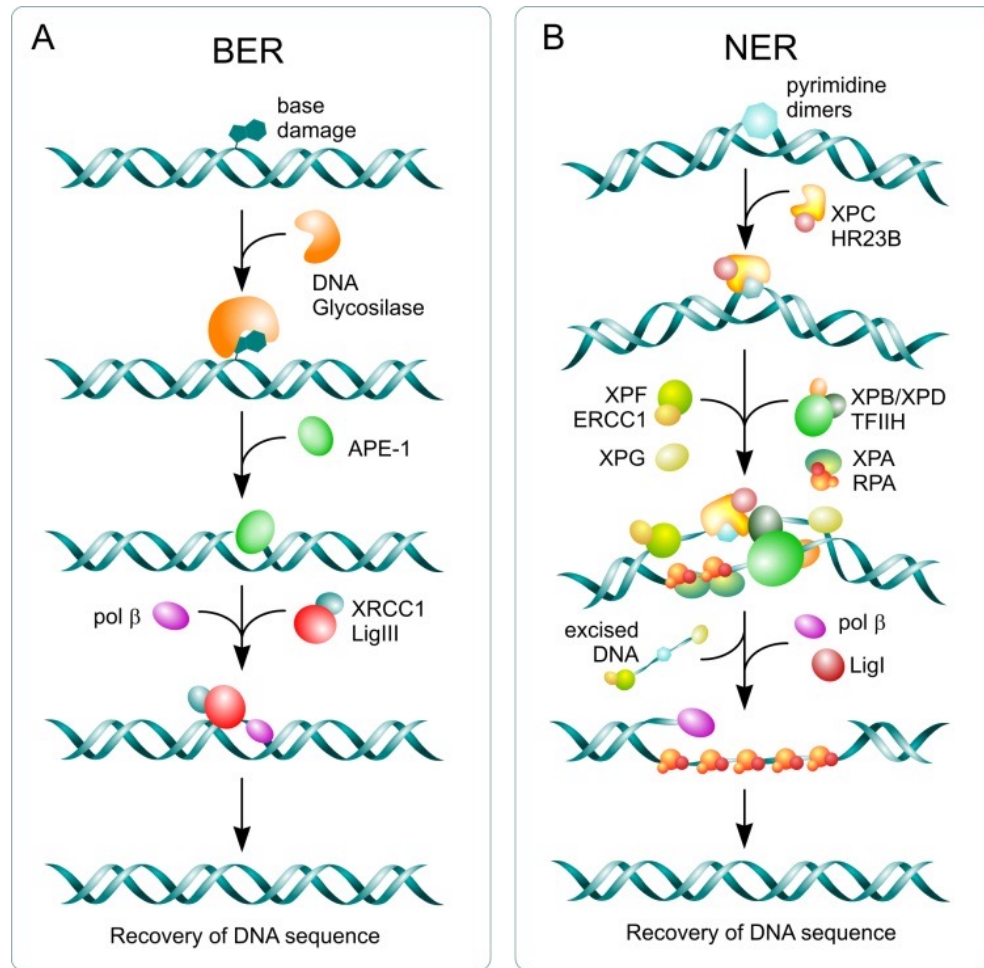
DNA damage repair – Single Strand Breaks

