$\mathop{\rm MP}_{\it pqi}$

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Welcome

This is my personal understanding and notes about Medical physics.

8 CONTENTS

Introduction

Physics constant 1.1

The constants are available from the website (http://physics.nist.gov/cuu/Constants/) supported by National Institute of Science and Technology (NIST).

The important constants used in medical physics are:

- Avogadro constant: $N_A = 6.022 \times 10^{23} \; \mathrm{mol}^{-1}$
- Speed of light in vacuum: $c = 2.998 \times 10^8 \text{ m/s}$
- Atomic mass constant: $u = 1.661 \times 10^{27} \text{ kg} = 931.5 \text{ MeV/c}^2$
- Elementary charge: $e = 1.602 \times 10^{19} \; \mathrm{C}$
- Electron rest mass: $m_e = 9.109 \times 10^{31} \ {\rm kg} = 0.5110 \ {\rm MeV/c}^2$
- Proton rest mass: $m_p = 1.673 \times 10^{27} \ \mathrm{kg} = 1.007 \ \mathrm{u} = 938.3 \ \mathrm{MeV/c}^2$
- Neutron rest mass: $m_n=1.675\times 10^{27}~{\rm kg}=1.009~{\rm u}=939.6~{\rm MeV/c}^2$ Planck constant: $h=6.626 \times 10^{34}~{\rm J\cdot s}=4.136\times 10^{15}~{\rm eV\cdot s}$

The SI system of units

The 7 base quantities and their units are

- Length *l* meter (m)
- Mass m kilogram (kg)
- Time t second (s)
- Electric current I ampere (A)
- Temperature T kelvin (K)
- Amount of substance mole (mol)
- Luminous intensity candela (cd)

1.2 Atomic Representation

Atoms = Nucleus (neutron and protons) 1 + Orbital electrons 2

¹Rutherford interpreted the results of the gold foil experiment or Geiger-Marsden experiment and established the Rutherford model of atom, which constitutes a tiny (10⁻¹⁵ m), heavy nucleus which consists of protons and/or neutrons. He also won the Nobel Prize in Chemistry 1908 "for his investigations into the disintegration of the elements, and the chemistry of radioactive substances". He discovered three types of radiation: α , β , and later γ radiation.

²In 1913, Bohr proposed a theory for the hydrogen atom based on **quantum theory** that (a) electrons orbit around the nucleus; (b) electrons orbits at a certain discrete set of distances from the nucleus without radiation and energy loss; (c) electrons can only gain and lose energy by jumping from one allowed orbit to another, absorbing or emitting electromagnetic radiation

$$_{Z}^{A}X$$

- A (mass number) the number of protons and neutrons
- Z (atomic number) the number of proton number
- X chemical symbol for the element

Atomcs can be classified in terms of the number of protons, neutrons, mass, and (meta)state.

- Isotope
- Isotone
- Isobar
- Isomer (same A, Z, N but different energy (meta)states; eg $^{99m}_{43}Tc$ is in metastable³ state and $^{99}_{43}Tc$ is in stable state)

1.3 Stability

The stability depends on the ratio of neutron and proton (see Figure 1)

1.4 Mass Defect

The mass of an atomic nucleus is less than the sum of the individual masses of the free constituent protons and neutrons. This "missing mass" is known as the mass defect.

For example, the mass defect of a ¹²C atom can be calculated by:

$$6 \times m_p + 6 \times m_n + 6 \times m_e - m_C = 0.0988 \ amu$$

where $m_C = 12$ (the ratio of mass to 1 amu). The complete list of mass number can be found in the NIST database.

The mass defect is closed related to nuclear bind energy. If we divide the above energy by 12 and times 931.5 MeV/amu, we obtain the bind energy per neucleio for ¹²C is 7.67 (see figure below)

A complete table of nuclear bind energies can be found on Lawrence Berkeley National Laboratory (link).

1.5 High energy charged particles

The mass of a moving particle (not a photon) depends on its velocity v and its rest mass m_0 .

$$E_{total} = mc^2 = \frac{m_0 c^2}{\sqrt{1 - \frac{v^2}{c^2}}} \tag{1.1}$$

or

$$E_{total} = E_{rest} + E_{K,E} \tag{1.2}$$

For an electron and a proton accelerated to the velocity of 0.96 c, the kinetic energy will be about 2 MeV and 2400 MeV. Therefore, high energy electrons coming out of linac head will have the speed close to the speed of light. To achieve similar high speed, you need give much more energy to a proton than an electron.

with a frequency: $v = \frac{E_m - E_n}{h}$. He won the Nobel Prize in Physics 1922.

³Metastable state is an excited state of an atom that has a longer lifetime than the ordinary excited states but generally has a shorter lifetime than the lowest, often stable, energy state, called the ground state. britannica

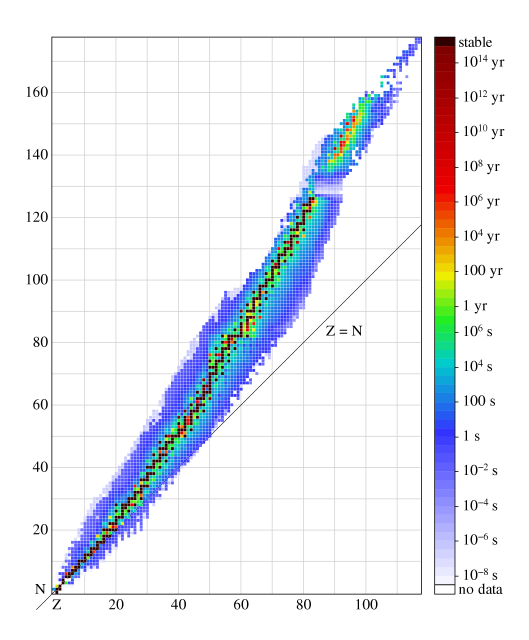


Figure 1.1: Stability of isotopes

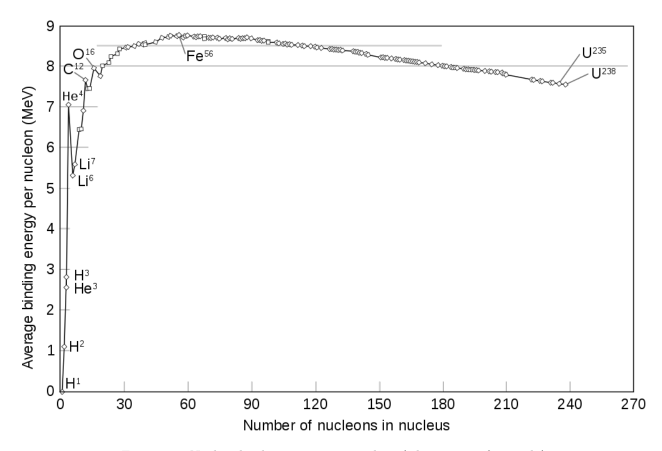


Figure 1.2: Nuclear binding energy per nucleon (The image is from wiki)

1.6 High energy photons

The energy of a photon is given by

$$E = h \cdot v \tag{1.3}$$

where h is the Planck constant, v is the frequency in unit of $s^{\{-1\}}$.

Or

$$E(eV) = \frac{1.24 \times 10^6}{\lambda(m)},\tag{1.4}$$

1.7 Electron Shell

• Principal quantum number (n = 1, 2, 3, ... or K, L, M, ...) – the main energy level (or shell) occupied by an electron. The energy can be calculated by

$$E_n = \frac{Z^2 \hbar^2}{2m_0 \alpha_B^2 n^2},$$

where α_B is the Bohr radius $(5.29 \times 10^{-11} m)$. The **K** shell (binding) energy for Lead, Tungsten, and Carbon are 88, 69.5, and 0.28 KeV.

- Secondary quantum number (l=s,p,d,...) the energy sublevel (angular momentum) occupied by the electron.
- Magnetic quantum number $(m_l = -l, -l + 1, ..., 0, ..., l 1, l)$ the number of possible orientations (projections) for each of energy sublevels.
- Spin quantum number $(m_s = -1/2, 1/2)$ the two possible orientations that an electron can have in the presence of a magnetic field.

1.8 Solutions

Q15: b) using Eq. (1.4) Q16: c) using Eq. (1.4)

```
Q1: a), c), (e) see Section 1.2
Q2: b), d); a) is wrong because they are isotopes; c) is wrong because they are isobars.
Q3: a), b), and c)
Q5: b), c); a) should be 6 neutrons and d) should be 12 times 931 MeV.
Q6: see above
Q7: b)
Q8: d) see Section 1.5
Q9: c)
Q10: a)
Q11: b)
Q12: a), b), c), e)
Q13: b), c)
Q14: c)
```

Nuclear Transformation

Radioactivity was discoverred in 1986 by A.H. Becquerel when he wa 44 years old. He received 1903 Nobel Prize in Physics along with Maria and Pierre Curie.

Radiation Sources (Siebers 2009 AAPM talk)

- Radioactive decay (Chapter 2)
 - Alpha-decay
 - Beta-decay (Section 23.4)
 - Electron capture
 - Isometric transitions
 - Gamma-ray
- Atomic energy transitions
 - Characteristic x-rays
 - Auger electrons
- Accelerated charge particles
 - Direct (electrons, protons)
 - x-ray generators (synchrotron radiation (magnetic field), Bremmstrahlung)
- Interaction products (?)

2.1 Decay (disintegration)

General balance equations of radioactive decay

$${}_{Z}^{A}P = {}_{Z-Z_{R}}^{A-A_{R}}D + {}_{Z_{R}}^{A_{R}}R + \sum Q, \tag{2.1}$$

where P and D stand for parent and daughter element, R for radiation, and Q is reaction energy ($\sum Q = M_P - M_D - M_R$). To find out the Q-value, you can use a online Q-calculator (http://www.nndc.bnl.gov/qcalc/).

Atoms found in nature are either stable or unstable. An atom is unstable (radioactive) if these forces are unbalanced if the nucleus has an excess of internal energy.² The instability of a radionuclide may result from an excess of either neutrons or protons. Radionuclides attempt to reach stability through

- 1. ejecting neutrons and protons (C area; Alpha-decay);
- 2. converting one to the other with ejection of a beta particle or positron (B area; Beta decay);
- 3. the release of additional energy by photon emission (Gamma decay).

 $^{^{1}{\}rm A~good~read~from~Wikipedia~(https://en.wikipedia.org/wiki/Henri_Becquerel)}.$

 $^{^{2} \}rm http://www.epa.gov/radiation/understand/radiation.html$

Alpha-decay occurs in nuclides with atomic numbers above 82 (only the first 92 occur naturally) and where the ratio of neutrons to protons is low, thus resulting in the repulsive coulomb force of the protons overcoming the attractive strong nuclear force.

Example
$${}^{226}_{88}Ra \rightarrow {}^{222}_{86}Rn + {}^{4}_{2}\alpha + \gamma + Q$$

Beta-decay, a neutron within the nucleus is converted into a proton, and an electron and an antineutrino are emitted, or a proton is converted into a neutron, and a positron and a neutrino are emitted. The forces responsible for the β -decay are weak (referred to as weak nuclear force) compared with both the strong nuclear force and the electrostatic force among the nucleons.

Example
$$\beta^-$$
 decay $^{27}_{60}Co \rightarrow^{60}_{27}Ni^* + \beta^- + \bar{\nu}$
 β^+ decay $^{18}_{9}F \rightarrow^{18}_{8}O + \beta^+ + \nu + 1.022 \text{ MeV}$

Neutrino (ν) and anti-neutrino $(\bar{\nu})$ results in spectrum of β energies, and they are non-ionizing particles so we don't consider them in dose calculation.

Electron capture (EC) is an alternative to positron decay. In this process, an electron, usually in the K shell, is captured within the nucleus and combined with a proton to create a neutron. Electron capture most often is followed by characteristic x-ray or Auger electron.

Gamma decay occurs when a nucleus undergoes a transition from a higher to a lower energy level. These γ -rays are identical to the x-rays emitted by excited atoms, except that γ -rays originate from within the nucleus and x-rays originate from outside the nucleus.

Example ${}^{60}_{27}Ni^*$ decay to stable ${}^{60}_{27}Ni$ by emitting two γ -rays with energies of 1.17 and 1.33 MeV.

The decay scheme can be found (http://atom.kaeri.re.kr:8080/gamrays.html)

2.2 Activity

The activity (A) of a sample is the average number of disintegrations (decay) per second,

$$A = \frac{\Delta N}{\Delta t} = \lambda N,\tag{2.2}$$

where λ is the decay constant which is the probability that a nucleus will decay per second. Remember that Radioactive decay is a **stochastic** process. We can find certain laws only by observing a large number of events (decays here).

