

Gamma spectroscopy

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Group B

Measurement date: November 4, 2021.

Hand in date: November 16, 2021

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1 Introduction

The goal of this laboratory class was to measure the gamma radiation emitted by radioactive isotopes. Alpha and beta decays release energy on the order of one up to a couple MeV. Based on the measured spectrums both the isotope type and the activity of the isotope can be determined.

1.1 Measurement setup

The two main parts of the measurement setup were the HPGe detector and the amplitude analyzer. HPGe stands for high purity germanium, this makes up the bulk of the detector. Incoming ionizing radiation creates electron-hole pairs. The number of created pairs is to good approximation proportional to the energy absorbed by the semiconductor. The detector is indirectly connected to a liquid nitrogen bath, which cools down the detector considerably. This makes it that thermal fluctuations don't create a significant number of electron-hole pairs, so the only source of current that is measured comes from electron-hole pairs created by the ionizing radiation. These pairs are prohibited from recombining by the high voltage connected to the germanium detector.

The other key component is the amplitude analyzer. It measures the charge carried by electric impulses. This charge is directly proportional to the number of electron-hole pairs created during the time interval of the pulse and therefore characteristic of the energy of the incoming radiation. Based on this charge the amplitude analyzer chooses the corresponding energy bin, and increments the counter corresponding to that bin by one. Finally this analyzer is connected to a computer, where the data is interpreted and displayed. The relation between the bin numbers and the energy of the radiation is expected to be linear, but the exact parameters of the relation are not known before calibration.

2 Calibration

To determine the parameters of the linear relation between bin number and energy peaks with known energy are needed to be measured. During our measurement we used ^{232}Th isotope. The energy of two easily identifiable gamma peaks is known. The first one is 238.6keV, this is the peak with the largest intensity. The other characteristic peak is at

2614.7keV, this is the highest energy peak observed. These peaks were identified in the spectrum. To validate the identification of peaks, after using the two peaks to calibrate the linear relation between energy and bin number, other peaks of the ^{232}Th isotope were identified and we checked if their energy matches up with the values read off from tables. These other peaks were at 580keV and 908keV according to the calibration. These are the 583.191keV and 911.316keV peaks respectively, both corresponding to ^{208}Tl , which is part of the thorium series. The linear fit for energy was

$$E = 1.3247\text{keV} \cdot n - 4.6\text{keV}, \quad (1)$$

where n is the bin number. From the validation we know the energy measured from the bin number is accurate within a couple keV. This is precise enough to identify peaks. The source of the error is that the current impulse coming from the HPGe detector isn't exactly in linear relation with the energy absorbed by the detector.

3 Granite sample

The first sample we examined after calibration was a granite rock sample. Table 1. contains the data required for the activity calculations.

Energy [keV]	Net area	Net area rel. error	Intensity	η	η rel. error	Isotope
351.3	36043	0.6%	3.689e-01	2.1975%	9.0%	^{214}Pb
999.1	620	14.2%	5.892e-03	0.8457%	7.6%	^{234}Pa
186.7	12230	1.1%	3.280e-02	3.1371%	9.1%	^{226}Ra
295.0	23525	0.9%	1.919e-01	2.0425%	6.6%	^{214}Pb
1765.1	4430	1.9%	1.620e-01	0.4518%	8.8%	^{214}Bi
1118.6	5706	2.0%	1.550e-01	0.7023%	9.3%	^{214}Bi
242.1	10554	1.7%	7.453e-02	2.6091%	8.7%	^{214}Pb

Table 1: Granite sample data

The spectrum was taken over a period of 1000s. From this the activity can be calculated using

$$A = \frac{N}{I\eta t}, \quad (2)$$

with error

$$\sigma_{\text{rel}}(A) = \sqrt{\sigma_{\text{rel}}(\eta)^2 + \sigma_{\text{rel}}(N)^2}. \quad (3)$$

These equations give the results in table 2.