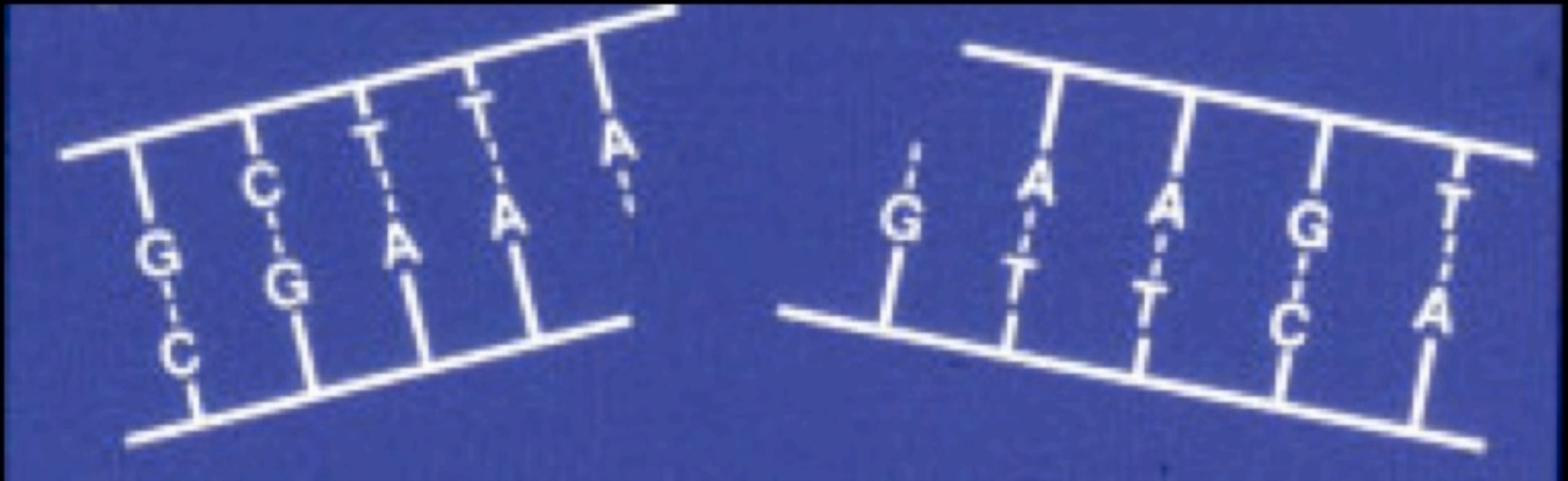
Base excision repair

- base damage
- induction of SSB
- synthesis of missing bases
- annealing

Nucleotide excision repair

- bulky lesions
- induction of SSB
- excision of 20-30 BP
- synthesis of missing bases
- annealing





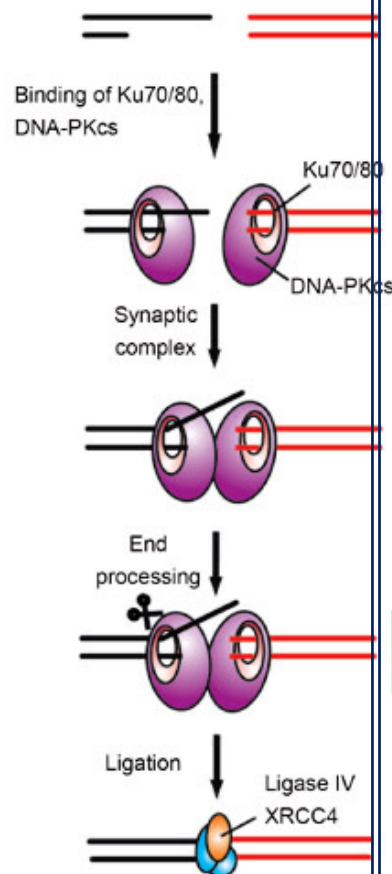
DNA double break triggers cell death

DNA Damage Repair – Double Strand Breaks

Nonhomologous endjoining Homologous recombination

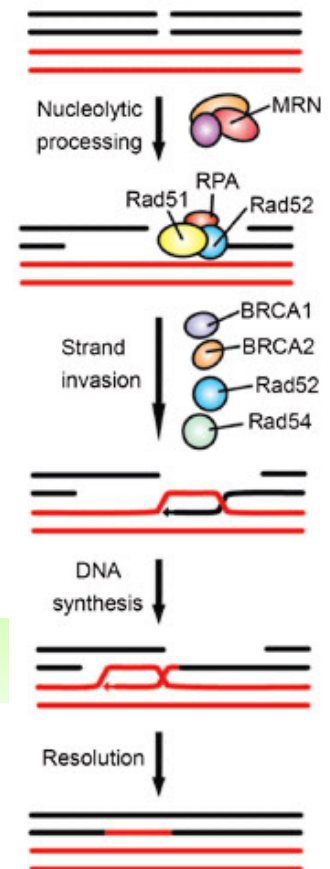
- >80 % of DSB
- G0 and G1 cell cycle phase

