



ROYAL INSTITUTE
OF TECHNOLOGY

Nuclear Fuel Cycle 2011

Lecture 5: Detection of Ionizing, Radiochemistry

Gas filled detectors

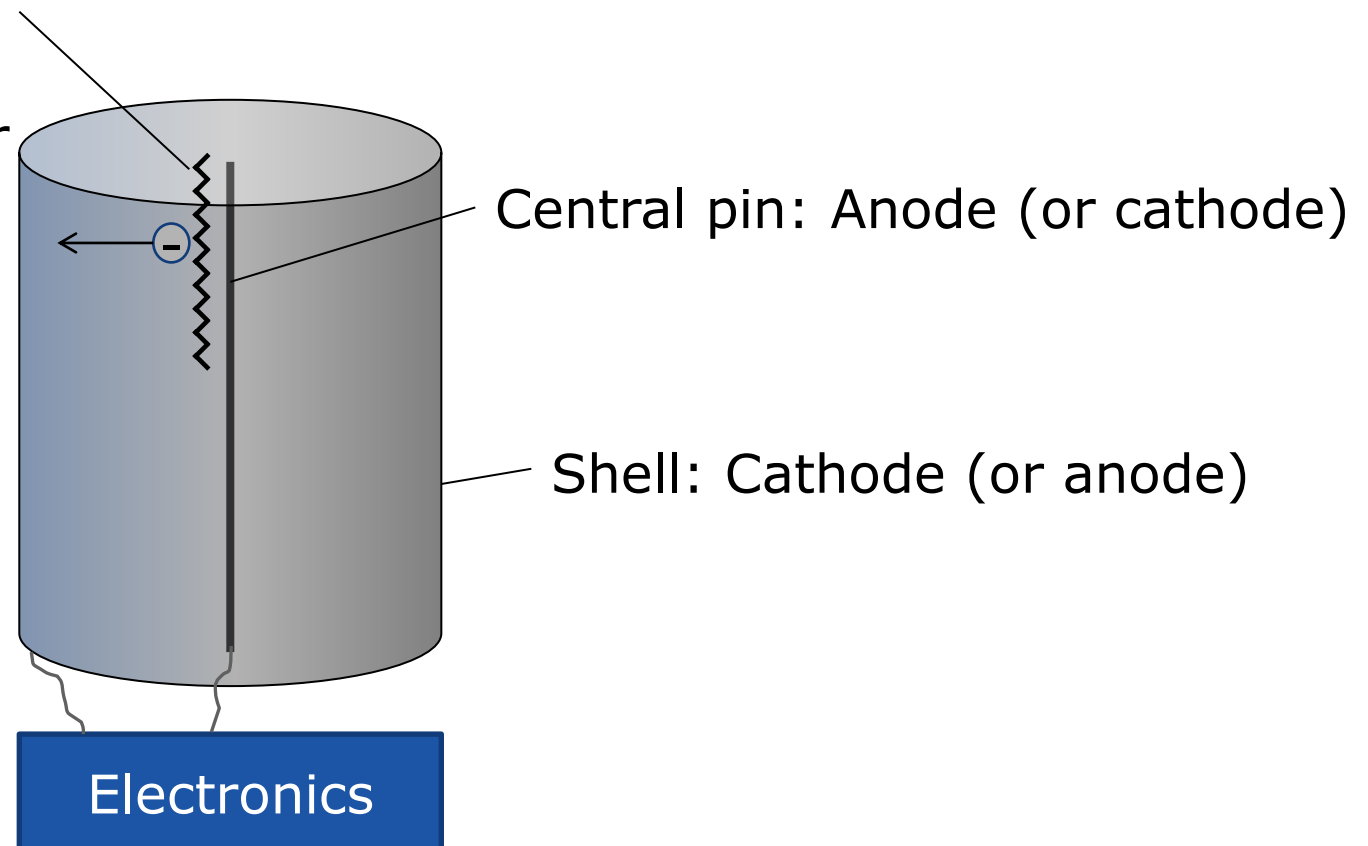
Principle for many of detectors
(GM counters for instance)

Radiation enters chamber

Ionizes gas in chamber

Ions travel towards
cathode/anode

=> Current detected
by electronics



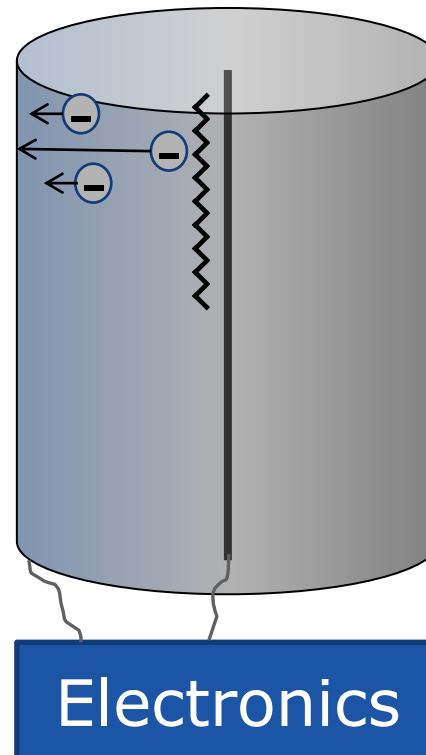
Gas filled detectors

- Energy of α -particle is about 4 MeV
- 100 000 ionizations
- The charge = $10^5 \times 1.602 \times 10^{-19} \text{ C} \sim 10^{-14} \text{ A}$
 - The signal needs to be amplified greatly