From the equation above, we can obtain the radioative decay law at a certain time t:

$$N = N_0 e^{-\lambda t},\tag{2.3}$$

or

$$A = A_0 e^{-\lambda t}. (2.4)$$

More frequently, we use half-life time $(T_{1/2})$ instead of the decay constant λ . Their regulationship is

$$T_{1/2} = \frac{\ln 2}{\lambda}.$$
 (2.5)

2.3. UNIT 17

The mean or average life is the (arithmetic) average lifetime for the decay of radioactive atomes.

$$T_a \equiv \frac{1}{\lambda} = 1.44T_{1/2}.$$
 (2.6)

2.3 Unit

The SI unit for radioactivity is *Becquerel* (Bq). The historic unit for radioactivity is Curie (Ci), and 1g of radium is 1 Ci. The relationship between Curie and Becquerrel is

$$1 Ci = 3.7 \times 10^{10} Bq (2.7)$$

In practice, the more frequently used formula is

$$1 \text{ GBq} = 27 \text{ mCi} \tag{2.8}$$

2.4 Solutions

Q1 Decays

Using Eq. (2.3) or (2.4) and (2.5), we get

Residual activity =
$$1 - 0.02 = e^{-\frac{\ln 2}{30}t} \rightarrow \boxed{t = 0.87 \text{ years}}$$

It is easy to solve the above equation, but it will be faster to find a good estimation using the Taylor's expansion with first two terms $e^{-\frac{ln^2}{30}t} \approx 1 - \frac{0.693}{30}t$. The caveat of using Taylor expansion is make sure the exponents are much smaller than 1. You can try this approach for question 3, but you will not get the correct answer.

Q2 b), e)

Q4 a) b)

Q5 c)

Q6 Calculation of total decay

Decay_{total} =
$$1.44 \times T_{1/2} \times A$$

= $1.44 \times 30 \times 3.15 \times 10^7 \times 3.7 \times 10^9$
= 5.04×10^{18}

Q6 Average life time

$$A = A_0 e^{-\lambda T_a} = A_0 e^{-\lambda \frac{1}{\lambda}} \to \frac{A}{A_0} = e^{-1} \approx \boxed{37\%}$$

For question 6, with 1 year = 31536000 s and Eq. (2.7), the total number of decays is equal to total activity of 10 mCi Cs-137 is

$$10e^{-\frac{0.693}{8.05}t} = 4e^{-\frac{0.693}{14.3}t} \xrightarrow{\text{take ln() on each side}} ln10 - \frac{0.693}{8.05}t = ln4 - \frac{0.693}{14.3}t \rightarrow t = \boxed{24.3 \text{ days}}$$

Q7 c)

 $\mathbf{Q8}$

b); For higher electrons coming out of linac head, the electron velocity is close to the speed of light.Q9 a)Q10 b)Q11 d)Q12 b) d)Q13 c) $^{\circ}$

Production of X-rays

3.1 History

- Cathod-ray tube (for example, Crookes tube)¹.
- 1895-11-08, Wilhelm Rontgen discovered x-ray by observing fluorsece when applying high volotage (using an induction coil) on a board-covered Crookes tube (wiki and The Laureates: William Roentgen.
- Coolidge developed the **hot cathode** x-ray tube in 1913, in which a wire filament was heated with electrical current to release electrons by the process of **thermionic emission**. This was the first major breakthrough as cathod-ray tubes cannot generate reliable and high-intensity x-ray.
- The next major breakthrough in x-ray tube design was the rotating anode, which was developed by Albert Bouwers in 1930.

The Coolidge tube was the prototype for x-ray tubes in use today.

3.2 Conventional x-ray tubes

The basic components of a useful x-ray tube include: (a) electron source, (b) high voltage supply, (c) target for x-ray production, (d) vacuum, and (e) collimator.

3.2.1 Electron Source

- Tungsten (melting point is 3370 °C).
- The filament is housed within a negatively charged focusing cup.
- With high voltage applied across the tube, thermonic electrons are attracted to the target (anode) without electrons pile-up around the filement. In this scenerio, the tube is operated in the mode of filament-emission limited.
- With lower voltage applied across the tube, thermonic electrons are not pulled immediately to the target. The accumulated electrons (*space charge*) will prevent additional electrons leaving the filament and therefor limit the tube current. Under this condition, the tube is operated in the mode of space-charge limited (e.g. mammography machine).
- Dual-focus x-ray tube

¹partially vaccum and low voltage; think about neon lighting; see wiki



Figure 3.1: First medical X-ray by Wilhelm Röntgen of his wife Anna Bertha Ludwig's hand. (The image is from wiki)

3.3. X-RAY SPECTRA 21

3.2.2 High voltage

The electric potential difference (voltage) between the filament (cathode) and target (anode) of an x-ray tube affects the x-ray output, **intensity** (see electron source part) and spectrum of x-ray.

high-frequency x-ray generators (1-100 kHz)

kVp: maximum or peak volotage

Applications

- \bullet CT-simulator
- kV-CBCT

3.3 X-ray spectra

'The efficience is

Clinical Treatment Generators

4.1 History

- Cyclotron Earnest O. Lawrence 1932
- Betatron Donald W. Kerst 1940
- Cobalt machine
- Linear accelerator 1950s
- Gamma knife Lars Leksell 1968

4.2 Waveguide

4.3 Microwave amplifier

The possibly best simple explanation about how a klystron amplifier and microwave oscillators work can be found on YouTube.

A Klystron is a microwave (300 MHz – 300 GHz) amplifier tube that makes use of two (or more for better bunching result) resonant cavities. For a simple two cavity Klystron,

- 1. The first resonance cavity is energized by very low-power microwaves through a coaxial cable.
- 2. The microwave will cause alternating "E" fields across the gap between left and right cavity wall.
- 3. As the electrons from the accelerated through the first cavity, half of them will be decelerated and the other help will accelerate (velocity modulation), and thus form electron bunches as they drift towards the second cavity.
- 4. The Catcher cavity is resonant at the arrival frequency of the bunch.
- 5. This will generates a retarding "E" field for slowing down electrons and in turn the electrons give their energies in the form of high-power microwaves (more electrons in a bunch \rightarrow more kinetic energy \rightarrow more EM energies induced in the 2nd resonant cavity).

A Magnetron is a device that produces microwaves.

- 1. The electrons emitted from the heated cathode are accelerated by the pulse electric field, EP, toward the anode across the evacuated drift space between cathode and anode.
- 2. A static magnetic field, H, is applied perpendicular to the cross section of the device.
- 3. The accelerated electrons induce an additional charge distribution shown on the anode poles and an electric field Em of microwave frequency between adjacent segments of the anode (similar to that in the catcher cavity of the klystron).

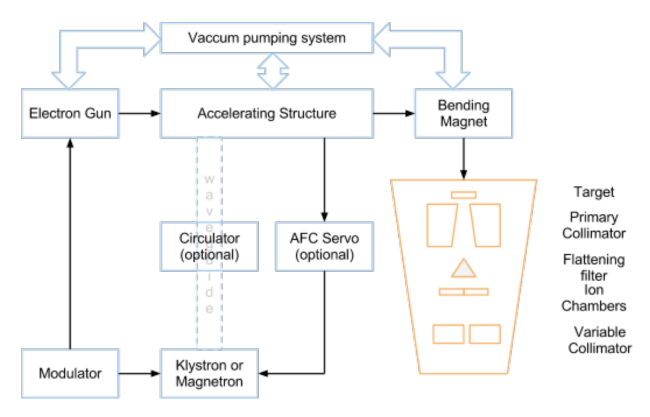


Figure 4.1: a simple schematic of a linear accelerator

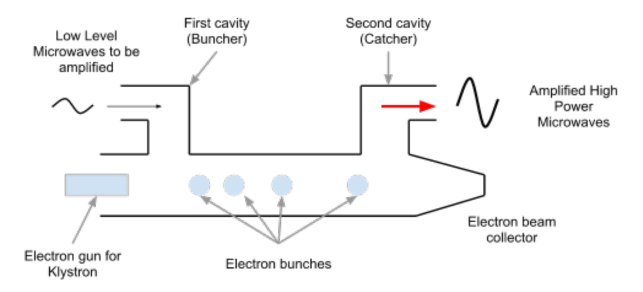


Figure 4.2: how a klystron works

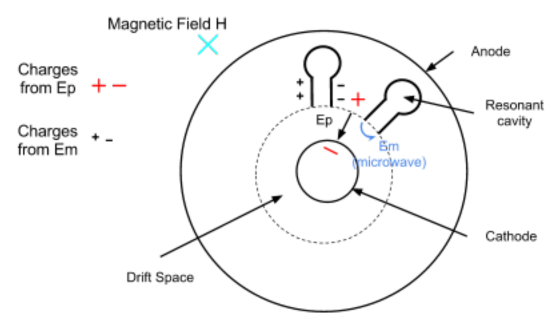


Figure 4.3: how a magnetron works

4.4 Microwave frequency

The microwave pulse frequency in most medical linear accelerators is about **3 GHz**, which falls into the category of IEEE S-band (2-4 GHz, Wiki). The Mobetron and Cyberknife machines use higher frequency (8-12 GHz, categorized as in IEEE X band, for compact design (Hanna 1999 Applications of X-band Technology in medical accelerators).

4.5 Penumbra

The term Penumbra means the region, at the edge of a radiation beam, over which the dose rate changes rapidly as function of lateral distance. The overall penumbra was contributed from three sources:

- Geometric penumbra is caused by the source (or focal spot) having a finite size and the location of the collimator. It can be reduced by decreasing the focal spot and move the collimator closer to the patient (e.g. Varian tertiary MLC).
- Transmission penumbra is caused by photons transmitted through the edge of the collimator. It can be reduced by aligning the collimator following the beam divergence (e.g. X and Y photon jaws).
- Physical (total) penumbra is the combination of transmission, geometric penumbra, and lateral scatter of radiation (photon and electrons) within the patient. Lateral electron disequilibrium (# of electrons projected laterally outward is not equal to # of electrons projected laterally inward). Because the range of these laterally projected electrons increases as energy increases, higher energy beams have a slightly greater penumbra than low energy beams.

Interaction

Follow the energy

5.1 Photoelectric interactions

The probability¹ of photoelectric interaction $\propto \frac{Z^3}{E^3}$.

- incident photon interact with bound atomic electron;
- all energy is given to electron;
- an orbital electron is ejected possessing most of incident photon, and a vacancy is present;
- Characteristic x-ray and Auger electron (The energy released by the downward transition is given to one of the outer electrons instead of to a photon).

5.2 Compton interactions

The probability of Compton interaction $\propto \rho_e$.

- interaction between incident high energy photons and loosely bound orbital electrons.
- With $\alpha = \frac{hv_0}{m_e c^2}$ and θ is the angle between incident and scattered photon, the scattered photon energy is

$$E_p = hv_0 \frac{1}{1 + \alpha(1 - \cos \theta)}.$$

- a. with $\theta = 0^{\circ}$ (glazing hit) electron acquires minimum energy, $\Delta \lambda = .00243 \times (1 \cos \theta)$;
- b. with $\theta = 90^{\circ}$ for megavoltage linars with $\alpha > 10$, scatter photons always have energy of about 0.5 MeV (shielding consideration);
- c. with $\theta=180^o$ (photon is scattered back) electron acquires maximum K.E and photon has an energy of 0.255 MeV.

Q2: d)

¹The basic quantity in collisional dynamics is cross section. The SI unit is cm^2 and the unit is barn (1 $b = 10^{-24} cm^2$) in nuclear physics.

5.3 Pair production

The probability of pair production $\propto Z \cdot E$.

- occurs when a photon approaches closely enough to the target nucleus;
- the incident photon energy may be converted directly into an electron-positron pair. When the positron comes to rest, it combines with an electron, and both particles then undergo annihilation, with the appearance of two photons with energy of 0.511 MeV traveling in opposite directions.

Q3: b) The threshold energy for pair production is 1.022 MeV.

5.4 Compton interactions

If the photon is scatter back at $\theta = 180^{\circ}$, the electron gains the maximum energy $hv \times \frac{2\alpha}{1+2\alpha}$.

$$\lambda' - \lambda = \frac{h}{mc^2} (1 - \cos\theta) \tag{5.1}$$

Attenuation of radiation is removal of photons or energy from a beam by different interactions including absorption and scatter. Like the process of radioactive decay, the attenuation is also a stochastic process

For a thin absorber, with absorber far away from the source (so effect of beam divergence is negligible e.g. ignore inverse square law²), or in a **narrow** beam geometry, we get $-\Delta N/N = \mu \Delta x$, where μ is linear attenuation coefficient which can be thought as the fraction of photons or energy removed from beam per cm of absorber beam per cm. Half-value layer (HVL) relates to the linear attenuation coefficient by

$$HVL = \frac{0.693}{\mu} \tag{5.2}$$

Mass attenuation coefficient is often used to remove the dependence of the physical density.

$$\left(\frac{\mu}{\rho}\right) \propto \frac{\sigma_{tot}}{\rho} = \frac{\sigma_{coh}}{\rho} + \frac{\sigma_{pe}}{\rho} + \frac{\sigma_{comp}}{\rho} + \frac{\sigma_{pair}}{\rho} + \frac{\sigma_{trip}}{\rho} + \frac{\sigma_{ph.n}}{\rho}$$
(5.3)

²The intensity of a point radiation source follows inverse square law. This a kind of geometric concept as the area of a sphere is $A = 4\pi r^2$. The inverse square law is valid under two assumptions: (1) point source, i.e. small enough compare to distance; (2) photon undergoes no interaction (e.g. TBI with spoiler).

Measurement of Ionizing Radiation

Attempts were made to measure ionizing radiation based on chemical and/or biological (skin) effects. But those measurements were not reliable. The ICRU adopted the roentgen, denoted by R, as the unit of measuring x- and γ -ray exposure.