error prone

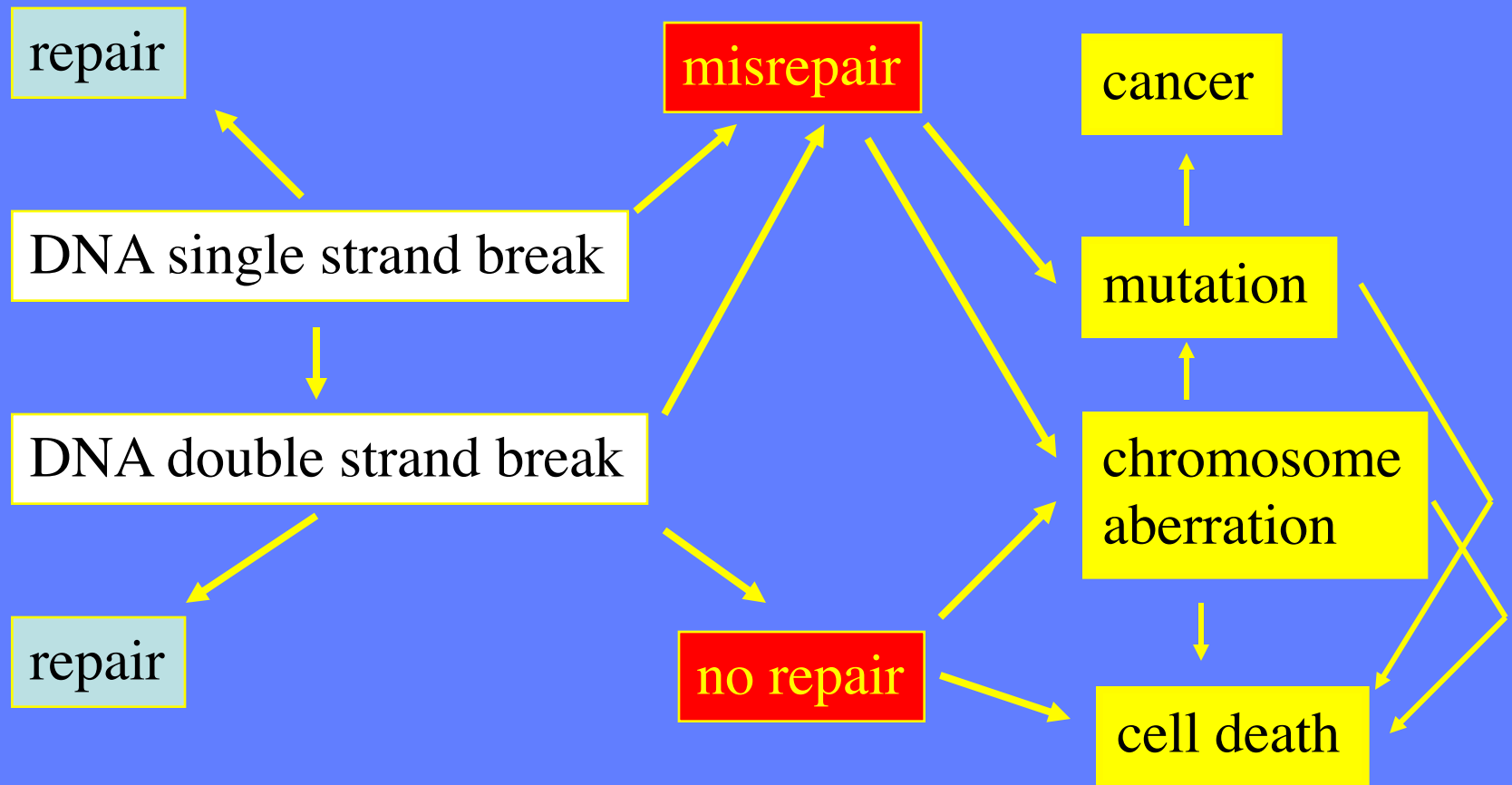


- < 20% of DSB
- S and G2 cell cycle phase

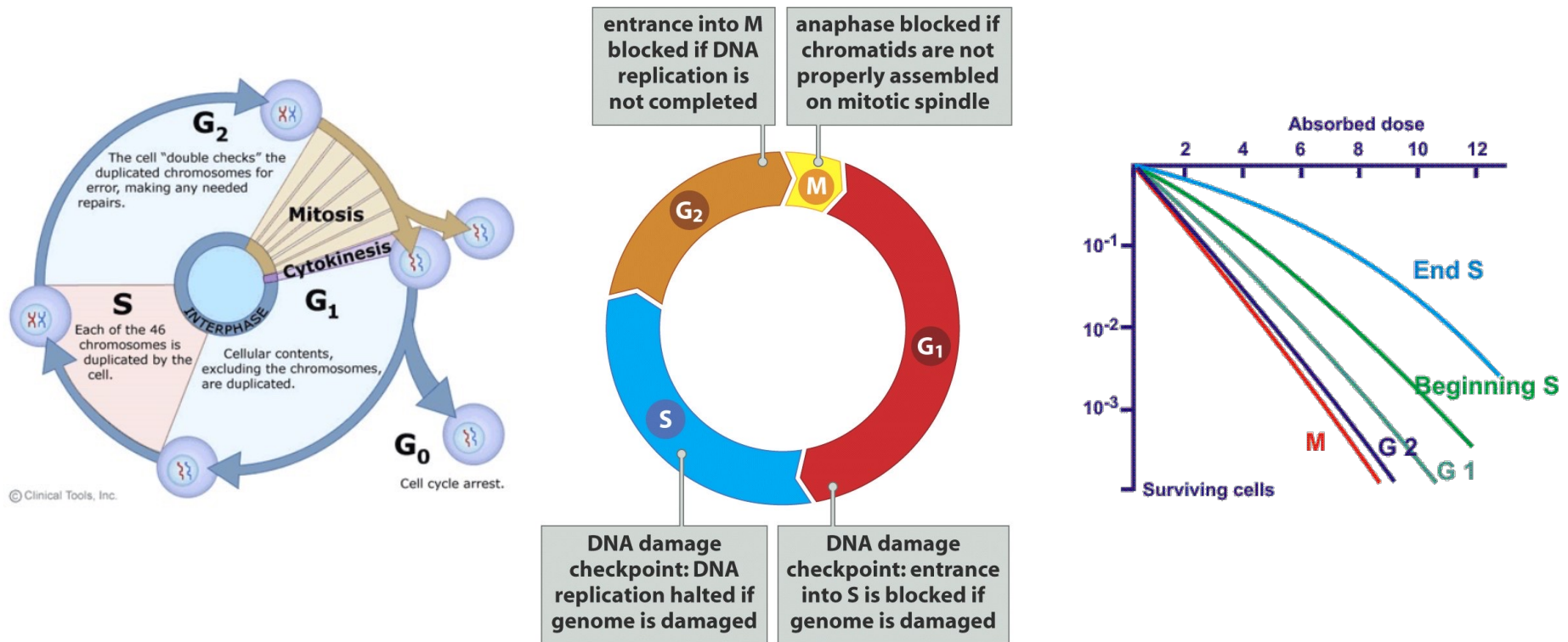
error free



DNA damage and its consequences