Energy [keV]	Activity [kBq]	Activity error [kBq]	Isotope
351.3	4.4	0.4	^{214}Pb
999.1	12.4	2.0	^{234}Pa
186.7	11.9	1.1	^{226}Ra
295.0	6.0	0.4	^{214}Pb
1765.1	6.1	0.5	^{214}Bi
1118.6	5.2	0.5	^{214}Bi
242.1	5.4	0.5	^{214}Pb

Table 2: The activities corresponding to peaks in the granite sample.

Corresponding peaks can be averaged. The correct way to calculate the average given that we know the uncertainties is the following,

$$\langle A \rangle = \frac{\sum_i A_i / \sigma_i^2}{\sum_i 1 / \sigma_i^2}. \quad (4)$$

The uncertainty of this averaged number is

$$\sigma_A = \frac{1}{\sqrt{\sum_i 1 / \sigma_i^2}}. \quad (5)$$

With the above formula I determined the combined activity of the lead and bismuth isotopes. To determine if a significant fraction of the radon gas escapes the granite sample I also calculated the weighted average of the isotopes before and after the radon in the decay chain. Only the radium and protactinium peaks correspond to isotopes which are before radon in the decay chain. The results are summed up in table 3. and in figure 1.

Group	Activity [Bq]	Activity error [Bq]
^{214}Pb	5282	244
^{214}Bi	5614	367
Before ^{222}Rn	12012	957
After ^{222}Rn	5384	203

Table 3: Averaged activities of the granite sample

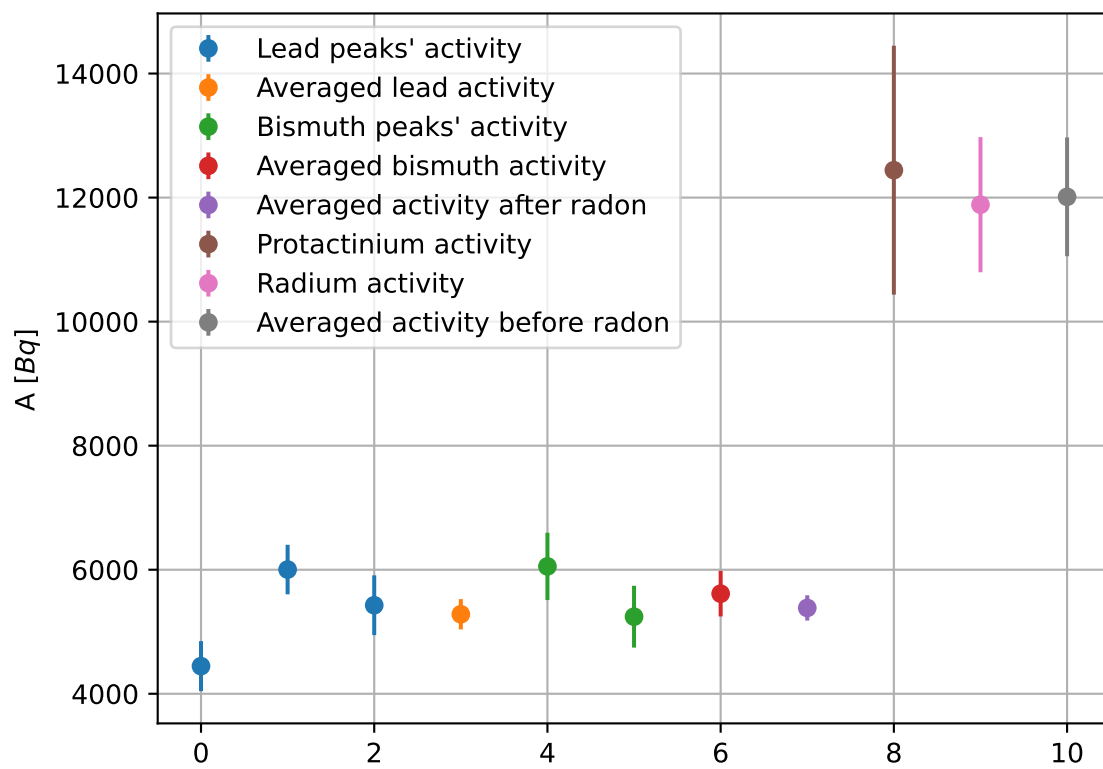


Figure 1: Graphical representation of activities and errors corresponding to peaks and their averages.