The ICRU No.33 (1980) (1980) definition of exposure:

$$X = \frac{dQ}{dm},$$

where dQ is the absolute value of the total charge of the ions of one sign produced in air when all the electrons (negatrons and positrons) liberated by photons in air of mass dm are completely stopped in air.

From the book "Fundamentals of Radiation Dosimetry" (chapter 5 - good chapter to read): the photons first interact with a defined mass of air. They will produce electrons by the photoelectric and Compton effect and both electrons and positrons by the pair production process. All those secondary charged particles must travel through the air until their energy is

6.1 Collection volume

A free air ionization chamber has a 10 mm diameter aperture, a plate separation of 90 mm, and a collection length of 70 mm. Calculate the mass of air in the collection region.

$$mass = \rho \cdot V = 1.293 \ kg/m^3 \cdot \frac{1}{4}\pi \times (10 \ mm)^2 \times 70 \ mm = 2.3 \times 10^{-7} kg$$

6.2 Signal of an ion chamber

The signal from an ionization chamber is proportional to the charge (ionization) collected (so to the numbers of gas molecules in the cavity. Combining the ideal gas law $(P \cdot V = nRT)$, we have

$$signal \propto \frac{P \cdot V}{T}$$

In this case, $V_{unsealed} = (1/2)^3 V_{sealed}$ and P_unsealed = 1atm = 1/3 P_{sealed}.

6.3 Temperature and pressure correction

Most likely, the local measurement condition will be different from the standard environment condition ($22^{\circ}C$ and 760 mmHg) under which the ion chamber (and possible its electrometer) is calibrated. Therefore we need to correct the reading with a factor (AAPM TG-51):

$$P_{TP} = \frac{(237.2 + T)}{(273.2 + 22.0)} \times \frac{760}{P},\tag{6.1}$$

wehre T is in the unit Celsus and P is in the unit of mmHg.

Pressure drops about 1 inch per 1000 feet.

So the pressure is $760 - 3600/1000 \times 25.4 = 668.6$ mmHg. $P_{TP} = \frac{273.2+24}{273.2+22.0} \times \frac{760}{668.6} = 1.14$ $P_{TP_wrong} = \frac{273.2+24}{273.2+22.0} \times \frac{760}{760} = 1.006$; If we used the wrong P_{TP}, the "corrected" reading (machine output) will be thought as 13% lower than the actual value. If we increase linac output to compensate this 13% difference, we will overdose the patient by 13%.

Q6: a); Q7: d); Q8: d); Q9: b)

C); but this is different from my calculation! The plate separation of 90 mm is not used here.

Chp6 c c d e b a d d b b d bcd b bc acd

Rogers's talk

6.4 Guard electrode

The guard electrode in a Farmer-type chamber can (1) prevent leakage from the high-voltage collector electrode; (2) define the ion-collecting volume; and (3) minimize polarity effect (?).

Good reading materials include Deward A good document can be found here.

Figure Radiographs (above) and drawings (below) of five Baldwin–Farmer-style ion chambers plus an Exradin A12 . In the drawings, the heavy black lines represent the extent of guarding, as also indicated by the arrows on the left. The grey blocks indicate the insulator in closest contact with the active air volume, indicated by the arrows on the right. The A12 has no insulator other than air in contact with the active air volume. (PMB 50 N121, 2005)

Quality of X-rays

We have finished a nice book.

Absorbed Dose

"Perhaps one of the greatest contributions physics has made to radiation oncology and radiology, x-ray imaging and all of its forms has been in developing ways to measure radiation accurately and precisely (commonly 'using ion chamber)."

— Peter Almond

To measure the absorbed dose from ionizing radiation within a medium, we need to know

- 1. The number of particles or photons, or the quantity of energy, passing through the medium (fluence)
- 2. The quantity of energy transferred from initial particles (often photons, which are uncharged) to charged particles in the medium (KERMA)
- 3. The rate at which energy is transferred from the charged particles in the medium, to the medium itself (stopping power, leading to absorbed dose).

Fluence is defined as the number of particles dN incident on a sphere of cross-sectional area da. The SI unit is m^{-2}

$$\Phi = \frac{dN}{da} \tag{8.1}$$

Energy fluence $(\Psi, \text{ unit: } J \cdot m^{-2})$ is defined as the energy dE incident on a sphere of cross-sectional area da. The SI unit is $J \cdot m^{-2}$.

$$\Psi = \frac{dE}{da} \tag{8.2}$$

If you have a fluence Φ of particles all of energy E, then the energy fluence is simply $\Psi = \Phi \cdot E$.

KERMA (Kinetic Energy Released per unit MAss) is defined as the mean kinetic energy transferred to charged particles from uncharged particles in a mass dm of a given material. The SI unit is J/kg, and the special name for the unit for Kerma is gray (Gy).

$$K = \frac{d\bar{E}_{tr}}{dm} \left(J \cdot kg^{-1} \text{ or } Gy \right)$$
(8.3)

The relation between Kerma and fluence can be expressed as

$$K = \int \Psi(E) \frac{\mu_{tr}(E)}{\rho} dE$$

Where $\frac{\mu_{tr}(E)}{\rho}$ is the mass energy transfer coefficient of the material for uncharged particles of energy E.

Unrestricted stopping power for charged particles (electrons) is defined as

$$S = \frac{dE}{dx}$$

- Collisional stopping power (S_{coll})
- Radiative stopping power (S_{rad}) cause by the interactions of charged particles with nuclear electric field bremsstrahlung radiation The relationship of fluence and stopping power to absorbed dose is given by:

$$D_{med} = \int \Phi_{med,E}(E) \frac{S_{coll}(E)}{\rho} dE$$

8.1 Optical density

The details about the radiographic films can be found in AAPM TG-69: Radiographic film for megavoltage beam dosimetry.

8.2 Q9 OD

A pivotal assumption in film dosimetry is that the dose to the film is reflected in the resulting "blackness" or optical density (OD) of that film.

$$OD = log_{10}\left(\frac{1}{T}\right) = log_{10}\left(\frac{I_0}{I_t}\right)$$

The details about radiochromic films can be found in AAPM TG-55 and its update AAPM TG-235 as well an excellent review article by Butson et al. "Radiochromic film for medical radiation dosimetry" (2003). Table 1 lists the radiation interaction processes and their variation with Z.

Relative dosimeter

- Diode (single or 2D diode array MapCheck)
- TLD
- OSL
- MOSFET
- Film

Dose Distributions

9.1 TAR

The first three factors are used for the source-to-axis distance (SAD) technique (mechanical isocenter and radiation isocenter roughly coincidence with the tumor centroid).

With d is the depth from the surface to the isocenter in a phantom and r is the field size at the level of the isocenter, we can define

• Tissue-air-ratio (TAR) is defined by

$$TAR(d, r_d) = \frac{Dose_{phantom}(d)}{Dose_{air}}$$
(9.1)

• Backscatter factor (BSF) is a special case of TAR, in which $d = d_m ax$

$$BSF = \frac{Dose_{phantom}(d_m ax)}{Dose_{air}}$$

$$(9.2)$$

• Scatter-air factor (SAR) can be calculated by

$$SAR(d,r) = TAR(d,r) - TAR(d,0)$$

$$(9.3)$$

THe Mayneord factor is used to find a new PDD from a known PDD value

$$f = \frac{PDD_2}{PDD_1} = \left(\frac{SSD_2 + d_{max}}{SSD_1 + d_{max}}\right)^2 \cdot \left(\frac{SSD_1 + d}{SSD_2 + d}\right)^2$$
(9.4)

Dose calcuation

We have finished a nice book.

Treatment Planning I: Isodose Distribution and Plan Evaluation

11.1 Penumbra

The dose distribution outside the field boundaries is significantly affected by geometric penumbra, depth, leakage radiation through collimator. The flattening filter mostly affect dose within the field boundary.

11.2 Wedges

- Physical wedge
 - External physical wedge
 - Internal physical wedge (aka motorized wedge, as in ElektaTM machines) typically consists of a single large wedge (e.g., 60 degrees) placed above the secondary collimating jaws. The smaller angle is form by combining the open (o) field and the 60° degree wedge field:

$$Dose_{\theta} = W_o Dose_o + W_{60^o} Dose_{60^o},$$

where $W_{60^o} = \frac{tan\theta}{tan60^o}$.

- Non-physical wedge
 - Virtual wedge (as in SiemensTM)
 - Enhanced dynamic wedge (EDW) in VarianTM, which is implemented by moving one of the collimating jaws from one end of the field to the other.

Wedge (isodose) angle is defined as the angle between wedged isodose curve (see figure wedge isodose) and the normal to the central axis at a specific depth (e.g., 10 cm). What we typically measure is wedge profile.

Wedge Commissioning

- Salk et al Physical aspects in the clinical implementation of the EDW 1D ion chamber.
- Fontanarosa et al Commissioning Varian EDW in the PINNACLE treatment planning system using Gafchromic EBT film.
- Njeh EDW output factors for Varian 2300 CD and the case for a reference database.
- Shao et al the accuracy of dynamic dose computation in the ADAC Pinnacle RTP system.
- Zhu et al Performance evaluation of a diode array for EDW dosimetry mapcheck.
- Ahmad et al Study wedge factors and beam profiles for physical and EDW

Treatment Planning II: Patient Data, Corrections, and Setup

12.1 Inhomogeneity

In the presence of inhomogeneity, the dose calculation needs to address two issues (https://www.utoledo.edu/med/depts/radther/pdf/JC%20Chapter%2011%20handout.pdf):

- 1. Change in primary fluence (see Eq. (8.1)) due to change in attenuation
- 2. Change in scatter contributions.

calculation either indirectly through a correction factor (CF) or directly inherent in the algorithm (Papanikolaou AAPM presentation)

12.2 Range

The energy loss of electrons in a medium can be evaluated using mass stopping power (S/ρ) in unit of $\frac{MeV}{g\cdot cm^2}$

$$\left(\frac{S}{\rho}\right) = \left(\frac{S}{\rho}\right)_c + \left(\frac{S}{\rho}\right)_r$$

$$= \frac{\frac{dE}{dl}}{\rho}$$

The detailed information about stopping power for electrons can be found on the NIST website (https://www.nist.gov/pml/stopping-power-range-tables-electrons-protons-and-helium-ions).

In the range of therapeutic energies, 4 MeV to 20 MeV, the total mass stopping power is almost a constant, e.g.,

$$\left(\frac{S}{\rho}\right) \approx 2\frac{MeV \cdot cm^2}{g} \tag{12.1}$$

For water, the stopping power (S) is equal to $S = \left(\frac{S}{\rho}\right) \times \rho \approx 2\frac{MeV \cdot cm^2}{g} \times 1\frac{g}{cm^3} = 2\frac{MeV}{cm}$.

For an electron beam of energy E, which is specified as the most probable energy at the surface $(E_P)_0$, the practical range of a broad electron beam in water can be estimated by $R_P = E/12 MeV$.

12.3 MRI

Shimony's Youtube video and more resources at 2:41.

Basics

- 1. A strong, uniform magnetic field B_0^1 is applied (clinical: 1.5-7 Tesla and research: 7-11.7 Tesla);
- 2. The magnetic field will algin protons (hydrogen atoms) which are normally randomly orientaed within human boday. This can also explained as the magnetic creates two separated energy levels, and the energy difference is $\Delta E = hf$, and the frequency f, the resonance (Larmor) frequency, can be written as

$$f = \gamma \cdot B_0$$
,

where γ is called gyromagnetic ratio and is equal to 42.6 MHz/T. For B=3.0~T, the Lamor frequency is 130 MHz.

- 3. As
- 4. To excite the atoms from lower to higher energy levels (RF coil) and an additional magnetic field is applied in the x-y plane to create a flip angle (90° or 180°);
- 5. The emitted RF waves can be picked by an antenna; to relate the spatial information with precisely controlled magnetic field (gradient foil)

The RF signal for a spin-echo sequence can be written as

$$Singal = \rho \cdot M_Z \cdot \left(1 - e^{-\frac{TR}{T1}}\right) \cdot M_{XY} \cdot e^{-\frac{TE}{T2}},$$

- ρ is the proton density;
- M_Z and M_{XY} areh the magnetization along the Z and XY direction
- TR: repetition time time between each RF pulse;
- TE: echo time time between delivery of RF pulse and receipt of the echo signal.
- T1: longitudinal relaxation time a measure of the time taken for spinning protons to realign with the external magnetic field; for example, T1 = 4,000 ms and 250 ms for water and fat;
- T2: transverse relaxation time a measure of the time taken for spinning protons to lose phase coherence among the nuclei spinning perpendicular to the main field; for example, T2 = 250 ms and 70 ms for water and fat;

T1-weighted image is called (fluid) dark image and T2-weighted image is called (fluid) bright image

It is all about water and fat

12.4 PET

Positron decay (see Section 2.1)

$$B = \mu_0 I N / L,$$

where B is field strength, μ_0 is the permeability constant of free space $(1.27 \times 10^{-6} mks^{-2}A^{-2};$ about the same as those for water, hydrogen, and human body), I is current per turn, N is the number of turns, L is the coil length. For B = 1 T, L = 1 m, N = 10,000, the current will be around 80 A Aarnink. We thus have to use superconducting technique - thanks to Fermilab Tevatron.