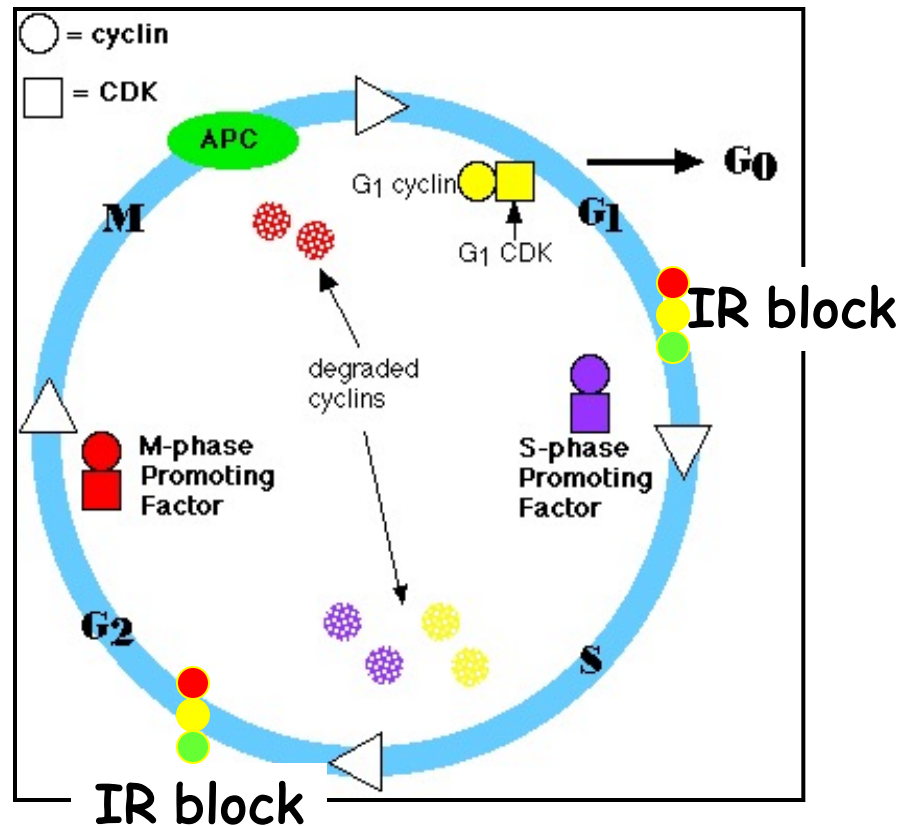
Cell Cycle Dependence Radiosensitivity



M>G₂>G₁>early S>late S for sensitivity

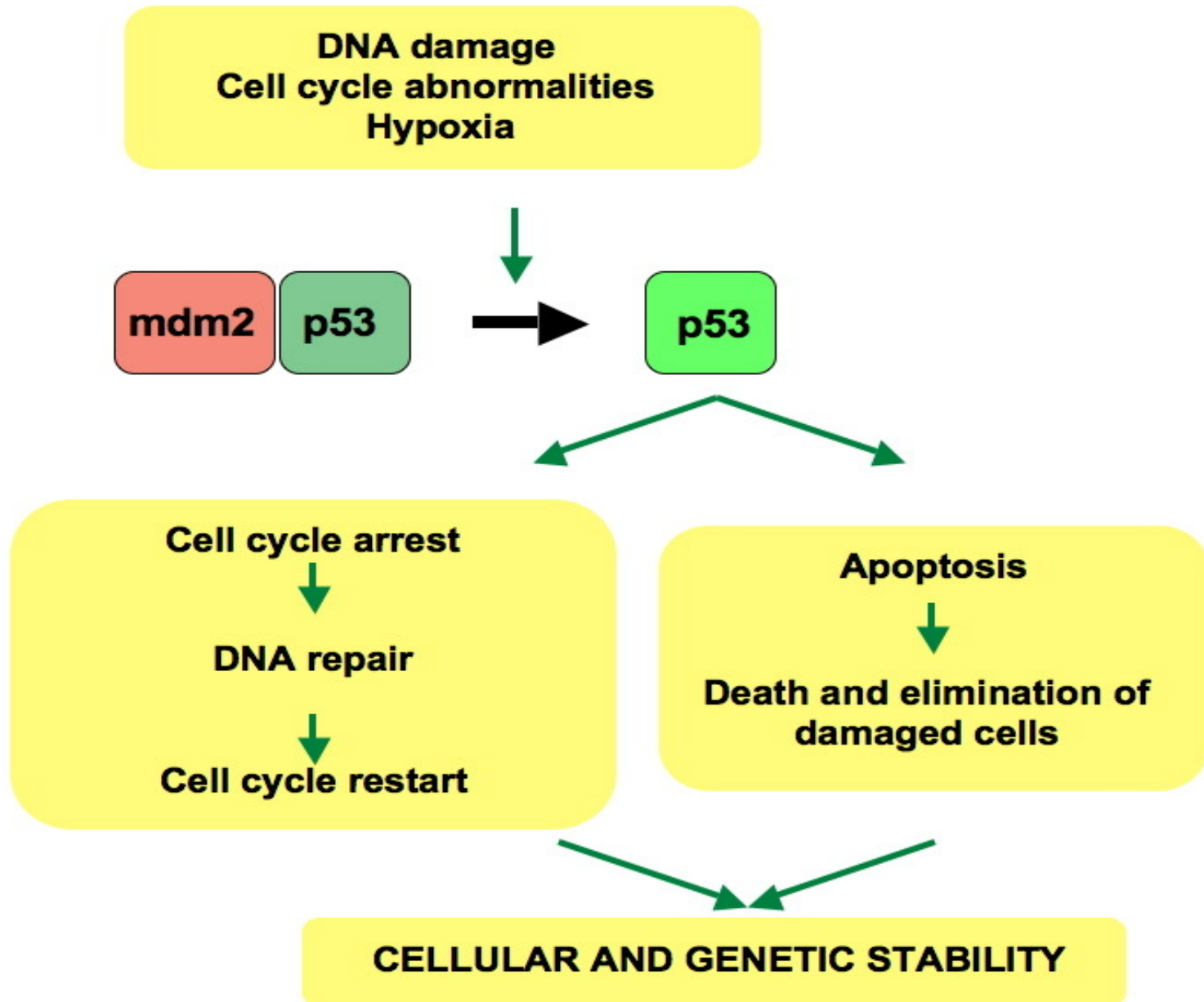
Difference caused by cell cycle are similar to difference caused by Oxygen effect

Regulation of the cell cycle



Cell cycle arrest  can occur in response to DNA damage (e.g. IR) in order to allow for DNA repair.