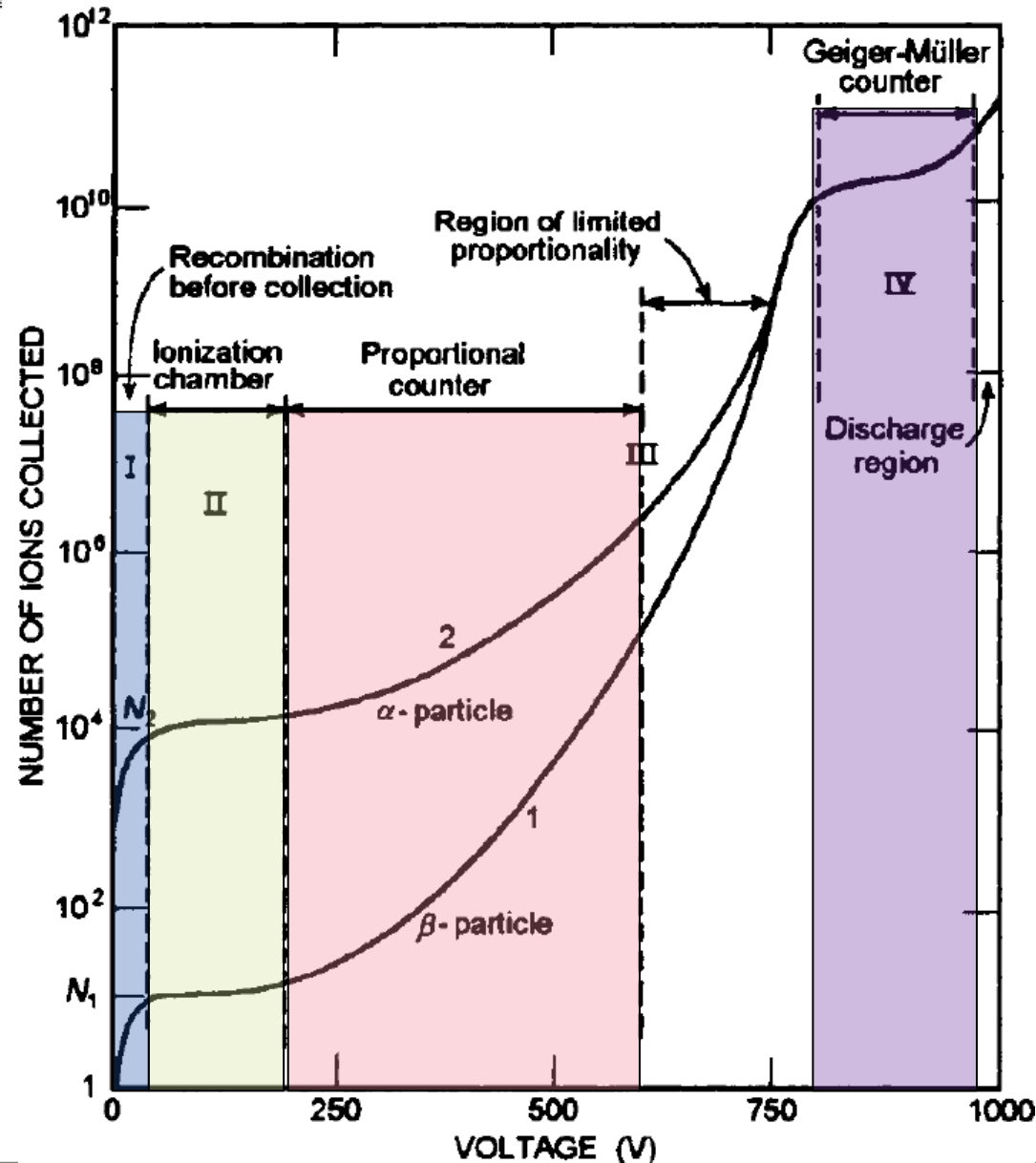
Gas filled detectors

If the potential between anode and cathode is large, the created ion will accelerate and initiate secondary ions

=> Stronger signal



Gas filled detectors



I: ions and electrons recombine before reaching electrodes

II: ionization chamber:
1 ionization = 1 signal

III: proportional region:
signal is proportional to primary ionizations

IV: Geiger-Müller counter:
tube is discharged by every ionization

γ -radiation

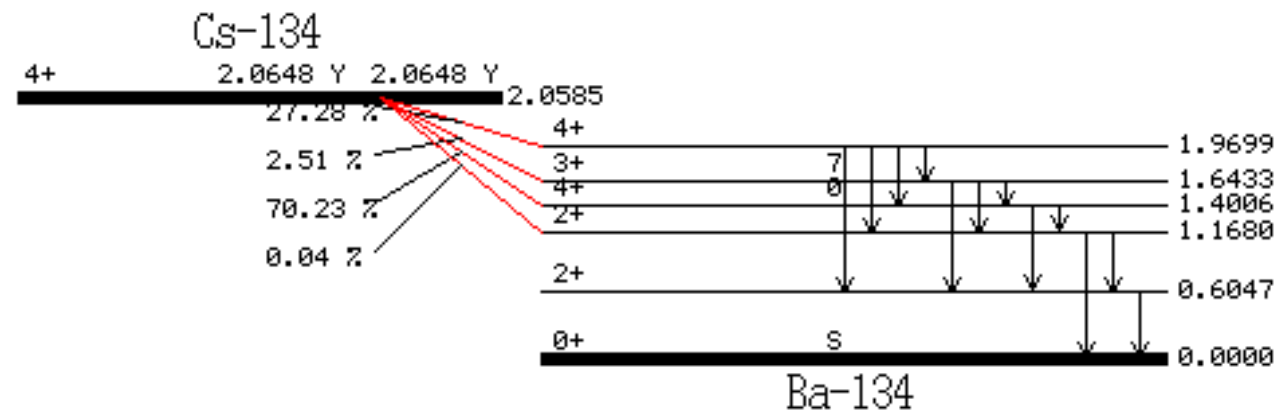
- Gamma radiation has much longer range (lower LET) and will travel through air-filled detector with very few interactions.
 - Instead: Use denser material in detector:
Semiconductor
 - When ionizing radiation interacts with the semiconductor an electron is excited to the conduction band.
The electron then travels towards the anode.
 - Gamma spectrometer (or Multichannel analyzer)
-