¹For a simple long solenoid with uniform winding density, the magnetic field will be

Treatment Planning III: Field shaping, skin dose, and field separation

13.1HVL

To calculate transimssion or attenuation problems, you can use one of three formula with given parameters

- 1. $2^{-t/HVL}$ given HVL 2. $10^{-t/TVL}$ given TVL
- 3. $e^{-\mu t}$ given linear attenuation coefficients

You can directly calculate the result from $2^{-n \times HVL/HVL} \le 0.02$. Or using 0.02 = 1/50, $2^{-(-5)} = 1/32$ and $2^{-}(-6)=1/64$, we can guess the result is d).

Related references Calibration: TG-21 (1983) to TG-51 (1999) + Addendum to the TG-51 (2014) Parallelplate chamber: TG-39 (1994) Clinical electron therapy: TG-25 (1991) to TG-70 (2009) Total skin electron therapy: TG-30 (1987) IORT - Mobetron: TG-72 (2006) Comprehensive: ICRU Report 71 (2004) IAEA Radiation Oncology Physics Chapter 8

Q2, 3, 4, and 7 Range 13.2

The energy loss of electrons in a medium can be evaluated using mass stopping power (S/ρ) in unit of $\frac{MeV}{g\cdot cm^2}$

$$\left(\frac{S}{\rho}\right) = \left(\frac{S}{\rho}\right)_{s} + \left(\frac{S}{\rho}\right)_{s} = \frac{\frac{dE}{dl}}{\rho}$$

The detailed information about stopping power for electrons can be found on the NIST website (https: //www.nist.gov/pml/stopping-power-range-tables-electrons-protons-and-helium-ions).

In the range of the rapeutic energies, 4 MeV to 20 MeV, the total mass stopping power is almost a constant, e.g.,

$$\left(\frac{S}{\rho}\right) \approx 2 \frac{MeV \cdot cm^2}{g}$$

For water, the stopping power (S) is equal to $\left(\frac{S}{\rho}\right) \times \rho \approx 2 \frac{MeV \cdot cm^2}{g} \times 1 \frac{g}{cm^3} = 2 \frac{MeV}{cm}$.

$44 CHAPTER\ 13.\ TREATMENT\ PLANNING\ III: FIELD\ SHAPING,\ SKIN\ DOSE,\ AND\ FIELD\ SEPARATION$

For an electron beam of energy E, which is specified as the most probable energy at the surface $(E_P)_0$, the practical range of a broad electron beam in water can be estimated by 12MeV.

Electron

14.1 History

- late 1930s Van de Graaff Accelerators (at MIT by Van de Graaff and Trump); low energy < 3 MeV
- late 1940 Betatron; beam quanlity is not good
- 1960s linear accelerators

14.2 Treatment Sites¹

A lot sites (located with 6 cm of the surface) but only accounts for 10-15% of treatment.

- Head (Scalp, ear, eye)
- Breast/Chest wall
- Skin
- extremities

However, the competing technology (VMAT, BT, ...), inaccurate dose calculation (account for bolus scatter, backscatter, eye shield,...), and most importantly, lack of motivation from the vendor have reduced the number of electron treatment in radiotherapy.

14.3 Interactions

With orbital electrons

- Elastic collision
- Inelastic collision (ionization and excitation to higher energies) dose deposition

With nuclei

- Elastic collision
- Inelastic collision (Bremsstrahlung)

¹Electron Radiotherapy, Past, Present, and Future (https://vimeo.com/78553521)

14.4 Delivery

- Double scattering foil system (spread + flattern)². Excerpt from Niroomand-Rad: In a Siemens machine, the electron beams pass through dual scattering foils. The first (primary) foil, made of stainless steel, serves to scatter the electron beam. Its thickness is 0.075 mm for 5-7 MeV beam and 0.030 mm for 10 MeV beams. The second (secondary) foil, made of 0.8 mm thick aluminum, for all the electron beams, produces a homogeneous radiation mainly by absorption.
- Collimation cones (typically multi-leveled to block electron spread at different distance)
- Jaws set at a much larger size than the cone sizes

14.5 Beam quality

PDD

- Surface dose (70%-90%)
- R90 (therapeutic range $\sim E/4$) is the depth for tumor edge
- R10 R90 for estimating dose fall-off to spare oARs
- Rp (practical range $\sim E/2$) where beam stops
- x-ray contamination (from linac and phantom and patient, about 50% each)

With energy, field size, and SSD increase, PDD will increase, decreases, and stays roughly the same.

Example: Electron treatment with cicular cutout of 3 cm and 2 cm diameter. the measured output factor is 0.85 and 0.67

$$OP\left(d_{max}(r), r, SSD\right) = \frac{D\left(d_{max}(r), r, SSD\right)}{D\left(d_{max}(r_0), r_0, SSD\right)}$$

where $d_m ax(r)$ and $d_m ax(r_0)$ are from PDDs of the custumized cutout or the reference cone. The reference cone size of 15 cm by 15 cm is recommended with higher energy is equipped.

The PDDs are normally measured using ion chambers and diode in an automated scanning system. The

$$PDD_w(d) = PDI_w(d) \times \frac{\left[(\overline{L}/\rho)_{air}^w \times P_{repl} \right]_d}{\left[(\overline{L}/\rho)_{air}^w \times P_{repl} \right]_{dmax}}$$

14.6 Internal shielding

is useful to protect the normal structures around the high Z shaping material. For electrons in the range of 1-25 MeV, the range of the backscattered electrons is about 1-2 g/cm2 of polystyrene (see TG-70 table IV below).

Example 3.1 A buccal mucosa lesion is treated with a 9 MeV electron beam incident externally on the cheek. Assuming cheek thickness including the lesion, to be 2 cm, calculate (1) the thickness of lead required to shield oral structures beyond the cheek; (2) magnitude of electron backscatter, and (3) thickness of bolus or aluminum to absorb backscattered electrons. (1) Electron energy at depth z, $E_z = E_0(1 - z/R_p) \sim 5$ MeV, lead thickness is 5/2 = 2.5 mm. (2) For the polystyrene-lead interface, the electron backscatter factor (EBF) can be calculated as , and thus EBF = 1.57 or 57% backscattering.

²Scanning electron beams have better beam quality but suffered from the Therac 25 incident; Scanning technque is widely used in proton beam delivery

14.7 Total skin electron irradiation (TSEI)

The total skin irradiation (TSI) is one of the most efficient techniques in the treatment of the cutaneous T-cell lymphoma (mycosis fungoides). (Diamantopoulos) Its purpose is to deliver the prescribed dose (average 36 Gy over 18 fractions) to patient skin, without damaging any healthy organ. The main prerequisite for TSE installation is a linear accelerator capable of producing large (200 cm x 80 cm) and uniform fields (acceptable variation of dose distribution: \pm 8% vertically and \pm 4% horizontally within the central 160 cm x 60 cm field area according to AAPM TG-30) of relatively low energy electrons (4-10 MeV at the exit window, 3-7 MeV at patient's surface) at an extensive SSD.

Our institutional experience

Treatment:

• Dose rate: 2500 MU/min (Truebeam High Dose Electron) or 900 MU/min (Artiste)

• Energy: 6 MeV

• technique: large-field technique - 6 patient positions, and two gantry angles per position

• Schedule: 6 beam per day;

wear paper short

• Protection: Finger and toe nail shields

• Internal eye shields

- TG-51 was performed on this beam, with the machine output adjusted to 1cGy/MU at the depth of 1.3 cm deep (dmax) with 100 cm SSD with a 15 x 15 cm² cone.
- The TG-51 setup was replaced with solid water and a PTW 23343 Markus chamber. A transfer factor was established for this chamber, 0.019675 nC/cGy.

The gantry was then angled to 270° and the chamber placed a varying extended distances from the isocenter. At 330 cm SSD an acceptable dose rate was found, 59 cGy/min, without reducing field size and uniformity. Film was placed on the back side of the scatter screen at 330 cm SSD. The film was irradiated with 450 MU, with varying sets of beam angles. A $\pm 10\%$ uniformity was achieved using beam angles of 253° and 287°, over a height of 200 cm (figure 1). The patient treatment will then be 12 beams. The patient will be treated by the two gantry angles at each of 6 positions, 3 one day, 3 the next, per fraction. The patient will face the accelerator (AP beams) and be irradiated by 253° and 287° gantry angle beams. The patient will then rotate 120° (RPO), receive the two beams again, then rotate 120° (LPO) for the last two beams. The next day the patient will face away from the accelerator (PA), then rotate 120° (LAO), and again (RAO). A 1 cm thick scatter screen will be placed at 310 cm SSD. A cylindrical phantom, 30 cm in diameter was then placed at 330 cm SSD, centered with the lasers, with the scatter screen 20 cm in front of it. Powder Thermo-Luminescent Dosimeters (TLD)s were placed around the circumference of the cylinder. Additional TLDs were placed around the circumference under 5mm of wax bolus. This phantom was irradiated with the two beams, (gantry angles of 253° & 287°), and then rotated 60, 120, 180, 240 and 300° about its vertical axis, irradiated at each position with the two beams. This simulates the patient treatment. Each beam was 450 MU, 6MeV at 900 MU/min, with a 33x33cm field size. The average TLD reading was 67.6 with a standard deviation of 2.8 cGy. This gives the beam calibration factor of 67.6 cGy/450 MU per beam. The TLD value was compared to the chamber measurement from the two beams, but no phantom rotation.

This is 26.2 cGy/450 MU. This implies B factor of 2.58 (expect 2.5-3). The B factor represents the increase in dose due to the overlapping of the surface exposed at each phantom rotation and the oblique angle of incidence. The Percentage Depth Dose (PDD) was determined using film (figure 2), chamber (figure 3) and TLD measurements. The chamber measurements are only based on directly incident beams, ie the phantom is not rotated. They give a PDD at 5 mm of around 93%. The film and TLD measurement used all 12 beams (6 phantom positions, 2 beams per position), therefore the surface dose is greatly increased by the oblique

angles of incidence of the electron beams. This reduces the PDD. The film and TLDs both see a PDD of 85% at 5 mm.

PDD at 5 mm is 85%. Calibration factor: 67.6 cGy / 450 MU MU = dose per fraction / (PDD * Calibration Factor) = 200 / (0.85 * 67.6/450) = 1566

Patient treated on TSET stand. 1 cm thick plastic scatter plate 20 cm in front of patient. 3 cm thick plastic shield for lower half of body 3 mm thick lead shields on fingernails and eyes Stand against wall* (how you make sure SAD and SSD setup?)

Every patient gets same MU!

Boost fields are required at various locations: Vertex of scalp; Mid forehead; Lt. (Rt.) Axilla; Sternum; Under Lt. (Rt.) Breast; Back at T5; Umbilicus; Lt. (Rt.) Gluteal fold; Middle gluteal fold; Under scrotum (perineum); Lt. (Rt.) Anterior Thigh; Lt. (Rt.) Anterior Finger; Lt. (Rt.) Anterior toe; Calibration TLDs 6 MeV boost with 1 cm bolus.

14.8 Solutions

```
Q1: a) see section 14.3
Q2: c)
Q3: c); the energy of clinical electron beams is specified as the most probable energy at the surface
Q4: c)
Q5: b)
Q6: a)
Q7: b)
Q8: a) b)
Q9: c)
```

Q10 Virtual SSD d); Virtual electron source-surface distance is not a physical distance. It is a distance with which the inverse square law could be used for different SSDs. In reality, however, this output and pdds are measured for different SSDs instead of using the method of virtual SSD.Q11: a) b) c) d)Q12 Photon and electron beam junction a); more scattering from the electron beam will enter the side of the photon beam.Q13 b)Q14 a) c)'

Brachytherapy

Brachytherapy began at the turn of 20th century, contemporary with external beam radiotherapy. Physicsits and physicians together developed the field. There has not been a period since the beginning that has not witnessed innovations and progress in brachytherapy

— B.R. Thomadsen in "Anniversary paper: past and current issues in brachytherapy physics"

Dose rates defined in ICRU 38 (1985) and 10 CFR 35

- Ultralow dose rate: 0.01-0.3 Gy/hr (prostate implants)
- $\bullet~$ Low dose rate: 0.4-2.0 Gy/hr
- Medium dose rate: 2-12 Gy/hr
- High dose rate: >12 Gy/h; modern HDR can deliver about 430 Gy/hr
- Pulsed dose rate

Brachytherapy category depending on placement of sources

- Plaques or mold eye plaques
- Interstitial prostate implants
- Intracaitary HDR cylinder
- \bullet Intraluminal IVBT

15.1 Isotopes

15.1.1 Radium sources

Radiumsources used for implant therapy contain 226 Ra in secular equilibrium (takes about 1 month) with its decay products (226 Ra to stable lead 206 Pb).

The radium was supplied in the form of a salt, which was mixed with an inert filler such as magnesium oxide or barium sulfate. The small crystals of radium salt and filler were contained within cylindrical cells about 1 cm long. The cells were made of gold foil 0.1 to 0.2 mm thick and were sealed to prevent the escape of radion gas. Each source of radium contained 1 to 3 cells surrounded by a wall of platinum, reinforced with iridium (10%). The thickness (usually 0.5 or 1 mm) of the platinum-iridium wall was sufficient to absorb α and β radiation from the source. Gamma rays were attenuated only slightly by the wall.