p53-guardian of the genome.....Lane,1992



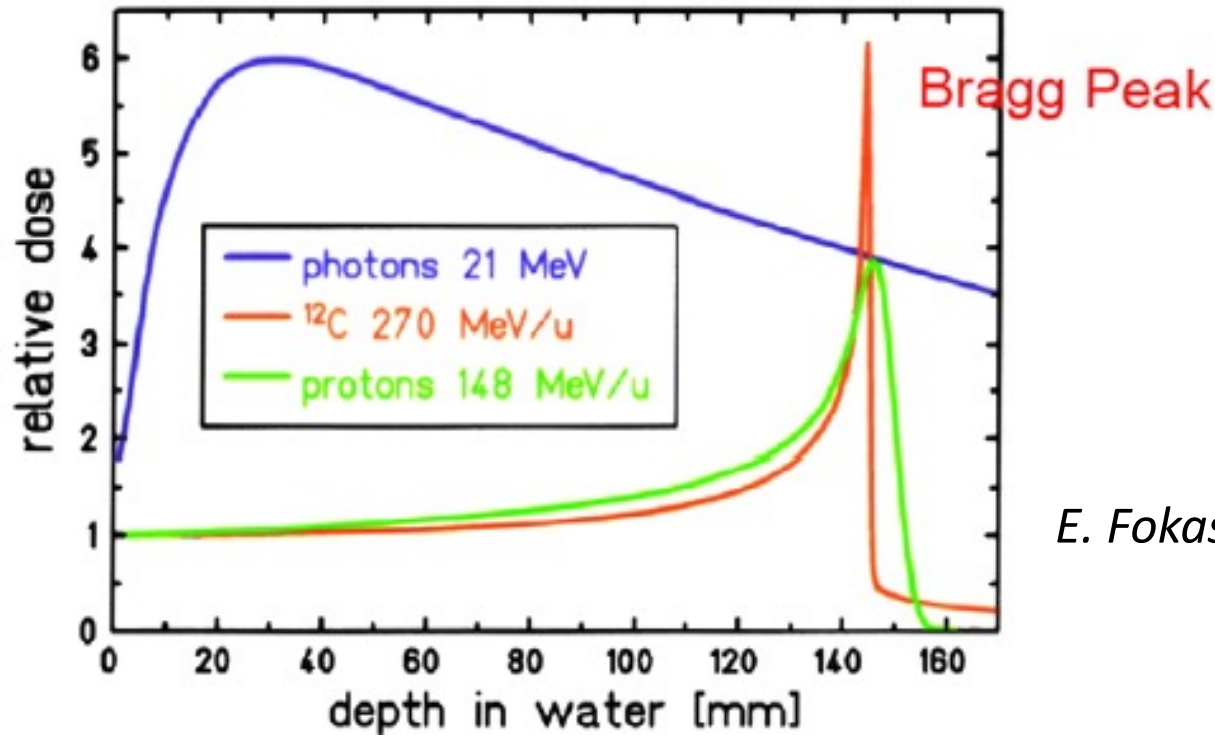
4 Rs of radiotherapy now 6 Rs

- Repair (few hours)
- Redistribution (few hours)
- Repopulation (5-6 weeks)
- Reoxygenation (hours to days)

Recently also added

- Intrinsic Radiosensitivity
- Reactivation of immune Response

Hadron Therapy vs classical RT



E. Fokas et al, 2009

In tumor therapy, hadrons, such as carbon produce a better depth-dose profile than protons.

The essential advantage of carbon ions is the higher biological effectiveness at the end of their range in the tumor. In the entrance channel the RBE is only slightly elevated.

In combination with the low dose in the entrance channel, as well as less more easily repairable damage is produced in normal tissue.