γ -counter

Almost every radioactive decay has quantified γ

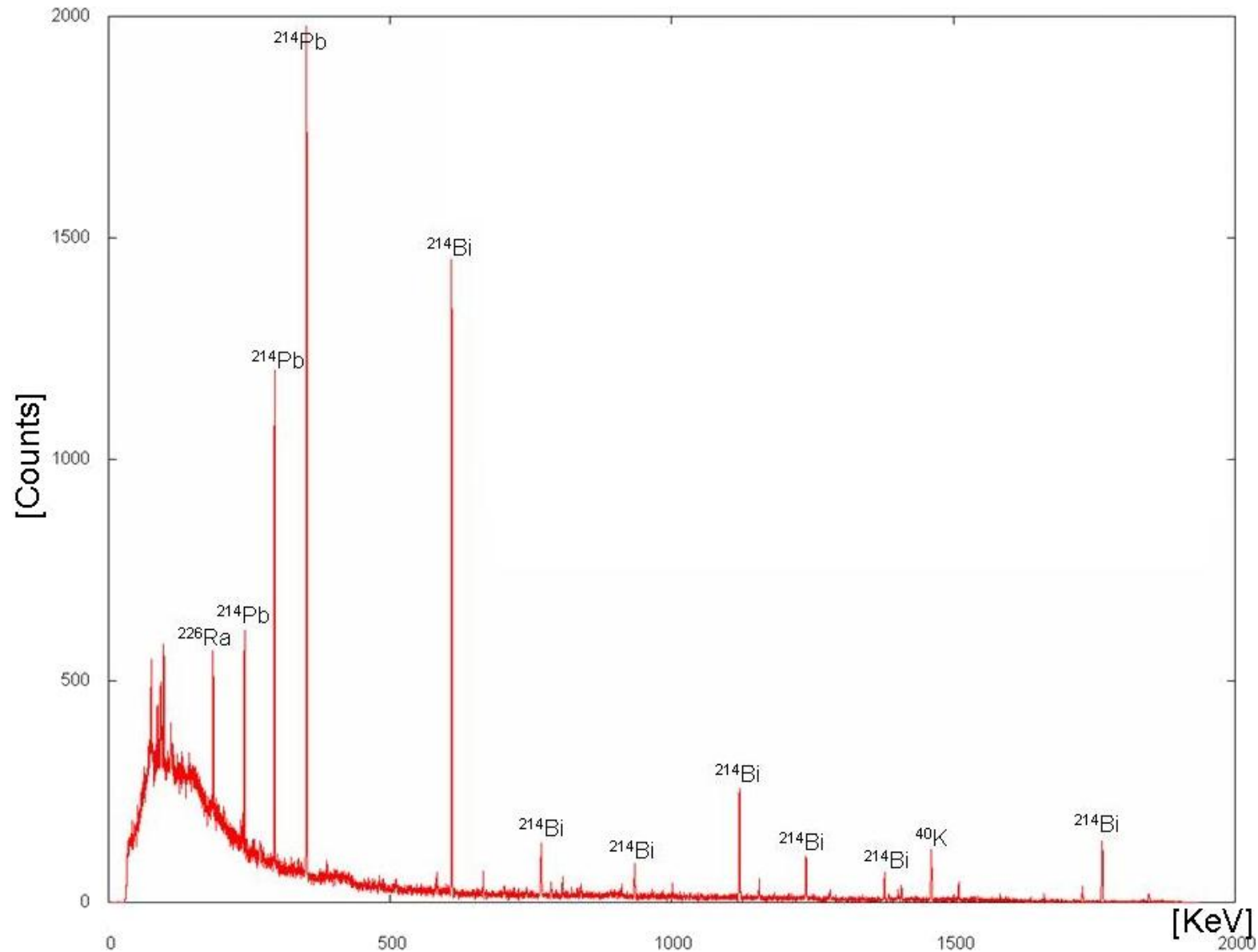


γ -Energy(keV)	Intensity(%)
232.6	0.0011
242.738	0.0272
326.589	0.0162
475.365	1.486
563.246	8.35
569.331	15.38
604.721	97.62
795.864	85.53
801.953	8.69
1038.610	0.988
1167.968	1.789
1365.185	3.014



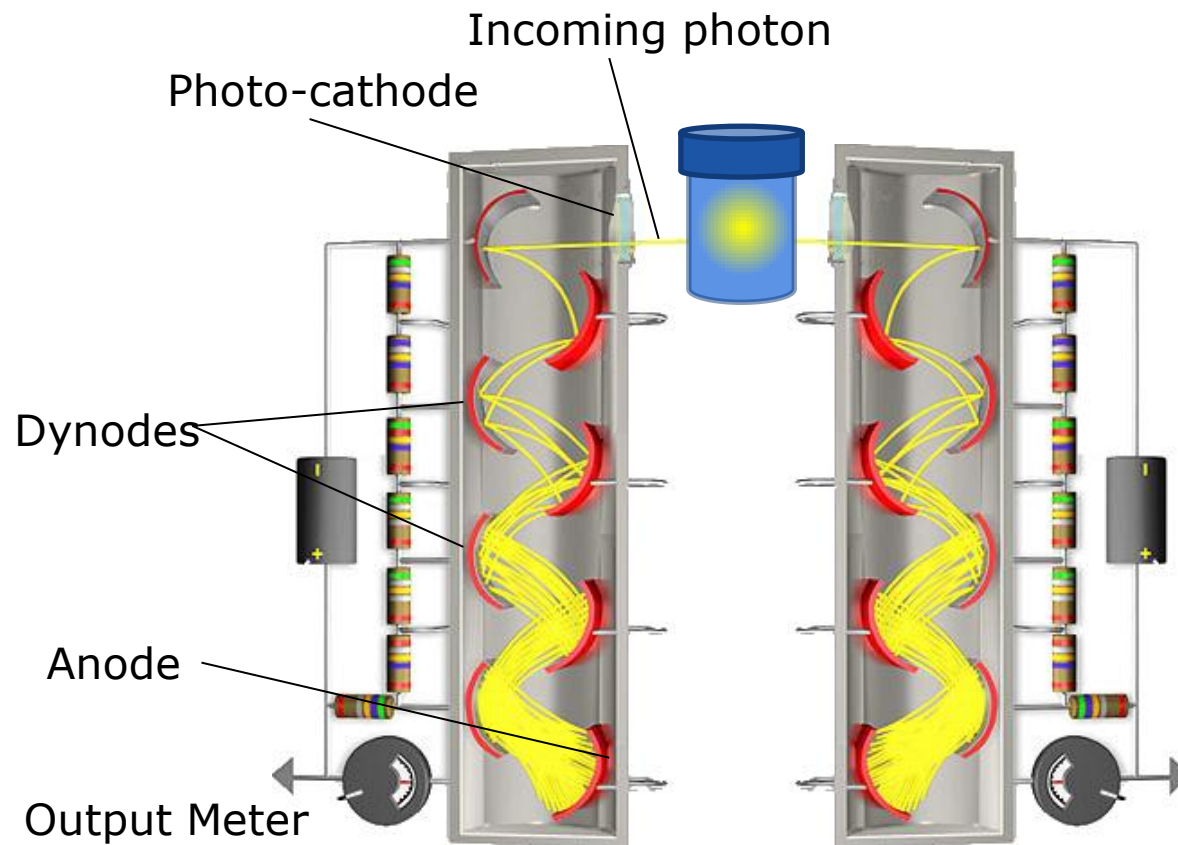
Two distinct peaks

Example γ -spectrum (mineral sample)