The exposure rate from a 1 mCi point source of 226 Ra that is in secular equilibrium with its decay products and enclosed within a 0.5 mmPt-Ir wall is 8.25 R/hr at a distance of 1 cm. The value of 8.25 R·cm²/hr·mCi is referred to as the exposure rate constant.

15.1.2 Radium substitutes

Isotope	$T_{1/2}$ (days)	Median E (KeV)	HVL (mm lead)	usage
Cs-137	30 years	660	5.5	-
Ir-192	74	400	2.5	interstitial implantation (in ribbon)
I-125	60	28	0.025	prostate implants (23), eye plaques
Pd-103	17	22	0.01	prostate implants (23)
Cs-131	10	29	-	intracavitary BT of uterine cervix
Au-198	2.5	400	2.5	eye plaque
Y90 (β emitter)	2.67	937	-	liver radioembolism

Table 15.1: Frequently used radioactive isotopes¹

Radiation safety of brachytherapy sources

- Storage shielding container; forceps, personal dosimetry
- Test for tumor distribution of activity autoradiography using films (exposure distribution + source physicial position and shape)
- Evaluation the safety leak test (the source is swapped with a moistened cotton swab, which is then placed into a scintellation counter)

15.2 Source strength

- Radium sources has been specified in terms of mass of radium
- Cesium sources was specified in terms of milligram equivalents of radium (mg·Ra·eq)
- Sources can also be specified in terms of activity; the apparent activity, of a source is determined from a measurement of the exposure rate at a distance; it describes the activity of that nuclide that would produce the same exposure rate when unencapsulated.
- AAPM recommend air-kerma strength

15.2.1 Unit: Air-kerma strength S_k

Air-kerma strength (S_k) of brachytherapy sources is is defined as the air-kerma rate at a reference distance (e.g. 1 m) from the source center along the perpendicular bisector. The air-kerma strength is related to the quantity exposure rate by²

$$S_k = \dot{K}_{\delta} \cdot d^2 = \dot{X}(d) \cdot \left(\frac{\bar{W}}{e}\right) \cdot d^2, \tag{15.1}$$

The exposure rate $\dot{X}(d)$ is measured using an ion chamber, a "reentrant"-type well chamber, or a dose calibrator supplied with a suitable standard source. The term (\bar{W}/e) is the averagy energy to create an ion pair $(\mathbf{0.876~cGy/R})$.

The international community uses the term of reference air-kerma rate (RAKR), which is defined as the air kerma rate at 1 m from the source in μGyh^{-1}

²The equation above is a revised definition of air-kerma strength (new cutoff energy 5 keV); experimentally, an aluminum filter is put in front of NIST wide angle free-air chamber (WAFAC) to get rid of photons with lower energies.

15.3 Radiation dose from brachytherpay sources

15.3.1 From exposure rate

Early prescriptions for brachytherapy treatments were expressed in terms of radiation exposure.

The exposure rate (in R/hr) at some distance, r (cm), from a point source of radioactive material is:

$$\dot{X} = \frac{\Gamma_{\delta} A}{r^2} \tag{15.2}$$

where A is the activity of the source, and Γ_{δ} is the exposure rate constant for the nuclide. For example, Γ_{δ} is 3.28, 4.62, 1.45, and 1.48 R·cm²/hr·mCi for ¹³⁷Cs, ¹⁹²Ir, ¹²⁵I, and ¹⁰³Pd, respectively.

The resulting Sievert integral is shown

$$\dot{D}(r,\theta) = \frac{\Gamma_{\delta} A f}{lr} \int_{\theta_1}^{\theta_2} e^{-\mu t/\cos\theta} d\theta \tag{15.3}$$

where f is the f factor (exposure to dose conversion), l is the length of source.

Although the effects of photon scatter and attenuation in tissue are neglected, dose calculations can be reasonably accurate because contributions of scattered radiation to a point very nearly compensate for the tissue attenuation of radiation reaching the same point.

15.3.2 TG-43

Most brachytherapy dose calculations are now based on TG-43 and its updates³ and are done using computer programs. It should be noted that most current dose calculation still assumes that all material is **water equivalent**.

The four principal factors influencing the relative dose distribution include: 1) **Distance** - a factor of 100 between the distances of 0.5 and 5 cm while the remaining factors over the same distance range rarely exceeds a factor of 2 or 3; 2) **Attenuation and scattering in source structure** - the dose near the longitudinal axis is usually smaller than on the transverse axis; 3) **Attenuation by surrounding medium** - compton scattering and photoelectric absorption; 4) **Accumulation of scattering in surrounding medium**.

Although the manual calculation is not needed, it is helpful to understand how the dose distributions are calculated in TG-43. In TG-43, the dose rate distribution around a sealed brachytherapy source can be determined in two-dimensions (2D) using the following equation:

$$\dot{D}(r,\theta) = \Lambda S_k \frac{G_L(r,\theta)}{G_L(r=1cm,\theta=90^\circ)} g_L(r,\theta) F(r,\theta), \tag{15.4}$$

or a simplified 1D version

$$\dot{D}(r) = \Lambda S_k \frac{1}{r^2} G_p(r) \phi_{an}(r), \qquad (15.5)$$

where r denotes the distance (cm) from the center of the active source to the point of interest and θ is the point of interest relative to the source longitudinal axis.

³Other TG 43 updates include: 1) eliminating apparent activity for specification of source strength, 2) eliminating the anisotropy constant in favor of the distance dependent 1-D anisotropy function, $\phi_a n(r)$, and 3) providing guidance on extrapolating tabulated TG-43 parameters to longer and shorter distance.

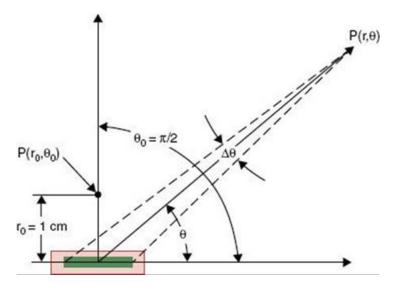


Figure 15.1: TG 43

Dose rate constant

The dose-rate constant is defined as:

$$\Lambda = \frac{\dot{D}(r_0, \theta_0)}{S_k}.\tag{15.6}$$

It can be thought as the dose rate in water at a reference point ($r_0 = 1$ cm for photon sources and 2 mm for beta emitters) along the transverse axis ($\theta = 90^{\circ}$) for source strength of 1 U.

Radial dose function and geometry function

The radial dose function g(r) represents the attenuation of radiation in tissue, defined as

$$g(r) = \frac{\dot{D}(r, 90^{\circ}) \cdot G(r_0, 90^{\circ})}{\dot{D}(r_0, 90^{\circ}) \cdot G(r, 90^{\circ})}.$$
(15.7)

where $G(r,\theta)$ is the geometry function which accounts for the effect of the distribution of radioactive material inside the source on the dose distribution at a given point. It is equal to $1/r^2$ for point source approximation, $\frac{tan^{-1}[(x+L/2)/y]-tan^{-1}[(x-L/2)/y]}{Ly}$ or $\frac{1}{x^2-(L/2)^2}$ for linear source approximation with $\theta \neq 0^\circ$ or $\theta = 0^\circ$ and x > L/2.

Anisotrpic function

 $F(r,\theta)$ is the anisotropy function, defined as

$$F(r,\theta) = \frac{\dot{D}(r,\theta) \cdot G(r,90^{\circ})}{\dot{D}(r,90^{\circ}) \cdot G(r,\theta)}.$$
(15.8)

It accounts for anisotropy of dose distribution around the source, including effects of absorption and scatter in medium, i.e., self-filtration in source, oblique filtration in walls, scattering and absorption in tissue. In TG-43U, typically calculated from Monte Carlo.

15.4. SOLUTIONS 53

		n for photon-emitting I t of the AAPM and ES			ii oco wii	ii arcias	e cherg	, mgi
/led	I. Phys. 39	(2012) 2904-2929						
	Λ =	1.113		cGy/(h U)				
	Internola	ated / extrapolated	l da	ta are hold	lface /	underli	ned V	alue
	Interpole	ited / extrapolated	ı ua	la ale bolo	ilacc /	unden	ileu. v	aruc
	$g_L(r)$			F(r,θ)				
	L=	0.35	cm		Dis	tance fr	om Acti	ve S
	r (cm)	g _L (r) L = 0,35 cm unbounded		Theta/deg	0	0.25	0.50	1.0
	0	0.991		0	0.672	0.672	0.654	0.6
	0.25	0.991	1	1	0.671	0.671	0.652	0.6
	0.50	0.997	1	2	0.669	0.669	0.651	0.6
	0.75	0.998	1	3	0.663	0.663	0.652	0.6
	1.00	1.000	1	5	0.671	0.671	0.665	0.6
	1.50	1.002	1	7	0.694	0.694	0.690	0.6
	2.00	1.004	1	10	0.735	0.735	0.731	0.7
	3.00	1.005	1	12	0.762	0.762	0.760	0.7
	4.00	1.003	1	15	0.803	0.803	0.799	0.7
	5.00	0.999		20	0.852	0.852	0.850	0.8
	6.00	0.991		25	0.892	0.892	0.887	0.8
	8.00	0.968		30	0.917	0.917	0.913	0.9
	10.00	0.935		35	0.936	0.936	0.933	0.9
				40	0.955	0.955	0.951	0.9
				45	0.964	0.964	0.962	0.9
				50	0.973	0.973	0.972	0.9
				55	0.986	0.986	0.979	0.9
				60	0.990	0.990	0.984	0.9
				65	0.993	0.993	0.989	0.9
				70	0.996	0.996	0.993	0.9

The example information about Elekta Flexisource can be seen in the figure below

In brachytherapy there is a rapid falloff in dose as distance from the source increases due to inverse square law. The dose within the tumor may much different from the prescription dose, thus the concept of equivalent uniform dose (EUD) was introduced by Dale et al. (1997). Mathematically, the generalized EUD is defined as

$$EUD = \left(\sum \nu_i D_i^a\right)^{1/a}$$

Here ν_i is the fractional organ volume receiving a dose D_i and a is a tissue-specific parameter that describes the volume effect.

- $a \to -\infty$, EUD = minimum dose;
- $a \to -\infty$, EUD = maximum dose (serial organs);
- a = 1, EUD = mean dose;
- a = 2, EUD = RMS dose.

The EUD model is parameterized by the single biological parameter a, which should be chosen so that the EUD reflects the intended biological properties for the given tumor or organ. Parameter a and the Lyman model parameter n are related by a=1/n Tumor: a is a negative number (e.g., a=-15) Normal tissues: a is a positive number

The volume-effect: very small normal tissue volumes (e.g. 1-2 cm³) can tolerate very high doses that larger volumes would not tolerate. There are a few exceptions to this such as spinal cord, though the dose as high as 167.3 Gy to the cord has been reported in very low dose rate brachytherapy of paraspinal tumor. Rogers et al. (2002) reported that the mean cord dose was 72.5 Gy (ranging: 53.1-167.3 Gy), combining the EBRT and I-125 brachytherapy.

15.4 Solutions

Q1 d)

As
$$D = \dot{D} \times \Delta t$$
, $\frac{\Delta t_{new}}{\Delta t_{old}} = \frac{\dot{D}_{old}}{\dot{D}_{new}} = \frac{A_{old}}{A_{new}} = \frac{A_0 e^{-10/30}}{A_0} = 0.79$

Q2 Shielding b)

Although the average energy of ⁶⁰Co is higher than that of ²²⁶Ra, there are gamma rays of 1.76 and 2.2

MeV emitted from 226 Ra sources. In shielding design, we need to consider their existence (although their contribution is small) and thus HVL for 226 Ra is greater than HVL for 60 Co. 4

Q3 a) b) c) A ¹³⁷Cs source is normally used for consistence check (like linac monthly QA) but not calibration.

Q4 a) Like external beam radiotherapy, the inverse square law is always the biggest factor for dose calculation.

Q5 a) Should c) and d) be correct?

Q6 Initial dose rate c)

The prescription dose or total dose for an prostate implant is

$$D = \int_0^\infty \dot{D}_0 \cdot e^{-\frac{0.693}{T_{1/2}}t} dt$$

Using an important definite integral, $\int_0^\infty e^{-ax} dx = \frac{1}{a}$, we can find that

$$D = \dot{D}_0 \cdot \frac{T_{1/2}}{0.693} \rightarrow \dot{D}_0 = \frac{D}{59.4/0.693} = \frac{14400 \text{ cGy} \times 0.693}{59.4 \text{ days} \times 24 \text{ hours/day}} = \boxed{7.0 \text{ cGy/hr}}$$

Q7 b)

Q8 The Paterson-Parker system c)

Q9 The Quimby system a)

Q10 The Paris system c) d)

The air kerma strength and apparent activity conversion is 1 U = 0.348, 0.243, 0.486, 0.787, and 0.773 mCi for 137 Cs, 192 Ir, 198 Au, 125 I, and 103 Pd, respectively.

15.5 Traceability

Calibrations of brachytherapy sources should be directly traced to NIST or to an Accredited Dosimetry Calibration Laboratory (ADCL) which is traced to NIST. Normally, we don't send sources to NIST or ADCL, but instead a well chamber with specific inserts designed for different isotopes. To calibrate Bard PS-1251L I-125 sources, for instance, the well chamber with an I-125 insert will be used, which was checked using Bard PS-1251L I-125 sources at NIST or ADLC.