HT increasingly used due to better tumour targeting

Hadrons

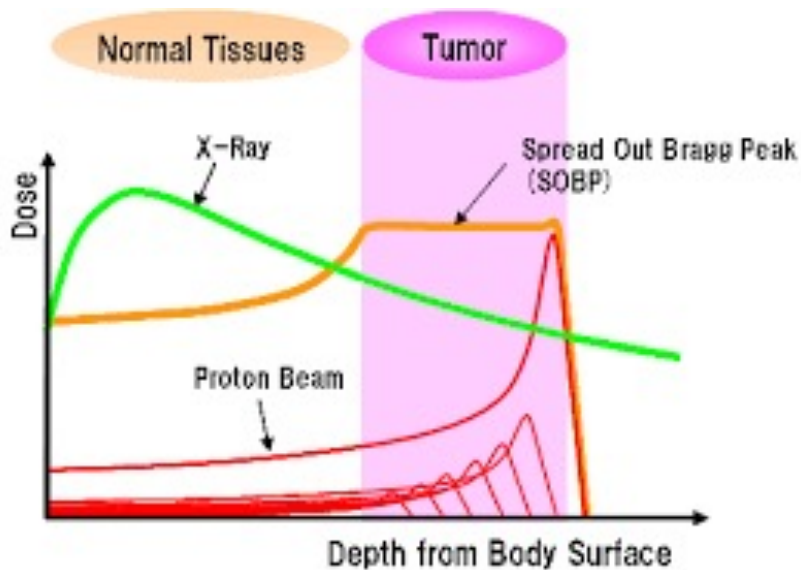
Dose highest at Bragg Peak

DNA damage not repaired

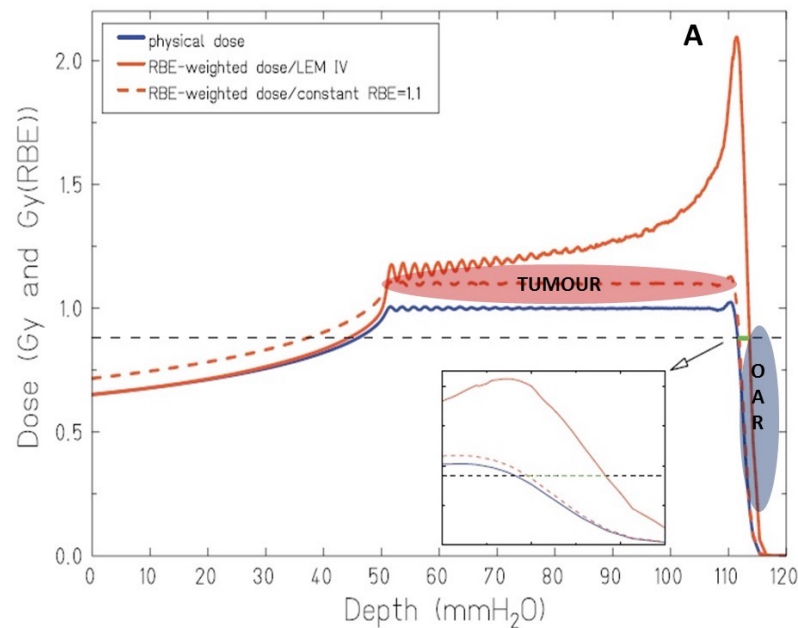
Biological effect high

Do not need oxygen

Effect is localised



Biological Range Uncertainty and actual RBE



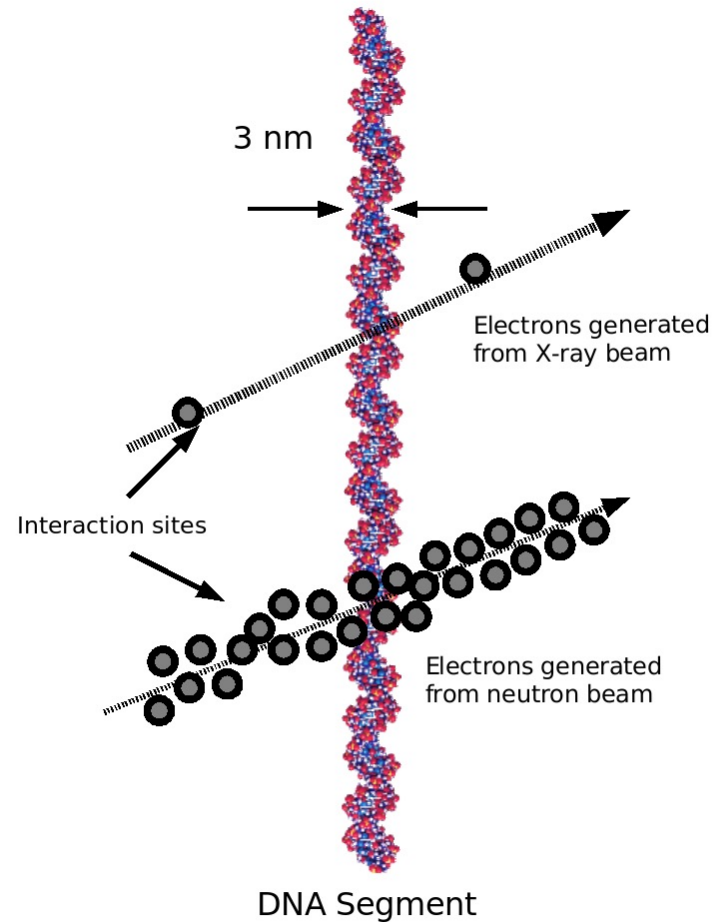
Distal SOBP

- Decrease of dose and increase in LET
- RBE is highly uncertain
- RBE can be greater than 1 in OAR
- Biological optimization needed

Linear Energy Transfer (LET)

It is the measurement of the number of ionisations which radiation causes per unit distance as it traverses the living cell or tissue

The LET depends on the charge and velocity of the ion: fast moving, light ions have low LET, and their biological effectiveness is close to that of X-rays; slow, heavy ions have high LET, and are more effective than X-rays for killing cells, as well as for other end points





FLASH: a new way of delivering Radiotherapy for treating cancer?