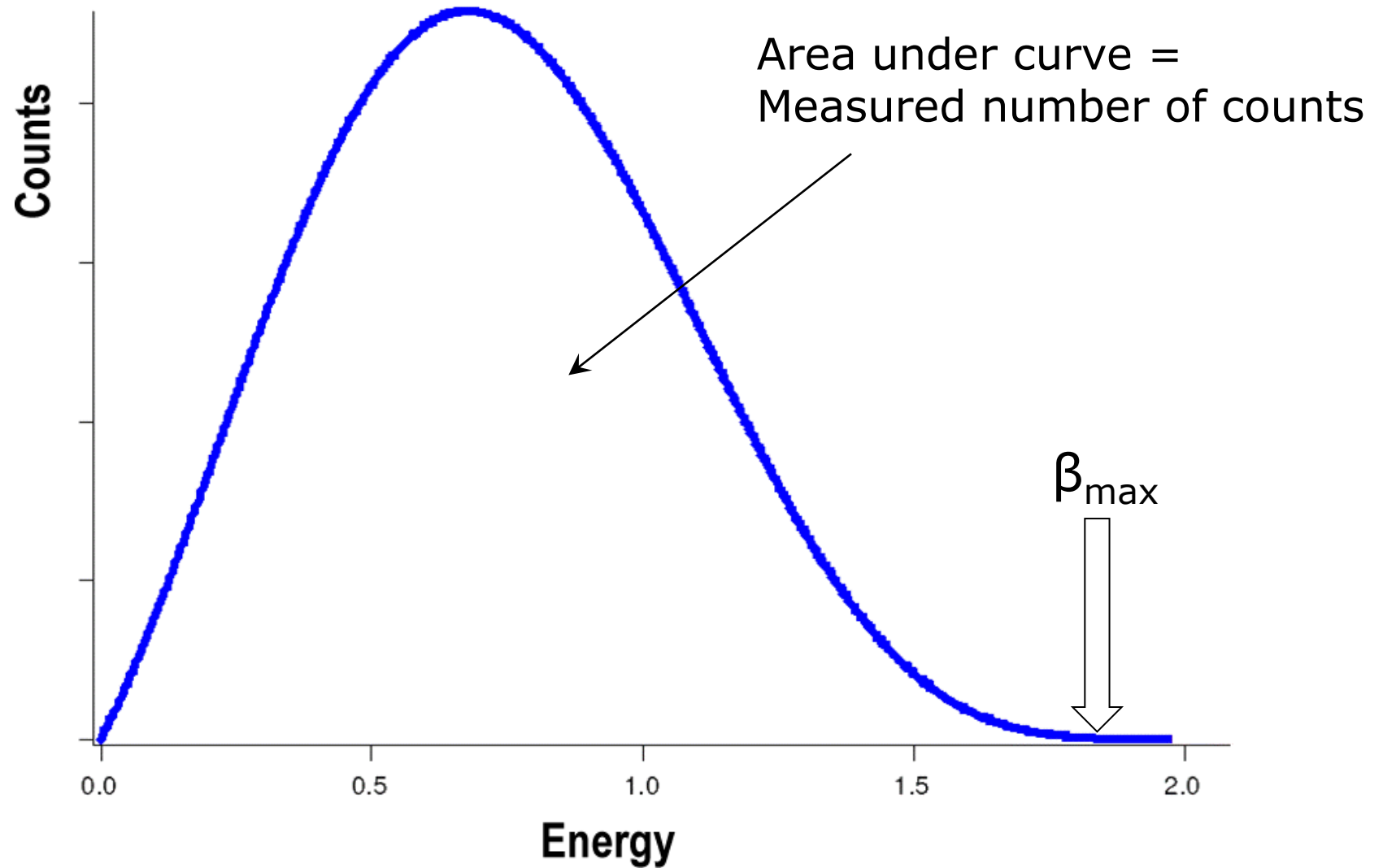


Liquid scintillation

α/β^- radiation is absorbed by scintillation liquid
The energy is given back as light



β -spectrum



Efficiency of a detector

- Some of the activity will never enter detector due to geometry reasons. This has to be considered but is NOT the efficiency
- Some radiation will not interact with detector

$$\text{Efficiency} = \frac{\text{What is detected}}{\text{What enters the detector}}$$

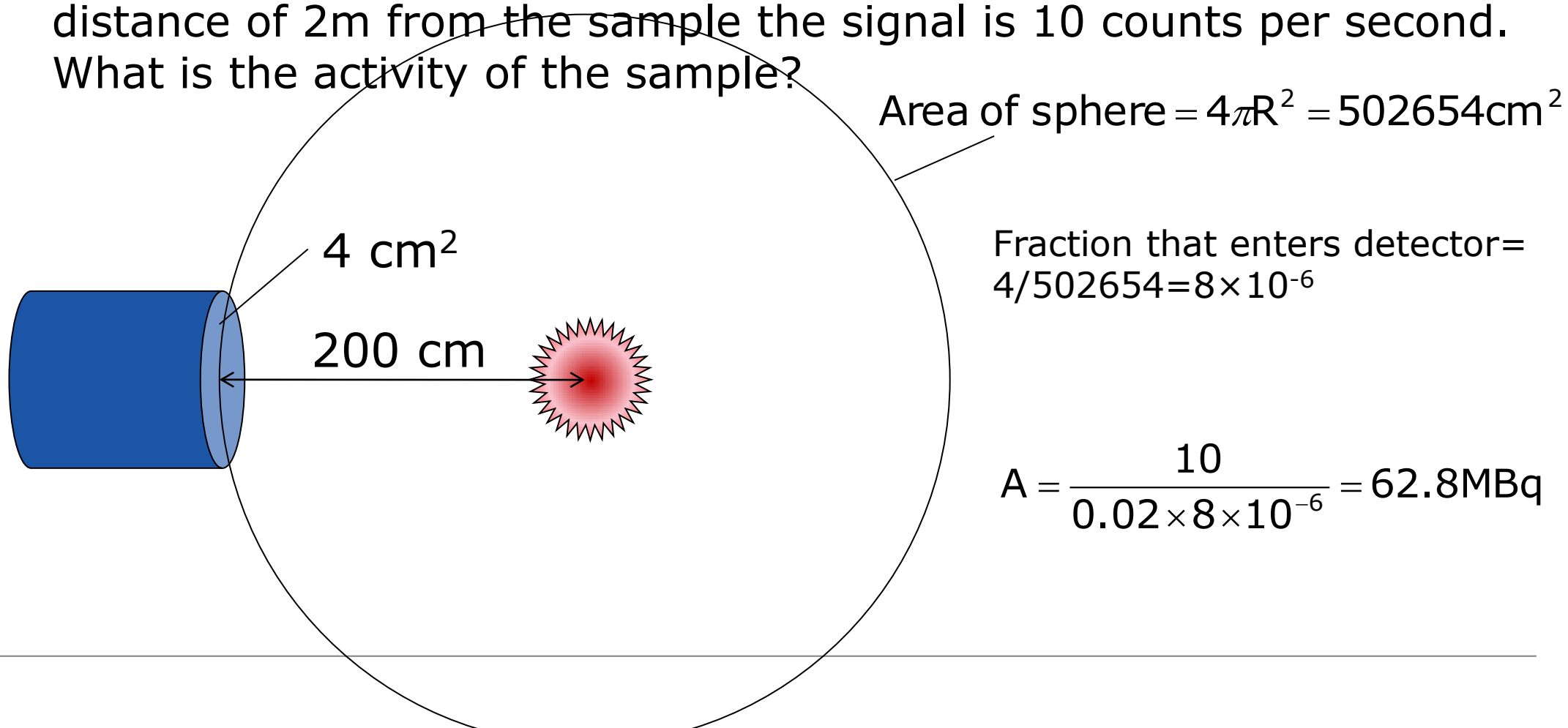
NOTE!

Bq is absolute activity.

cps (counts per second) is measured value!

Example

A portable detector is used to measure the activity of a sample. The surface area of the detector is 4 cm^2 and the efficiency is 2% . At a distance of 2m from the sample the signal is 10 counts per second. What is the activity of the sample?



$$A = \frac{10}{0.02 \times 8 \times 10^{-6}} = 62.8 \text{ MBq}$$



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Radiochemistry I

The use of radioactive isotopes (radionuclides) as tracers to follow chemical processes

Why are radionuclides used?