TG-40

- all long half-life sources should be calibrated;
- at least 10% or 2 ribbons (whichever is larger) should be calibrated for a large number of loose seeds with **short** half-life.

If the institution's verification of source strength disagrees with the manufacturer's data by more than 3%, the source of the disagreement should be investigated. We further recommend that an unresolved disparity exceeding 5% should be reported to the manufacturer.

Radioactive materials must be under control by the facility at all times. This means under direct control or by securing in a locked area.

 $^{^{4} \}rm https://www.nrc.gov/docs/ML1122/ML11229A721.pdf$

Radiation Protection

Historically, the most commonly used unit in US is millirem (mrem) where rem stands for *Roentgen Equivalent Man*. The SI unit of effective dose and equivalent dose is *Sievert* (Sv). Because 1 Sv, equal to 1 Gy numerically, is rather large quantity, the milliSievert (mSv) is commonly used in practice. The relationship between mSv and mrem is

1 mSv = 100 mrem

.

16.1 Sources of radiation exposure

According to the National Council on Radiation Protection and Measurement (NCRP) report 160 (2009), the average annual radiation dose per person in the U.S. is about 6.2 mSv, in which medical imaging contributes about 50% (e.g. CT: 24%, NM: 12%, interventional fluoroscopy 7%, conventional radiography 5%). Naturally occurring sources of radiation include cosmic radiation (5%), radioactive minerals in the ground and in your body (5%), and terrestrial radiation emitted by naturally occurring materials such as uranium, thorium, and radon (37%) in earth. The pie chart of sources of radiation exposure from NCRP 160 can be found here.

jdkhfakjdhfkljadsfdf

16.2 Stochastic and deterministic event

Although the severity of the stochastic effect is independent of the dose, the probability of having such effects is proportional to the dose **without dose threshold**. The examples of stochastic effects include radiation induced cancer and genetic mutation. Skin erythema, epilation (hair loss), lens opacification, and tissue necrosis are best described as non-stochastic or **deterministic** events. For deterministic effects, there is a threshold and the severity of the effect depends on the dose.

To avoid unacceptable complications, normal tissue should be below a **tolerance dose** (TD) (Emami et al.) Complications is categorized as fatal, severe (e.g. grade 3-4 pneumonitis), and quality-of-life complications. TD5%/5 and TD50%/5 are used to imply complications in 5 years.

Answer: e)

16.3 TDS rule

Time $(D \propto \dot{D} \times \Delta t)$, distance (inverse square law), and shield (attenuation) measures are major factors in consideration of minimizing the unavoidable radiation exposure. Other procedures to minimize the exposure are containment and NRC's system for radiation protection according to NRC guidelines. The NRC's system for protection includes (1) dose limits for radiation workers and members of the public; (2) monitoring and labeling radioactive materials; (3) posting signs in and around radiation areas; and (4) reporting the theft or loss of radioactive material. In addition, the NRC imposes penalties for failures to follow the agency's regulations.

If the licensees' can limit the radiation to 1 mSv to the public and 50 mSv to adult radiation works in a year, the NRC may enter into an agreement with a State governor to give the State authority for regulating radioactive materials. States that meet these conditions and agree to regulate materials using the same standards as the NRC are called **Agreement States**.

Answer: d)

$\mathbf{Q}\mathbf{A}$

Uncertainties in Radiation Medicine: An Oncologist's Perspective

CTV margin for subclinical tumor - ML

Medical physics practice guideline 4.a

17.1 Patient-specific QA

At CC, we do PSQA using 2D ion-chamber array and EPID.

Question: compared to film/chamber, what are the limitations of EPID-based QA?

• not true composite (TC)?

.

What are the action limit and tolerance limit at CC?

For EPID-based QA, how do you make sure the physical properties (sag, position, ...) of imager are still in tolerance?

• IsoCal Calibration

17.2 Quality and safety

AAPM quality and safety sources

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TBI

TG-17 (1986) "The physical aspects of total and half body photon irradiation"

The reported D_0_ value - the amount of ionizing radiation necessary to eradicate a particular cell type—of hematopoietic stem cells is 0.5 to 1.4 Gy, while those of human leukemia cell lines are 0.8 to 1.5 Gy, indicating that both cells are radiosensitive.

Fractionated TBI has been shown to lead to a higher incidence of graft rejection than the same dose delivered in a single fraction, possibly due to DNA repair during interfraction intervals.4,7,12 However, fractionation decreases the eradication of bone marrow stromal cells, which are necessary for successful hematopoietic stem cell engraftment, and is, therefore, considered the standard of treatment

 $https://appliedradiationoncology.com/articles/total-body-irradiation-a-practical-review \\ https://www.ted.com/talks/daniel_kraft_invents_a_better_way_to_harvest_bone_marrow/transcript why TMR is SAD independent http://www.npl.co.uk/upload/pdf/20140513-dart-pres-byrne.pdf$

If only high-energy photons are available and superficial structures would be underdosed, spoilers may be used. The ideal is to maintain a low skin dose and increase dose in the build-up region, to emulate a lower energy beam. However, while it is impossible to exactly mimic a lower energy beam with a spoiler, the build-up characteristics may be preferable to using bolus.

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Three-dimensional conformal radiotherapy

19.1 ICRU reference point

The ICRU reference point is the point in the center (or center parts) of the PTV

19.2 Image registration

- Brady: Geometric (and Photometric) alignment of one image with another Images may be of same or different types (MR, CT, and etc.)
- ITK: The process of determining the spatial transformation that maps points from one image to homologous points on an object in the second image.
- Elastix: The task of finding a spatial one-to-one mapping from voxels in one image to voxels in the other image.

19.3 Image segmentation

http://www.cs.uu.nl/docs/vakken/ibv/reader/chapter10.pdf: the division of an image into meaning structures. Wiki: image segmentation is the process of partitioning a digital image into multiple segments (sets of pixels) Khan: slice-by-slice delineation of targets and organs-at-risk

19.4 Cumulative DVH

DVH See Chapter 11 Q6

19.5 Differential DVH

The choice of c): a certain dose within a specified dose interval as a function of dose. The differential DVH is similar to conventional histogram in statistics.

IMRT

IMRT provides an ability to deliver many beamlets (smallest element to be modified) of varying radiation density within one treatment field.

20.1 IMRT

The number of photons was modulated by **blocking the photon beams** (fluence) at specific location and/or time with MLC or **changing the dose rate**¹.

20.2 IMRT QA

AAPM TG-218

20.3 Transmission or leakage

For current machines, In IMRT, the relative contribution to the target dose from collimator transmission scatter is greatest for: a) leaf transmission; b) round edge transmission; c) X-ray jaws; d) overall head scatter Intra- and inter-leaf transmission: (Varian manual) Average intra-leaf and maximum interleaf leakage for the Varian HD-120 MLC is and 2.0% and 2.5% (up to 10 MV). Based on Bedford et al. (2013), the maximum intra- and inter-leaf leakage for the Elekta Agility MLC (9 cm height) is 0.5% and 0.2%. Leaf (round) end transmission is not reported anymore, and should be in the range of 10%-20%. Jaw transmission is about 1% and 1.5% for the Edge and Versa machine, respectively.

Q1: b), c)

20.4 Q4 MU: IMRT vs. 3DCRT

Compared to the four-field box technique, an IMRT plan could require *substantially* more monitor units (MU). The MUs of an IMRT plan largely depend on the degree of dose modulation within a target and/or a

¹It is fancinating that how dose rates are changed in linacs. A 2013 PMB paper, Radiobiological effects of altering dose rate in filter-free photon beams, showed that altering radiotherapy dose rate through either changing pulse repetition frequency or instantaneous dose rate does **not** have an effect on cell survival. An increase in survival was seen in both modes upon protracting dose delivery to 15, 30 or 60 min rather than delivering acutely. Is this important to PLDR? Should we increase the prescription dose?

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proximity between a target and nearby OARs. With the improvement in optimization algorithm and electromechanical performance of linac, the difference of MUs between an IMRT plan (especially using VMAT technique) and 3D-CRT plan has decreased.

Q2: c

20.5 Q5

In generating an intensity-modulated profile in minimum time with the dynamic MLC: a) the opposing pair of leaves should move with equal but variable speed; b)the leading leaf should move at the maximum speed and trailing leaf should provide the required intensity modulation, if the gradient of the intensity profile is positive (increasing fluence); c) the trailing leaf should move at the maximum speed and trailing leaf should provide the required intensity modulation, if the gradient of the intensity profile is negative (increasing fluence); d) the two leaves should move with equal and maximum speed, if the spatial gradient of the intensity profile is zero.

Q5: b), c), d)

20.6 Shielding for IMRT

If majority of the patients are to be treated with IMRT instead of conventional radiation therapy, the total MUs will be largely increased despite delivered dose remains the same. Therefore, the major concerns would be the increased leakage radiation so is the design of the secondary barrier. Solution: c and d

20.7

The difference between an IMRT and 3-D CRT delivery typically include: a) Non-uniform (modulated) beam intensities; b) Patient-specific beam-shaping c) Inverse planning for dose optimization; d) Dosimetric or biological objectives with relative weights; e) Significantly more complex dose calculation algorithm Solution: a c d

20.8 Q8

IMRT delivery technique include:
IMAT
Conformal arc therapy
Helical tomotherapy
DMLC delivery
SMLC delivery

Solution: e

20.9 Q9

The term step-and-shoot is sometimes used to describe which IMRT delivery technique: Helical tomotherapy Serial tomotherapy IMAT Segmental MLC-IMRT Dynamic MLC-IMRT

Q9: d)

20.10. Q10 65

20.10 Q10

For a step-and-shoot IMRT treatment delivery, an MLC controller system introduces 50 millisecond delay between the monitor chamber signal reach a control point and beam termination. If the initial segment of a field is set to receive 2 MU, what percent error does this delay introduce for this segment if the linac's output is set to 600 MU/min? $<1.5 \times 10.25 \times 250$

Q10: d)

20.11 MLC test(s)

Which MLC test(s) are unique to dynamic MLC delivery? Linac performance for small MU delivery Leaf positional accuracy Inter- and intra-leaf leakage Tough-and-groove effect Leaf speed accuracy Solution: e

Generally speaking, the MLC delivery can be categorized into two types: static (e.g., step-and-shoot) and dynamic (e.g., vmat and conformal-arc). Linac performance on small MU delivery has been a serious issue for static MLC delivery. Xia et al. (2002) has used a simple formula to relate the dose error (Δ) with dose rate (R), communication time (T), and MU/segment (M): $\Delta = RT/M$ For example, if dose rate is 600 MU/min, T = 100 mS, and M = 1 MU/seg, the dose error 1 or 100%. Therefore, larger dose errors are expected for smaller MU segments with certain dose rate and communication time. Recent progress in increase of sampling rate (e.g., from 100 ms to 20 ms) and integration of MLC controller with the linac have significantly improved the dose delivery accuracy for step-and-shoot IMRT Li et al. (2012). In addition, as the optimization algorithms improved, the use of increased minimal MU per segment (> 4-8) further reduce the dose errors caused by smaller MU segments used in the plan. Leaf positioning error impact also

20.12 Q12

The contribution of MLC leakage to the total dose from an IMRT field: Is the largest contribution of the dose Increases with increase in leaf speed Increases with increase in leaf gap width May be neglected in the final dose calculation None of the above

Solution: e

More Inverse planning procedures Clinical objectives (goals) are specified first in terms of desired (physical or biological) dose or DVH goals. Field's fluence map (the set of beamlet weights) is optimized The optimize fluence is converted to deliverable MLC positions and further optimization is continued It should be noted that the step 2 and 3 are integrated into the direct machine parameter optimization (DMPO). Consideration of beam number and placement (shortest path to irradiate targets and avoid OARs) Complexity of the target shape Proximity to critical organs Collimator angle (minimize leakage and maximize coverage)? Previous RT? Non-coplanar beams? Parallel opposed beams?

idea EPID triggered imaging with mlc position?

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SBRT

The great difficulty in the world is not for people to accept new ideas, but to make them forget about old ideas. -John Maynard Keynes

21.1 Milestones

- 1908 Horsley and Clarke coined the term stereotaxis and fabricated an apparatus that can be rigidly clamped to the skull;
- 1947 Spiegel and Wycis frame¹;
- 1949 Talairach defined anterior commissure posterior commissure (AC-PC) line and a brain atlas;
- 1951 Lars Leksell developed a frame exclusively for human beings;²
- 1967 First SRS treatment on a Gamma Knife (GK) machine.
- 1991 Lax and Blomgren "Extracranial sterotactic radiation therapy" at Karolinska.
- 1999 Adler et al. IGRT-SRS on a Cyberknife (CK) machine.
- 2003 Timmerman Phase-I trial on lung cancer at Indiana U.
- 2005 SBRT CPT code added.
- 2018 ZAP system

The SRS treatments have been delivered using GK, CK, tomotherapy machine, and linear accelerators (Lutz et al. (1987))³.

21.2 The definition of SRS and SBRT

ACR/ASTRO definition of SBRT: an external beam radiation therapy method used to very precisely deliver a high dose of radiation to an extracranial target within the body, using either a single dose or a small number of fractions.