From figure 1. it can be seen that the error in the activity corresponding to the 186keV radium peak and the 1001keV protactinium peak is large compared to their difference. This means the amount of ^{235}U , which would contribute to the 186keV peak can not be determined from this measurement. 0.7% ^{235}U contributes

$$\frac{0.7\%\lambda_{235}}{0.7\%\lambda_{235} + 99.3\%\lambda_{238}} = 4.3\% \quad (6)$$

to the 186keV peak. The relative error in the activity of the 186keV peak is 9.2% . This means 0.7% ^{235}U is consistent with the measurement, any concentration between 0% and roughly 2% is.

Based on the averaged activity before and after the radon, we can estimate the amount of radon gas that escapes the rock sample before decaying.

$$r = \frac{A_1 - A_2}{A_1}, \quad (7)$$

where r is the ratio of radon gas that escapes before decaying, A_1 and A_2 are the averaged activities measured from elements before and after the radon in the decay chain, respectively. The error of this ratio is given by

$$\sigma_r = \sigma \left(1 - \frac{A_2}{A_1} \right) = \frac{A_2}{A_1} \sqrt{\left(\frac{\sigma_{A_1}}{A_1} \right)^2 + \left(\frac{\sigma_{A_2}}{A_2} \right)^2}. \quad (8)$$

The result of this calculation is that $(55.2 \pm 4.0)\%$ of the radon gas escapes the granite sample before decaying.

From the activity of the isotopes before the radon gas the amount of ^{238}U can be calculated using its decay rate. This is because those isotopes are guaranteed to be in secular equilibrium granted the rock formed sufficiently long ago.

$$N = \frac{A}{\lambda}, \quad (9)$$

$$\sigma_N = \frac{\sigma_A}{\lambda}, \quad (10)$$

where $\lambda = \frac{\log(2)}{T_{1/2}} = 4.92 \cdot 10^{-18} \frac{1}{s}$ is the decay rate of ^{238}U . From this the number of ^{238}U atoms in the granite sample is $(2.44 \pm 0.19) \cdot 10^{21}$, which corresponds to a mass of $(0.966 \pm 0.077)g$ ^{238}U in the granite sample. With the mass of the granite sample The concentration of ^{238}U could be given in gram/ton units.

4 Soil sample

The data of the measured peaks from the soil sample are summarized in table 4.

Energy [keV]	Net area	Net area rel. error	Intensity	η	η rel. error	Isotope
658.5	407	5.7%	8.998e-01	1.1082%	8.6%	^{134}Cs (^{134}Ba)
120.9	437	6.6%	2.843e-01	3.3051%	7.7%	^{152}Eu
343.4	172	9.9%	2.649e-01	1.6480%	8.2%	^{152}Eu
1405.3	27	22.8%	2.075e-01	0.4523%	8.9%	^{152}Eu
776.7	50	22.4%	1.274e-01	0.9176%	8.9%	^{152}Eu
244.9	65	14.6%	7.494e-02	2.5473%	9.5%	^{152}Eu

Table 4: Soil sample data. The 658.5 keV peak is emitted by ^{134}Ba after a ^{134}Cs beta decays.

The first peak is emitted by a transition of the barium atom, after a cesium atom decays into it. The activities can be calculated in the same ways as for the granite sample. The results are listed in table 5.