- The exact same element is used, it has the same chemical and physical properties
 - Radionuclides are independent of pressure, temperature, chemical and physical state
 - Radionuclides are easy to detect and are measured with high precision
 - Does not affect the system (if activity is not too high)
 - No interference of other elements
 - Cheap (compared with for instance ICP-MS)
-

With radionuclides low amounts can be detected

$$A = N\lambda \quad \longleftrightarrow \quad m = A \frac{M}{\ln 2 N_A} t_{1/2}$$

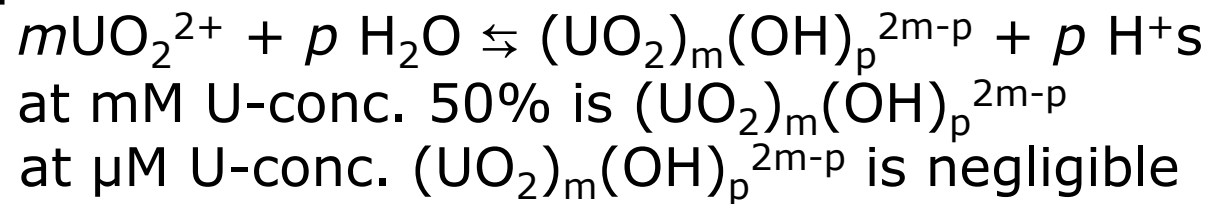
- Assume that 1 Bq can be measured with sufficient accuracy:

$t_{1/2}$	Number of atoms	mol
1 h	5 200	8.64×10^{-21}
1 d	125 000	2.08×10^{-19}
1 y	4.55×10^7	7.55×10^{-17}
10^5 y	4.55×10^{12}	7.55×10^{-12}
10^9 y	4.55×10^{16}	7.55×10^{-8}

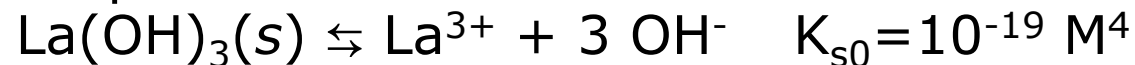
Issues to keep in mind when working with very low concentrations

- Adsorption to walls of vessel
- Formation of radiocolloids

- Equilibrium reactions



- Precipitation



At $[\text{OH}^-] = 1 \text{ mM}$ does 100 MBq/l $^{130}\text{La}^{3+}$ not exceed solubility product

Working with radionuclides

Carrier

A non-radioactive carrier is usually added to a system to ensure normal chemical behavior

Isotopic exchange



is required

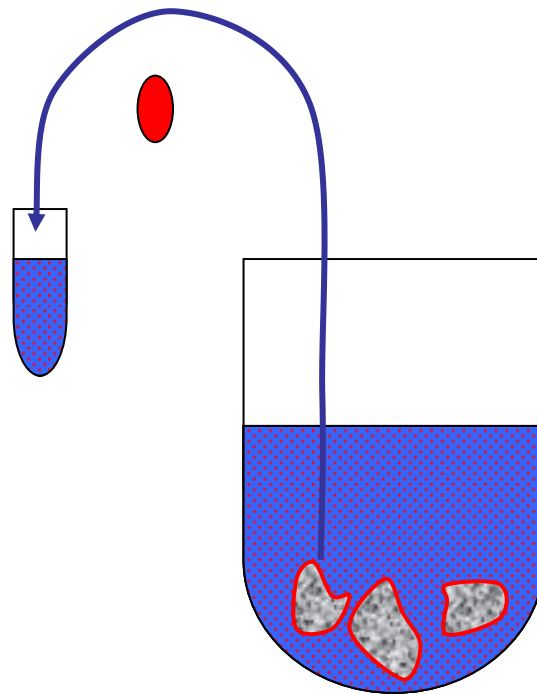
Working with radionuclides

ALARA

- **A**s
 - **L**ow
 - **A**s
 - **R**easonable
 - **A**chievable
-

Example continued

- Determination of Sr-distribution between granite and solution



Water

Add granite

Add tracer (Sr-90)

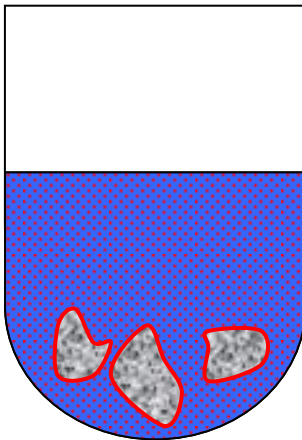
Wait for equilibrium

Take sample from solution

Sr-90 is a β^- -emitter
with β_{max} at 550 keV
 $t_{1/2} = 28.5 \text{ y}$

Example continued

Determination of distribution of cation between granite and solution



To be able to detect the radionuclide, a reasonable concentration of radionuclide would be very low (nM or lower, depending on $t_{1/2}$)

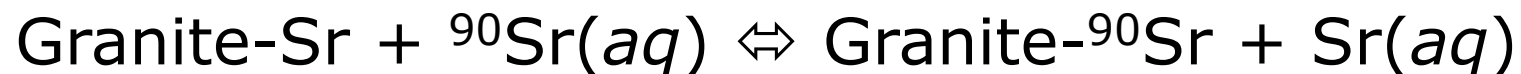
⇒ Very low concentrations: any sorption to glass wall or other loss of radionuclide would have large impact

Adding 10^6 times more radionuclide to obtain a reasonable concentration would cause significant water radiolysis which would change the system

Example continued

Carrier

- We need to add a carrier to our radioactive solution to ensure normal chemical behaviour and obtain reasonable concentrations.
- Isotopic exchange
- $AX + BX^* \rightleftharpoons AX^* + BX$



Working with radionuclides

Selecting radionuclide

- Has sufficient $t_{1/2}$ for the process to be studied to take place
 - Same oxidation state as carrier (isotopic exchange)
 - When very low activities are used the background has to be carefully attended
 - Examine the nature of any radioactive daughters
-

Working with radionuclides

Be careful with daughters

- ^{90}Sr (β^- $E_\beta=0.55$ MeV, $t_{1/2}=28.5\text{y}$)



β -measurement will
include ^{90}Sr and ^{90}Y

- ^{90}Y (β^- $E_\beta=2.3$ MeV, $t_{1/2}=64\text{h}$)