¹Pneumoencephalography is an painful procedure that requiring a spinal tap so that pressured air could be introduced into the cerebrospinal fluid space. This created two large air bubbles filling the patient's ventricles so that they could be imaged radiographically. (Interesting) the patient was suspended from the ceiling by a full body harness, which was rotated in 3D to assist the introduction of air into the brain. - Steven J. Goetsch "Historic developement of SRS and SBRT" in Stereotactic Radiosurgery and Stereotactic Body Radiation Therapy

²The location of the desired target was determined from radiographic procedure, and then translated to Leksell coordinates. The frame is still used for Gamma knife.

 $^{^3}$ A system for stereotactic radiosurgery with a linear accelerator. Extensive performance tests have shown that a target, localized by CT, can be irradiated with a positional accuracy of **2.4 mm** in any direction with 95% confidence. This number has not been decreased much in last 30 years. The geometric accuracy of isocenter localization of ± 1 mm is acceptable.

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To simulate the head frame used in the SRS, a body frame was used in early SBRT treatments (Sweden and Japan) in early 1990s. With advances of technologies, however, most current SBRT treatments do not use body frames, because the localization accuracy is comparable between frame-based and frameless SBRT and worse than that in the SRS.

21.3 Features of GAMMA Knife PerfexionTM and IconTM

9/13/18 from GK manual

As part of the imaging process, it is essential to provide exact points of reference by means of which the shape and position of the targets can be ascertained with respect to the patient's skull. Moreover, during the subsequent radiosurgery session, the head of the patient must be entirely immobilized to maintain the accuracy of the shots.

21.3.1 Stereotactic reference

- use an indicator box during image acquisition
- use the CBCT (only available for Icon)

21.3.2 Steps to create new plan

- Creat a new patient file
 - Have to complete the Radiological Examination field
- Plan | New Plan to make a new treatment plan
- Open a patient file Tomographic imaging acq

Leksell Coordinate System

21.4 GK Troubleshooting

- When doing CBCT verification, the Frame docking did not work
 - Ask engineering to fix the sensor which reads whether frame/mark dock is in position (Docked) and which frame is in position (Docking).
- The frame (one screw) became loose, what should be do?
 - Tighten the screw, do an CBCT vertication scan
 - Switch to mask case, you need to change fixation from frame to mask, and then do Request CBCT - Stand.
- If there are "out of range" warning messages when putting shots, what can you?
 - to add measurement information and change gamma angle from 90° to 70°.

Leksell Coordinate Frame G Gamma Knife Perfexion and

- The patient positioning system (PPS) is the treatment couch
- A frame adapter attaches the Leksell coordinate frame (known geometry) to the treatment couch (known relative position to radiation isocenter)
- Built-in collimation system, and 3 collimator size: 16, 8, and 4 mm
- 192 Co-60 sources; (the model U, B, and C, still in use at some centers, have 201 sources.)
- Souces are not fixed in space; they reside on 8 independent sectors
- Each sector has 5 positions (4 mm, blocked, 8 mm, 16 mm, ?)

21.5. QA OF GK 69

21.5 QA of GK

Paula Patti: QA for the Leksell Gamma Knife PerfexionTM

Basic tests and measurements (following NRC licensing guidelines 10 CFR 35.1000)

1. Coincidence of the **mechanical isocenter of the PPS** with radiation-focal point (or radiation isocenter, or unit center point)

- 2. Agreement of measured beam profiles with GK calculation for all collimator sizes in XY, YZ, and XZ planes
- 3. Measurement of the absolute dose rate calibration for largest collimator
- 4. Confirmation of the relative output factors for smaller collimators

21.5.1 Prescision and accuracy

GK (radiation) isocenter: the center of the smallest sphere through which all beam axes pass as the radiological Unit Center Point (UCP) or isocenter. The radius of this sphere may then be seen as a measure of the spread of the beam axes or the uncertainty of their location. This uncertainty is called the precision of the Gamma Knife. [Arndt GK Dosimetry and treatment planning]

GK mechanical isocenter:

The mechanical accuracy of Gamma Knife radiosurgery based on single-isocenter measurement has been established to within 0.3 mm.

The following radiophysical data is pre-stored in Gamma Plan:

- 4 beam profiles (OARs), one for each beam sizes measured at **400 mm** distance from source center and at 80 mm depth in polystyrene.
- One data set to analytically calculate Percentage Depth Dose (PDD)
- 4 measured output factors, one for each beam size

21.6 Commissioning of GK

The ion chamber is widely used for the measurement of the depth dose in conventional photon beams. The correction for the depth dose measurement depends on the ratio of the mass attenuation coefficient of the detector material and water, $(\mu/\rho)_{water}^{detector}$. In another word, if the detector is near tissue (water) equivalent, i.e., near equivalent Z number, there is no necessity for depth correction. Although silicon diodes and radiographic films typically need corrections, the diamond diodes (unlike silicon diodes) and radiochromic films are both near tissue-equivalent.

21.7 Preparation of GK Treatment Planning

21.7.1 Frame Application: model C

The Leksell frame runs from 40 mm to 160 mm in the X axis using the automatic positioning system (APS), runs from 25 mm to 175 mm in the Y axis, and the Z axis values are not marked on the frame but are calculated by the treatment planning software from fiducial markers.

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21.7.2 Frame application: Perfexion

With the Perfexion model a new instrument has been introduced called the **frame cap**. Special care is needed if a lesion is very posterior or very anterior.

21.8 GK Plan Indices

21.8.1 Systematic Errors

Based on the rule for sums and differences in error propagation,

$$\varepsilon_{total} = \sqrt{\varepsilon_x^2 + \varepsilon_y^2 + \varepsilon_z^2}$$

21.9 Linac-based SRS

21.9.1 Cone-based

For linac SRS systems, the fields are mostly shaped by tertiary cone collimation system. The tertiary cones are precisely machined, closer to patient (smaller geometric penumbra), and diverging beam shaping further minimizes penumbra (see Yenice (2011) AAPM presentation, page2).

- Smith et al. (1993) Role of Tertiary Collimation for Linac-Based Radiosurgery. They found that the geometrical penumbra (how they separate dosimetric and geometrical penumbra?) of tertiary cone for **2 mm focal spot** in only **0.6 mm**, which is much smaller than 5.1 mm and 3.3 mm from the upper and lower jaw.
- Novotny et al. (2008) Dosimetric comparison between the GK Perfexion and 4C. They found good agreement between dosimetric parameters of those two models for 4- and 8-mm collimators.
- Wen et al. (2015) Characteristics of a novel treatment system for linac-based SRS. They found that the penumbra is about 1.2-1.8 mm for 6FFF and 2.3-5.1 mm for 10 FFF beams (80%-20%).

21.10 Diesease sites treatment with SRS

Commonly treated tumors using the SRS technique include:

- 21.10.1 Acoustic neuoromas
- 21.10.2 Arteriovenous malforamtions (AVM)
- 21.10.3 Brain metastases
- 21.10.4 Malignant gliomas

21.10.5 Menningiomas

Keeping the margin dose > 12 Gy seems to be effective

?? Much of the current assessment of control rates is based on Kaplan Meier statistics rather than raw data. In the future there will be more raw data to give a more reliable assessment.

21.10.6 Pituitary tumors

21.10.7 Unilateral Vestibular Schwannomas

Using 12-13 Gy with a noticeable improvement in the rate of complications, particular hearing loss (pay attention to cochlear nerve) and facial .

21.10.8 Uveal melanomas

One of the odd features of this tumours is its well known ability to prove deadly from systemic metastases, years after an enucleation of an affected eye; metastases usually in the liver. Where the metastases reside in the interval is not known.

Interesting treatment technique: fix the eye motion; check the positions in the dose plan against physical measurements using polymer gel with MR images.

21.10.9 Trigeminal neuralgia

The recommended doses can be found here. The dose typically depends on the tumor size (the smaller the tumor, the higher the prescription dose).

a) wrong; c) wrong; Rhabdomyosarcoma is mostly treated with linac.

21.11 Dose fall-off

Currently higher X-ray, γ -ray, and protons are used to treat SRS/SRT.

a); d)

SRS treatment are characterized by steep dose gradients, for example, >50%/cm, at the target periphery. If the spatial accuracy of the treatment delivery is \pm 1 mm, the dosimetric uncertainty in this region will be >50%/cm times 1 mm, which equals to >5%/cm.

d)

b); c)

21.12 Required measurements for commissioning a SRS/SBRT program

The measurements required for commissioning a SRS/SRT program are similar to conventional external beam radiotherapy with the exception of transmission measurement which is typically very small due to the construction of the cone collimator.

```
a); c); d)
```

b) is wrong as the average energy from Co-60 sources is 1.25~MeV not 6~MV; c) is wrong because sources move in the translational mode; d) is wrong because the Gamma knife is more accurate than a linac-based SRS system.

Mindermann (2015) Gamma Knife, CyberKnife or micro-multileaf collimator LINAC for intracranial radiosurgery?, is a good read. The author questioned a few dosimetric studies in comparing several delivery systems, and stated that the questions of dosimetry needs to be answered, which include: the source of the 72 CHAPTER 21. SBRT

photon radiation (cobalt-60 or linear accelerator); the nature of the collimators (fixed aperture, iris, micromultileaf, etc.); moving or stationary radiation sources during beam-off time or beam-on time; the fixation of the head; the planning software; the imagery used; the way the images are acquired (dedicated protocols, head fixation, etc.); the number of beams; the number of arrival angles; the exit dose; the scatter factor of a given beam; the distance source to target; the time period over which a dose is delivered; the system's overall accuracy; dose rates; the nature and the size of the lesion; the shape of the lesion; its proximity to organs at risk; the experience and the neurosurgical and anatomical knowhow of the radiosurgeon.

SBRT prostate using spacer spaceoar.com

HDR.

22.1 HDR vs. LDR

LDR: well-established treatment; standard doses, plan, and treatment time

HDR: Outpatient treatment, short administration time, minimal staff exposure, standard source strength, and dose optimization

 ${\bf Good\ read:\ Current\ controversies\ in\ high-dose-rate\ versus\ low-dose-rate\ brachytherapy\ for\ cervical\ cancer.}$ Showalter 2014 AAPM

22.2 Common indications in practice

- **GYN** (cervical¹, uterine, vaginal, vulvar)
- Prostate (monotherapy or boost)
- Breast (accelerated partial breast irradiation)
- possible Sarcoma, skin, esophagus, and bile duct

?? Theoretically, HDR has a lower therapeutic ratio than LDR because of the short duration of the treatments. How? - Practical ROP chapter "Intracavitary Brachytherapy"

22.3 HDR-QA

22.3.1 Daily QA

According 10 CFR35.643, the AMP needs to review the daily QA within 15 days.

22.3.2 Pretreatment QA (TG-59)

- 1. Two people (therapists?) should check proper **connection of catheters** to the HDR unit and that the transfer tubes are free of kinks.
- 2. The emergency kit and source container are available.

¹Cochrane review and its update: there is no difference in OS, DSS, LC, nodal occurrence, distance occurrence was found between LDR and HDR (from a meta-analysis of 4 clinical trials in *Cochrane database* with a total of 1265 patients with advanced cervical cancer), but HDR is more convenient and accurate.

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3. Survey meter and/or GM-counter is present and operational. (The patient may have had a nuclear medicine scan prior to the treatment, causing an elevated reading. thus a **pre-treatment survey** is conducted though not listed in TG-59).

- 4. The **length** of transfer tube and applicator (catheters) are correct.
- 5. Check applicator positioning. How do physicians check this item without image verification?
- 6. Treatment documentation review.
 - a. Signed prescription and plan.
 - b. Second check has been performed. (use emipircal values)
 - c. Plan agrees with prescription.
 - d. Plan is consistent with previous fractions if applicable.
 - e. Dwell positions and times in plan agree with what is programmed on the treatment console.
- 7. Patient identity confirmed by two methods.

At current practice, a **check-list** is used by physicists for pretrement plan QA and a time-out is conducted prior to initiating the treatment.

22.3.3 Source change

The half-life time is about 74 days, so the old source is sawpped with a new source about every 3 months. The activity of the new source is normally about 10 Ci. According to Eq. (15.1) and Eq. (15.2), it is equal to 41100 U $(S_k = 10,000 \ (mCi) \times 4.69 \ \left(\frac{R \cdot cm^2}{mCi \cdot hr}\right) \times 0.876 \ \left(\frac{cGy}{R}\right)$). This quantity will be verified by an autheried medical physicist (AMP) using NIST tracable well chamber and electrometer, and then enterred in the treatment planning system for dose calculation. The engineering from HDR afterloader vendor also verifies the source using their own equipment.

- 1. Verify the source cable **positioning accuracy** at two different programmed positions (1205 mm and 1400 mm) before the vendor engineering leaves (using GYN transfer tube).
- 2. Although the well chamber and electrometer is still within 2-year calibration period, we always do **consistence check** using a NIST-traceble Cs-137 source (we actually checked with 2 Cs-137 sources provided by our RSO).
- 3. Switch a physics QA transfer tube and insert a catheter into Ir-192 insert.
- 4. Measure current at 5 positions (1195 mm, 1200 mm, 1205 mm, 1210 mm, and 1215 mm) and take an average
- 5. Check time-dose linearity
- 6. Check stopwatch accuracy (100 s)
- 7. Check transfer tube connection error
- 8. Switch emergency power switch
- 9. Check

22.4 Medical Events

- Errors on NRC website
- Wisconsin

22.5 Source

A comprehensive seed data source can be found from a database provided by Carleton University

Because Ir-192 has much higher special activity than most other isotopes, it is now the mostly used radio-isotope for HDR treatment. The higher the special activity means that the Ir-192 can be made with small physial dimension but still provide high radioactivity.