UNIVERSITY OF
OXFORD

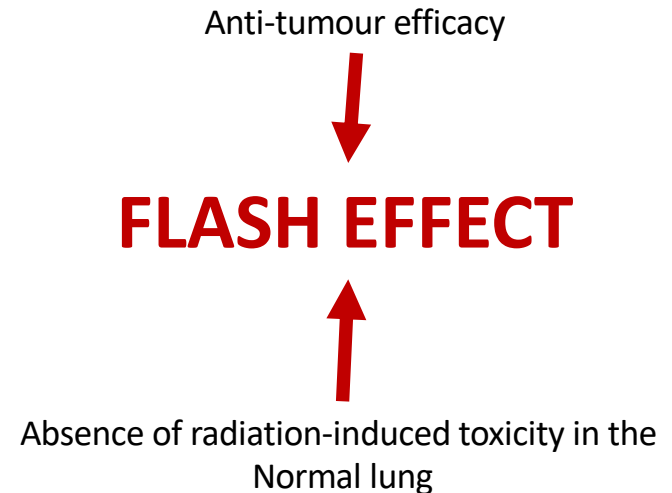
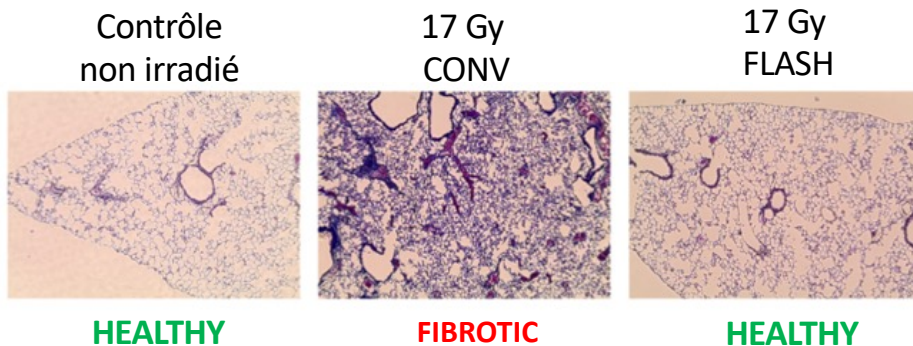
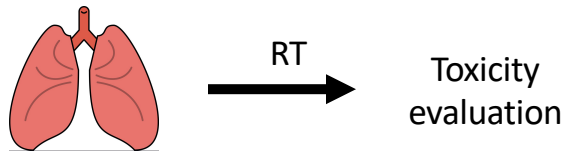
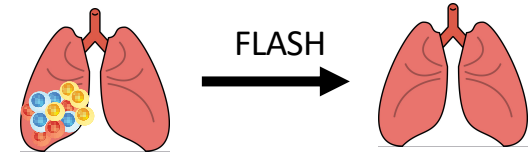
FLASH radiotherapy is based on the observation that healthy tissue is less damaged if treatment occurs very fast

RESEARCH ARTICLE

RADIATION TOXICITY

Ultrahigh dose-rate FLASH irradiation increases the differential response between normal and tumor tissue in mice

Vincent Favaudon,^{1,2*} Laura Caplier,^{3†} Virginie Monceau,^{4,5‡} Frédéric Pouzoulet,^{1,2§}
Mano Sayarath,^{1,2¶} Charles Fouillade,^{1,2} Marie-France Poupon,^{1,2||}
Isabel Brito,^{6,7} Philippe Hupé,^{6,7,8,9} Jean Bourhis,^{4,5,10} Janet Hall,^{1,2}
Jean-Jacques Fontaine,³ Marie-Catherine Vozenin^{4,5,10,11}



Treatment of a first patient with FLASH-radiotherapy

5.6 MeV linac adapted for accelerating electrons in FLASH mode

15 Gy with 10 pulses in **90 ms**

3.5 cm diameter tumour, multiresistant cutaneous

Appears that instantaneous dose
Induces a massive oxygen consumption
and a transient protective hypoxia in
normal tissues



Fig. 1. Temporal evolution of the treated lesion: (a) before treatment; the limits of the PTV are delineated in black; (b) at 3 weeks, at the peak of skin reactions (grade 1 epithelitis NCI-CTCAE v 5.0); (c) at 5 months.



First Patient Treated in FAST-01 FLASH Proton Therapy (November 2020) Transmission-shoot through

FeAsibility Study of FLASH Radiotherapy for the Treatment of Symptomatic Bone Metastases). The clinical trial involves the investigational use of Varian's ProBeam particle accelerator modified to enable radiation therapy delivery at ultra-high dose rates (dose delivered in less than 1 second) and is being conducted at the Cincinnati Children's/UC Health Proton Therapy Center with John C. Breneman M.D.

The study will assess Varian's ProBeam particle accelerator modified to deliver an advanced non-invasive treatment for cancer patients. *(Credit: Bokskapet from Pixabay)*

SUMMARY: Role of Radiobiology in Therapy

- Radiobiology research is essential in improving therapy
- The RBE is defined as the ratio of doses to reach the same level of effect when comparing two modalities and depends on multiple physics- and biology related parameters
- RT-induced DNA-damage and its repair is key in achieving better outcomes
- Role of 4Rs which are now 6Rs in treatment: repair, repopulation, redistribution, reoxygenation, intrinsic radiosensitivity and now reactivation of immune response
- New emerging modalities such as VHEE and FLASH

Many thanks to:

- U. Amaldi, CERN & TERA
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- M Durante, GSI, Germany; Kevin Prise, Queens, UK
- K. Prise, Queens
- A. Faocetti, CNAO, Italy
- Martin Pruschy, Zurich, Switzerland
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