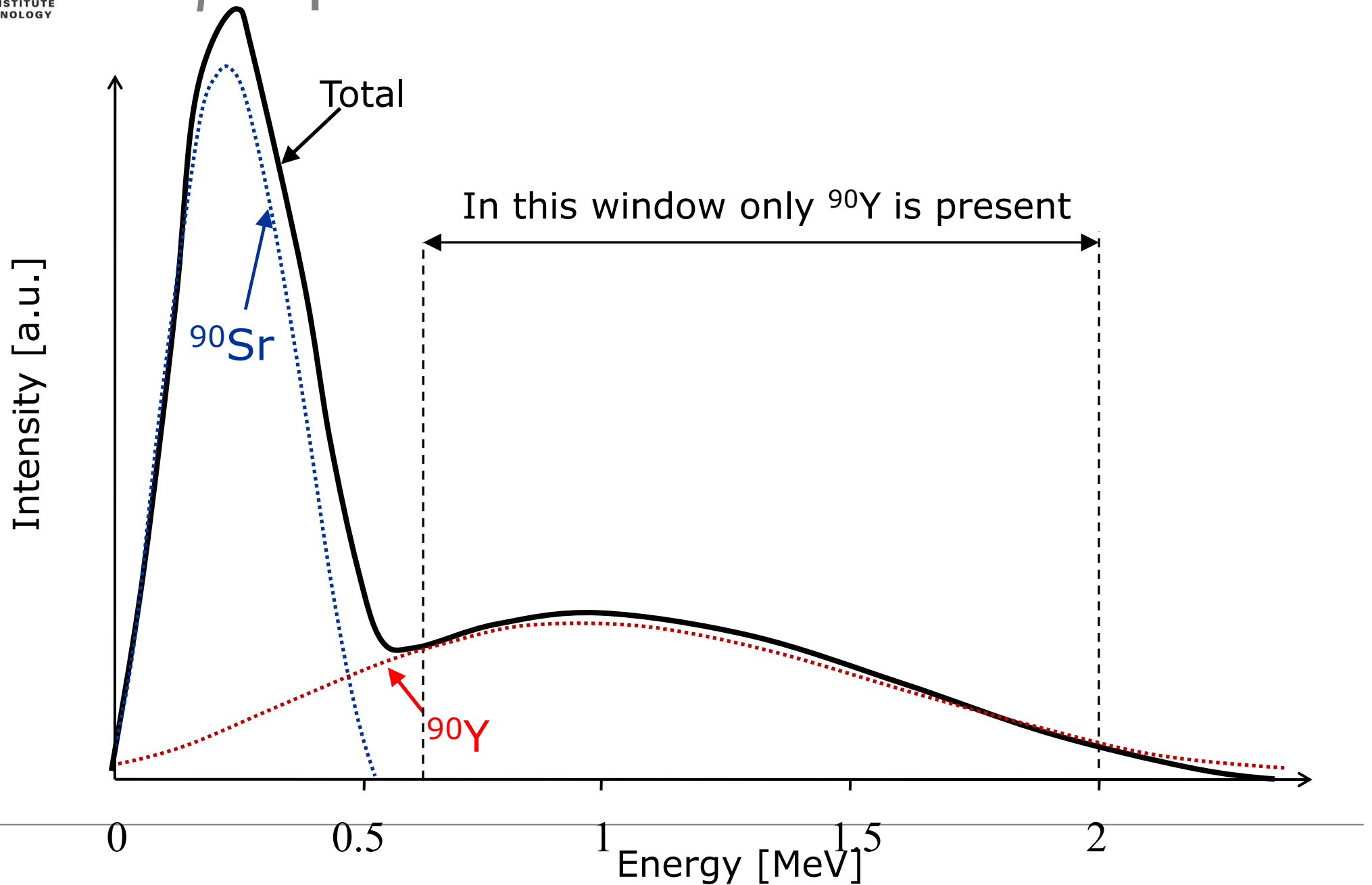
- ^{90}Zr (stable)

Better wait for
secular equilibrium and
measure daughter
via energy discrimination