The special activity (SA) is defined as the activity per mass. It depends on half lifetime and atomic number, $SA \propto \frac{1}{T_{1/2} \cdot A}$. For example, $\frac{SA_{Co}}{SA_{Ir}} = \frac{74~days \times 192}{5~years \times 60} \approx 0.13$. Wait a second, how about SA of I-125? Although I-125 can have higher SA than Ir-192, the energy of I-125 is just too low for enough tissue penetration.

22.6 Treatment sites

22.6.1 Endometrial cancer

ABS consensus guidelines for adjuvant vaginal cuff brachytherapy after hysterectomy

- Dose fractionation: 7Gy × 3 prescribed to 0.5 cm is a common fractionation scheme with active length of 5 cm (Are we treating vaginal cuff or the whole vigina?)
- the standard applicator is a segmented cylinder with one central catheter; the **largest diameter** cylinder that patient can tolerate is used to minimize the air gap between cylinder and vagina and to avoid rapid dose fall-off.

22.6.2 Cervical cancer

- ABS consensus guidelines for locally advanced carcinoma of the cervix. Part I: General principles
- ABS consensus guidelines for locally advanced carcinoma of the cervix. Part II: High-dose-rate brachytherapy

Cervical cancer is mostly treated with HDR brachytherapy.

- 1903 Stockholm and Paris
- 1938 Manchester point A
- 1953 Point A revision
- 1985 ICRU 38
- 1987 more point A updates
- 2000 GEC-ESTRO
 - D90, D100 for dose prescription
 - D2cc bladder, rectum, and sigmoid
- 2004 GTV and CTV delineation (MRI)
- 2005 GEC-ESTRO recommendation for IGRT brachytherapy

Improvement occurred only in tumours >5 cm: OS 28% versus 58% (p = 0.003) - R. Potter (2007) in "Clinical impact of MRI assisted dose volume adaptation and dose escalation in brachytherapy of locally advanced cervix cancer".

GEC-ESTRO target volumes²

- Gross tumor volume (diagnosis) (GTV_D)
 - macroscopic tumor extension at diagnosis
 - detected by clinical examination and as visualized on MRI (high signal intensity mass(es) at fast spine echo (FSE) sequences T2 in cervix/corpus, parametria, vagina, bladder, and rectum)
- Gross tumor volume (brachy) $(GTV_{B1},\,GTV_{B2},\,...)$
 - macroscopic tumor volume at time of brachy
 - detected by clinical examination and as visualized on MRI
- High risk CTV (HR CTV_{B1} , HR CTV_{B2} , ...)
 - includes GTV_{Bx} and the whole cervix or MRI grey zones?
 - represent **macroscopic** tumor load
- Intermediate risk CTV (IR CTV_{B1} , IR CTV_{B2} , ...)

²Schwarz 2015 AAPM Spring Clinical Meeting "Defining Targets for Brachytherapy" https://www.aapm.org/education/vl/vl.asp?id=4077

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- areas with a significant microscopic disease
- IR CTV = HR CTV + 5-15 mm margin for limited diseases
- based on GTV_D for extensive disease

Applicator imaging

Idea: availability of commercial dummy sources for MRI is limited.

Based on GEC-ESTRO recommendations³, the choice of MR sequence is essential for optimal visualisation of the applicator. There are difference between plastic and titanium applicators⁴

- Plastic has weak signal on T2; use of markers
- Titanium has (induced) susceptibility artifact, and thus more distortions for higher magnetic strength; worse on T2; T1 is more suitable (? Why Titanium is MR compatible?)
- If an applicator has been shown to be MR conditional for a 1.5T MRI, then it does **not** mean that it can be safely used in a 3T system without the need for further testing. CT still provides best imaging for applicator in terms of spatial accuracy (1 mm on CT vs. 1-2 mm on MRI for the localization of first dwell position) and artifacts.

Prescription Dose

- HDR prescription: $5.5 \text{ Gy} \times 5$, $6 \text{ Gy} \times 5$, or $7 \text{ Gy} \times 4$; once a week
- HR CTV: total dose > 85 Gy Can we go higher? or fewer fractions
- IR CTV: 60 Gy

22.6.3 Breast

ABS acceptability criteria for APBI

- Age: ≥ 50 year old
- Size: ≤ 3 cm
- Histology: All invasive subtypes and DCIS
- Estrogen receptor: +/-
- Surgical margin: -
- Lymphovasucular space invasion: not present
- Nodal status: -

Treatment planning

- 34 Gy in 10 fractions twice daily
- $PTV_{Eval} + D90\% >= 90\% + V150 < 50 \text{ cm}^3 + V200 < 10 \text{ cm}^3 + Skin dose < 145\% of prescription$

They are slightly different from ASTRO Consensus Statement 2009.

22.6.4 Prostate

ABS consensus guidelines for high-dose-rate prostate brachytherapy

Monotherapy: $13.5 \text{ Gy} \times 2 \text{ fractions (NCCN)}$

³Recommendations from Gynaecological (GYN) GEC-ESTRO Working Group: Considerations and pitfalls in commissioning and applicator reconstruction in 3D image-based treatment planning of cervix cancer brachytherapy

⁴Haack et al 2009 Applicator reconstruction in MRI 3D image-based dose planning of brachytherapy for cervical cancer. https://doi.org/10.1016/j.radonc.2008.09.002

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22.8 Solutions

Q1 Dose rate c)

Q2 a)

Q3 TG43U d)

Using Eq. @ref(eq.tg43) or TG-43U1 2D Brachytherapy dosimetry formalism,

$$\dot{D}(r,\theta) = \Lambda \cdot S_k \frac{G_L(r,\theta)}{G_L(r = 1cm, \theta = 90^o)} \cdot g_L(r,\theta) \cdot F(r,\theta)
= 1.12 \ cGy/(h \cdot U) \cdot 4.11 \times 10^4 \text{U} \cdot 1.023 \cdot 1
= 13.1 \ cGy/s$$
(22.1)

Q4 Afterloader QA a)

Q5 Shiedling b)

Q6 Impact of decay on treatment timee

The half-life time of Ir-192 is about 74 days, so activity after 90 days (Eq. ((2.4))) is

$$A_2 = A_0 2^{-t/T_{1/2}} = A_0 2^{-90/74} = 0.43 A_1$$

To maintain the prescribed dose $(\dot{D}_1 \Delta t_1 = \dot{D}_2 \Delta t_2 \text{ and } A \propto \dot{D}$, the dwell time Δt_2 will be

$$\Delta t_2 = \frac{\dot{D}_1}{\dot{D}_2} \Delta t_1 = \frac{\dot{A}_1}{\dot{A}_2} \Delta t_1 = \frac{1}{0.43} \times 16 \text{ min } \times 80\% = \boxed{29.7 \text{ min}}$$

The total treatment time will be $29.7 + 16 \times 20\% = 33$ minutes.

Q7 c)

 $\mathbf{Q8} \stackrel{\frown}{\mathrm{c}}$

 $\mathbf{Q9}\ \mathrm{b)}\ \mathbf{Q10}\ \mathrm{a)}$ but esophagus cancer is also treated with HDR but with less indication'

Q11 b)

Q12 d)

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Implants

23.1 Isotopes

good reference (https://aapm.org/meetings/amos2/pdf/42-11873-3201-79.pdf)

Isotope	$T_{1/2}$ (days)	Median E (KeV)	90% dose delivered (days)	Rx (Gy)
I-125	60	28	204	145
Pd-103	17	22	58	120 or 125
Cs-131	10	29	33	115
Y90	2.67	937	11	120 - 150

$$\dot{D}(r,\theta) = \Lambda S_k \frac{G(r,\theta)}{G(1,\pi/2)} g(r) F(r,\theta)$$
(23.1)

where

- $\dot{D}(r,\theta)$ is the dose rate at point P in a medium
- Λ is the dose rate constant
- S_k is the air kerma strength of the source
- ullet G is the geometry factor
- g is the radial dose function
- F is the anisotropy function

23.2 Patient Release¹

NRC-NUREG-1556 in Table U.1

Isotope	Activity threshold (GBq)	Activity threshold (mCi)	Dose rate at 1 m (mSv/hr)	Dose rate at 1 m (mrem/hr)
I-125 Pd-103	0.33 1.5	9 40	0.01 0.03	1 3

¹Wendt 2013 AAPM https://www.aapm.org/education/VL/vl.asp?id=2439

Isotope	Activity threshold (GBq)	Activity threshold (mCi)	Dose rate at 1 m (mSv/hr)	Dose rate at 1 m (mrem/hr)
Y90	NA	NA	NA	NA
Tc-99m	28	760	0.58	58

1 mR/hr = 1 mrem/hr for gamma and x-ray

The activity at which patients could be released was calculated by using, the method discussed in the NCRP Report No. 37, "Precautions in the Management of Patients Who Have Received Therapeutic Amounts of Radionuclides."

$$D(t) = 34.6 \times \frac{\Gamma Q_0 T_P \left(1 - e^{-0.693/T_P}\right)}{r^2},$$
(23.2)

where

- 34.6 = Conversion factor of 24 hrs/day times the total integration of decay (1.44)
- D(t) = accumulated exposure at time t, in R (It assumed that 1 R = 10 mSv = 1 rem)
- Γ = Specific gamma ray constant for a point source, R/mCi-hr at 1 cm
- Q_0 = Initial activity of the point source in mCi, at the time of the release
- T_P = Physical half-life in days
- r = Distance from the point source to the point of interest, in cm
- t = Exposure time in days

23.3 Prostate implants

Good Pre-Plan (Seattle Prostate Institute Criteria)

- Modified uniform loading
- V100: 98-100%
- V150:
 - I-125: 30-40%
 - Pd-103: 40-50%
- V200: 10-20%
- Uretha max: 100-125% (definitely<150%)
- Rectum point: <80%
- Margin: 3-5 mm

23.4 TheraSphere

⁹⁰Y-microsphere therapy usually target the liver, taking advantage of the unique circulatory system in the liver Portal vein (normal liver) and hepatic artery (tumor).²

SIR-Sphere is not discussed here but more detailed descriptions about both microsphere can be found from 2017 AAPM Annual meeting talk, ⁹⁰ Y-Microsphere Therapy: Emerging Trends and Future Directions (link).

Patient selection (an example)

- 62 year-old female with cirrhosis and HCC
- BSA = 1.78

 $^{^{2}}$ http://amos3.aapm.org/abstracts/pdf/68-19792-237349-87867.pdf

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- Child-pugh B
- UNOS T3
- ECOG perfomance status = 1 (Fatigue)
- AFP 809

Before treatment

To avoid radiation pnumanitis, the lung dose for TheraSphere should be less than **30 Gy**. The *lung shunt* (LS) percentage can be calculated from the signals (counts) in Tc-99m (normally 2-4 mCi) MAA planar scintigraphy or SPECT/CT³,

$$lung \ shunt \ (\%) = \frac{Lung \ Counts}{Lung \ Counts + Liver \ Counts} \times 100$$
 (23.3)

where $GMcounts = \sqrt{ANTcount \times POSTcount}$.

Delineation of target volumes is based on digital segmentated angiography - DSA, CT, C-arm CBCT, SPECT/CT. The treatment volume is then converted to mass, using a conversion facotr of 1.03 g/cc. The required activity can then be calculated

Standard model

$$A_{Totoal} = A_{Liver} + A_{Lung}$$

$$D_{Lung} = \frac{50(J/GBq) \times A_{Lung} \times LSF}{M_{Lung}}.$$
 (23.4)

$$D_{Liver} = \frac{50(J/GBq) \times A_{Lung} \times (1 - LSF)}{M_{Liver}}.$$
 (23.5)

Issues of the standard model

- Hetergeneous uptake distribution
- unknow Tumor dose and normal tissue dose

Partiion model

After treatment

Typical patient exposure rates

- Maximum surface: 5 25 mR/hr
- At 1 m: 0.1 0.3 mR/hr

Residual measurement at 30 cm on a template.

The radioactive 90 Y-label microspheres (20-30 μm) are injected via a catheter trans-aterially.

ddcdab

³why we need CT? similar to the function of CT in PET/CT?

Intravascular BT

We have finished a nice book.

IGRT

References

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Several publications about QA issues associated with image-guided radiation therapy

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- TG-154 (2011) QA of US-guided External beam radiotherapy for prostate cancer
- TG-135 (2011): QA for Robotic Radiosurgry
 - planar kV
- TG-179 (2012): QA for image-guided radiation therapy utilizing CT-based technologies
 - kV- and MV-CBCT; fan beam kVCT and MVCT
- TG-147 (2012): QA for nonradiographic RT localization and positioning systems
- Jaffrey (2012) Assuring safety and quality in image-guided delivery of RT

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Cyber Knife

A linear accelerator in CyberKnife generates a 6 MV x-ray beam. The microwave frequency it uses for accelerating electrons is in the range of: a) 500 to 1,000 MHz b) 2 to 4 GHz c) 8 to 12 GHz d) 15-20 GHz

Proton RT

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28.1 Clinac couch

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