Energy [keV]	Activity [kBq]	Activity error [kBq]	Isotope
658.5	44.8	4.6	^{134}Cs (^{134}Ba)
120.9	51.0	5.2	^{152}Eu
343.4	43.3	5.5	^{152}Eu
1405.3	31.6	7.7	^{152}Eu
776.7	46.9	11.3	^{152}Eu
244.9	37.4	6.5	^{152}Eu

Table 5: The activities corresponding to peaks in the soil sample. The 658.5 keV peak is emitted by ^{134}Ba after a ^{134}Cs beta decays.

Averaging the europium peaks' activities as in the case of the granite sample I got $(43.2 \pm 2.9)Bq$. These activities are visually represented by figure 2.

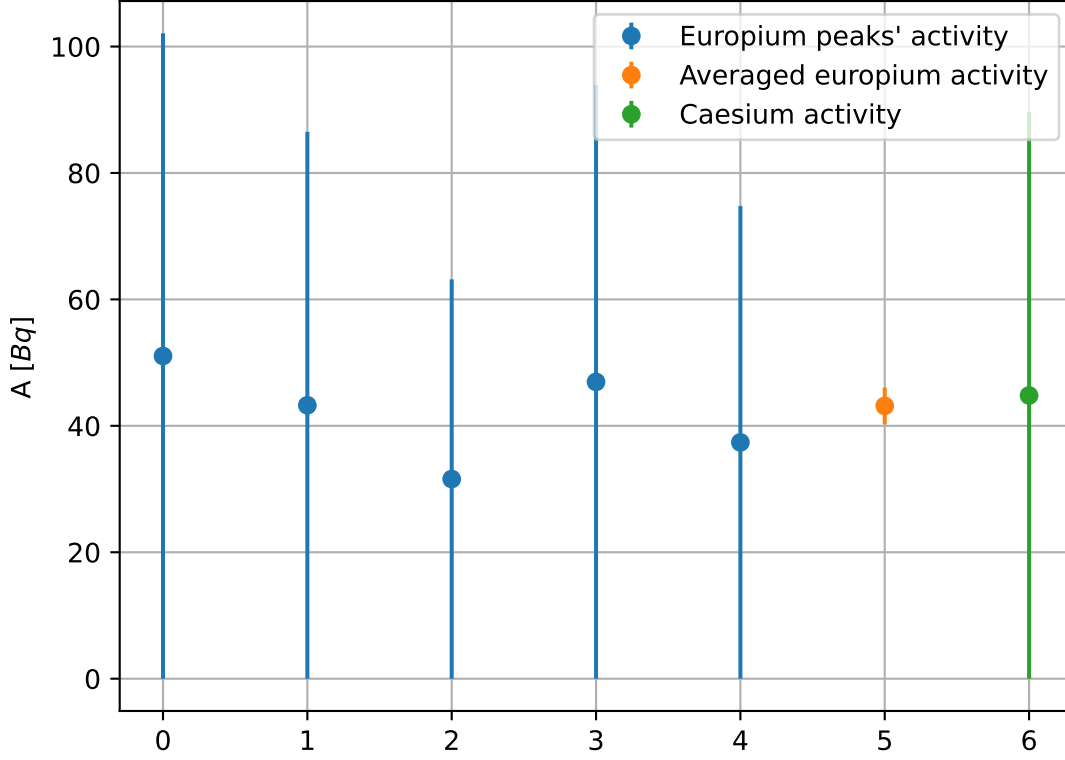


Figure 2: Graphical representation of activities and errors corresponding to peaks and their averages.

5 Conclusion

The lab introduced us to gamma spectroscopy and analyzing the data obtained. In the first part we calibrated the measurement with a thorium sample. The next part was about taking the spectrum of a granite sample and identifying the peaks seen. During evaluation I concluded that a significant portion of the radon gas escapes the sample and that the 186keV radium peak is consistent with a 0.7% ^{235}U isotope content (out of all the uranium atoms). I also determined the uranium content of the sample. Finally I repeated some of the calculations for a soil sample from near a nuclear bomb test.