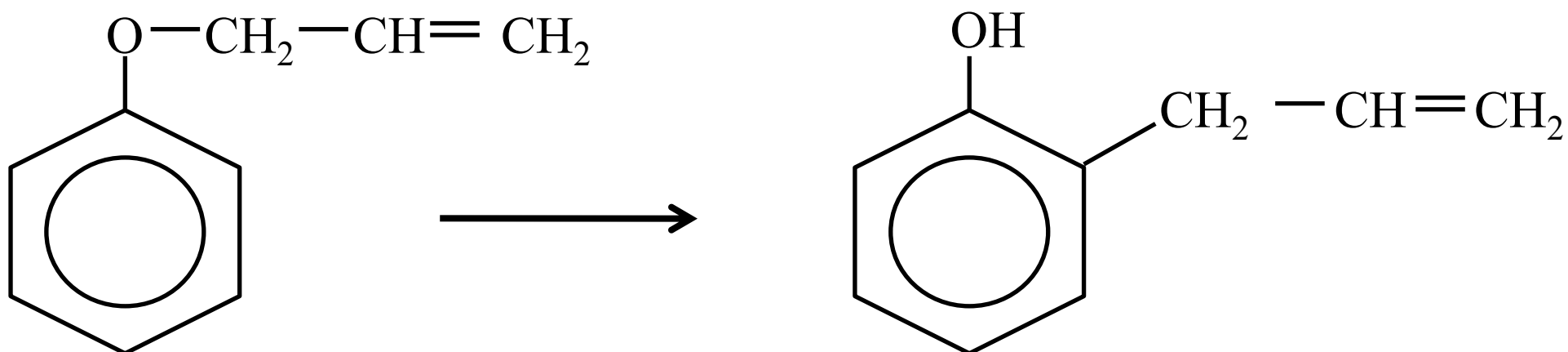
At secular equilibrium

$$A_{\text{Sr-90}} = A_{\text{Y-90}}$$

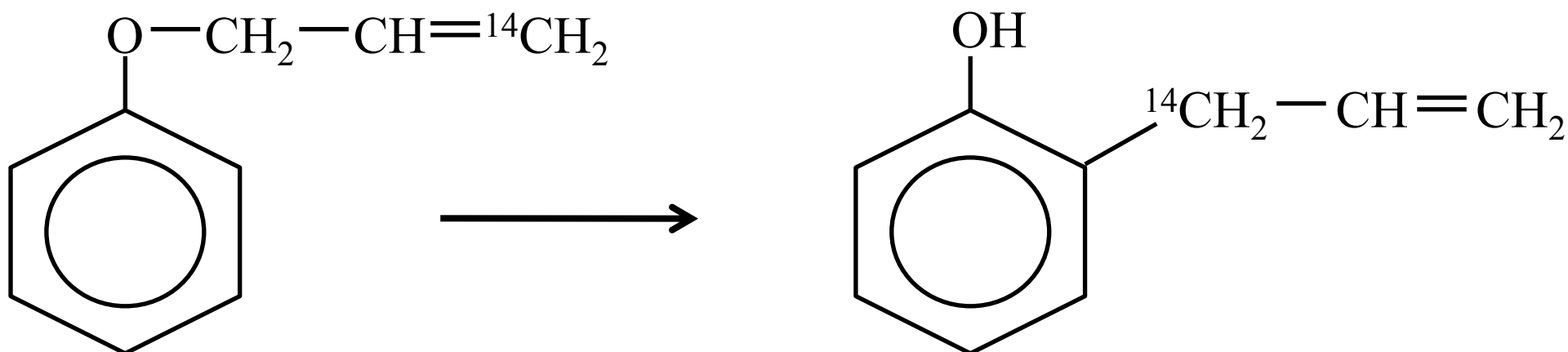
β -spectrum for ^{90}Sr and ^{90}Y



Determine reaction mechanisms
-For instance, the Claisen allyl rearrangement

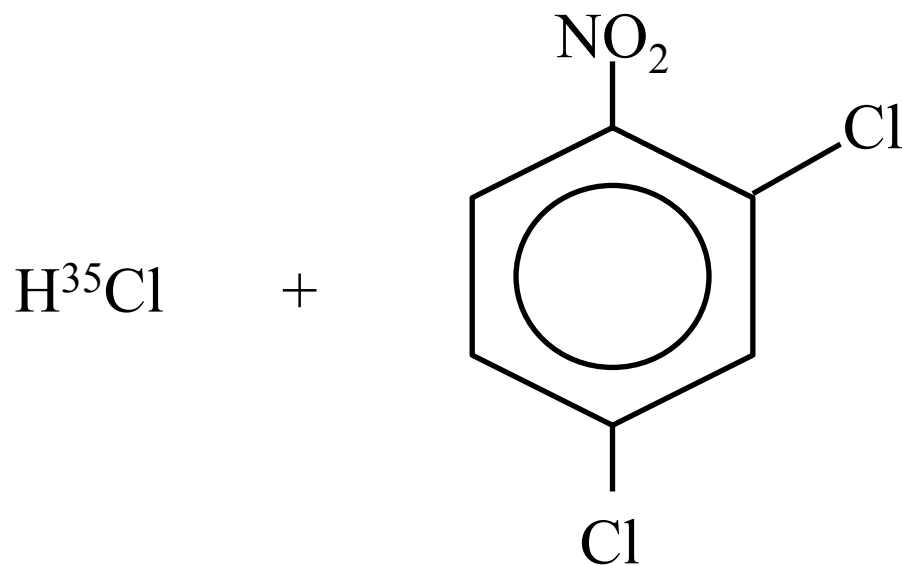


Determine reaction mechanisms
-For instance, the Claisen allyl rearrangement



- From decomposition products the mechanism can be determined

Using isotope exchange rates to determine characteristics of a compound



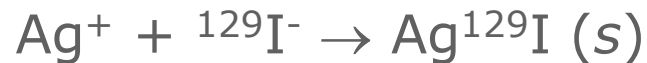
- The rate of exchanging the Cl at the
- ortho and para positions differs

Radiometric analysis

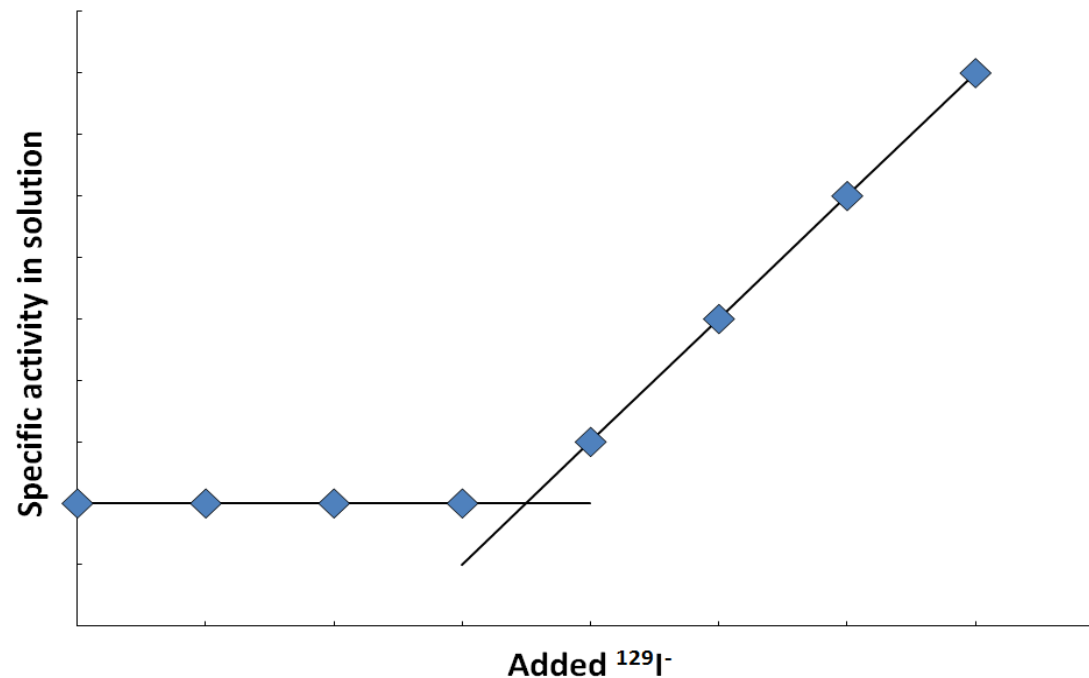
Radiometric titration

- Two phase titration in the presence of radionuclide

For instance determine Ag^+ concentration in a solution



Add ${}^{129}\text{I}^-$ and monitor activity in solution



Radiometric analysis

Radiometric titration

- Very low concentrations can be detected
 - Used as calibration for other instrumental methods
-

Isotope dilution

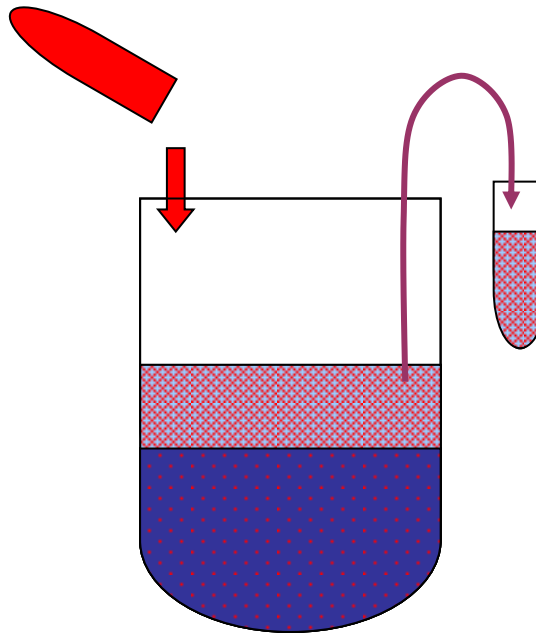
- Used when quantitative separation of one compound is not possible
- Qualitative separation is needed, though

System with element of
unknown mass w_u

Add known weight (w_0) RN
(same element)

Selectively separate element
for instance via extraction

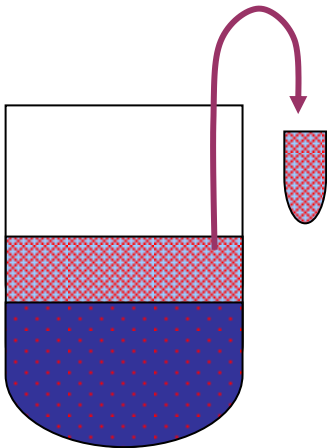
Take sample and measure
activity and mass
(Specific activity, S_m)



Isotope dilution

The specific activity is the same in the whole system.

$$S_m = S_{system} = \frac{\text{Total activity}}{\text{Total weight}} = \frac{w_0 S_0}{w_u + w_0}$$



And the unknown weight
can be calculated from

$$w_u = \left(\frac{S_0}{S_m} - 1 \right) w_0$$

Take sample and measure
activity and mass
(Specific activity, S_m)

Isotope dilution

Applications

Determine

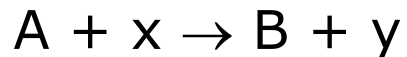
- The naphthalene concentration in tar
 - Fatty acids in mixtures of natural fat
 - Amino acids in biological material
-

Activation Analysis

- By irradiating a sample with neutrons, a small amount of the atoms in the sample will take up a neutron and become radioactive.
- The sample has been “activated”

Neutron Activation Analysis Equation

Consider a general nuclear reaction:



A, B are elements, x and y are elementary particles.

- The production of N_B can be expressed as

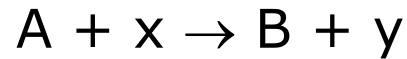
$$\frac{dN_B}{dt} = \Phi_x \sigma N_A$$

- Φ , Particle flux [neutron/cm²,s]
- σ , Cross-section for accepting particles [barn]
(1 barn=10⁻²⁴ cm²/neutron)
- N_A , Number of atoms A

ASSUMPTIONS:

- Flux and energy of particles constant through the target.
 - N_A is constant (an insignificant amount is transmuted).
 - σ is small (so that flux & energy remain constant).
-

Neutron Activation Analysis Equation



$$\frac{dN_B}{dt} = \Phi_x \sigma N_A$$

When the nuclide produced is radioactive, its decay has to be taken into account.

$$\frac{dN_B}{dt} = (\Phi_x \sigma N_A) - \lambda N_B$$

Integrate between $t=0$ ($N_B=0$) and $t=t$

$$N_t = \frac{\Phi_x \sigma N_A}{\lambda} (1 - e^{-\lambda t})$$

or writing it as activity ($A = N \lambda$):

$$A_t = \Phi_x \sigma N_A (1 - e^{-\lambda t})$$

Neutron Activation Analysis Equation

$$A_t = \Phi_x \sigma N_A (1 - e^{-\lambda t})$$

The produced radionuclide will decay and we have to account for the activity loss:

$$A_{t,t'} = A_t e^{-\lambda t'}$$

We get

$$A = \Phi \sigma N_A (1 - e^{-\lambda t_{\text{irr}}}) e^{-\lambda t_{\text{cool}}}$$

NAA Example

- 5 g. Chrome is irradiated for 3 hours in a neutron flux of 10^{13} n/cm²,s.
- What is the activity 48 hours after the irradiation?

$$A = \Phi \sigma N_A (1 - e^{-\lambda t_{\text{irr}}}) e^{-\lambda t_{\text{cool}}}$$

$$\Phi = 10^{13} \text{ n/cm}^2, \text{s}$$

$$\sigma = ?$$

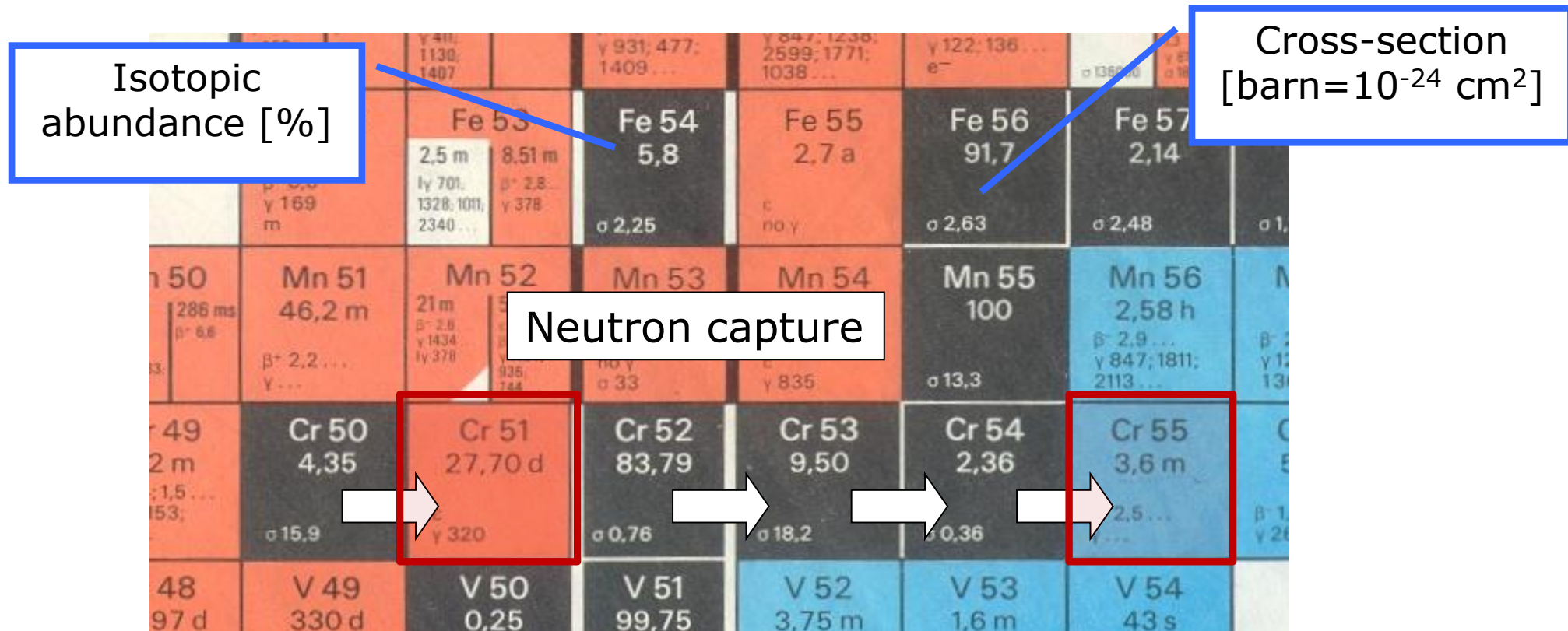
$$N = ? \text{ (total amount Cr} = 5/52 * 6 * 10^{22} \text{)}$$

What is the amount of each isotope of Cr?

} Segré or
<http://atom.kaeri.re.kr/ton/>

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Stable Isotopes	Cr-50	Cr-52	Cr-53	Cr-54
Isotopic abundance [%]	4.35	83.79	9.5	2.36
$t_{1/2}$ of neutron activated product	27.7 d	stable	stable	3.5 min
Cross section [barn]	15.9	0.76	18.2	0.36

Only Cr-51 will be radioactive after 48 hours

NAA Example

- 5 g. Chrome is irradiated for 3 hours in a neutron flux of 10^{13} n/cm²,s.
- What is the activity 48 hours after the irradiation?

$$A = \Phi \sigma N_A (1 - e^{-\lambda t_{\text{irr}}}) e^{-\lambda t_{\text{cool}}}$$

$$\Phi = 10^{13} \text{ n/cm}^2, \text{s}$$

$$\sigma = 15.9 * 10^{-24} \text{ cm}^2/\text{n}$$

$$N = 5 / 52 * 6 * 10^{23} * 0.0435 = 2.5 * 10^{21}$$

$$\lambda = \ln 2 / (27.7 * 24 * 3600) = 2.9 * 10^{-7} \text{ s}^{-1}$$

$$t_{\text{irr}} = 3 * 3600 = 10\,800 \text{ s}$$

$$t_{\text{cool}} = 48 * 3600 = 172\,800 \text{ s}$$

$$A = 1.19 \times 10^9 \text{ Bq}$$

$$A = 1.2 \text{ GBq}$$

Advantages with NAA

- Highly sensitive
 - Nondestructive
 - Determination of elements in complex samples;
 - Environmental samples
 - Mineral samples
 - Archeological samples
-

Example

Environmental history of waters in Sweden

1. Mussels were collected from rivers and lakes in Sweden. Mussels build shell thicker each year; The composition of the shell reflects the water chemistry.
 2. Shells were sliced.
 3. Sent to neutron irradiation source.
 4. Sample was measured directly at arrival from reactor (short lived nuclides dominate spectrum).
 5. Sample was measured 2 weeks after irradiation (short lived nuclides not present anymore).
 6. Evaluation of the spectra.
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Example

Environmental history of waters in Sweden

