

# Identifying nuclides by their gamma ray spectra

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## Abstract

We analyze the gamma ray spectra of known nuclides and compare our results to the expected values in order to calibrate our set-up for identifying unknown samples by their gamma ray spectra. We determine that an unknown nuclide's gamma ray spectrum has a singular photopeak corresponding to an energy of 842(31) keV. This is most likely  $^{54}\text{Mn}$ .

## 1 Introduction

In this lab, we investigate the process known as Gamma Ray Spectroscopy, whereby we are able to identify various radioactive nuclei by the signature gamma rays that are emitted when they decay. This is done using an inorganic scintillator. We are interested in gamma rays with energy between about 50 and 2000 keV as these are fairly unique to the nuclei that produce them.

## 2 Theory

When a nuclide decays, it can decay in one of a few ways. The decays we are interested in are those where the nucleus either gains a proton and loses a neutron, or loses a proton and gains a neutron, changing elemental structure. When this happens, in order to conserve the charge of a proton being created or destroyed, we see a  $\beta$  particle (positive or negative) emitted along with the corresponding neutrino. For this reason we call this decay  $\beta$  decay. After the decay, if we sum up the masses of the resulting nuclide,  $\beta$  particle, and neutrino, we see that the total is less than the mass of the nuclide before the decay. This missing energy is usually accounted for by the resulting nuclide being in an excited state and when it transitions to a lower energy state, it will emit a photon. These are the photons we are detecting.

Our detector is a scintillator detector, meaning it absorbs the incident gamma rays and emits rays of proportional but lesser energy, to be absorbed by a photomultiplier tube, whose signal is amplified by an array of amplification devices. In a perfect world this system would produce current pulses that are proportional only to the incident gamma rays, but since we don't live in a perfect world, we see pulses corresponding to secondary processes that end up on our spectra. The main processes of concern are photoelectric absorption, Compton scattering, pair production, and backscatter, on which more details can be found in Gilmore Ch2.2 [1].

The detector and amplifier set-up leads into either an oscilloscope or a Multi Channel Analyser (MCA), allowing us to observe the current pulses. The MCA counts the number of times that a pulse of a certain voltage is detected and plots that on a histogram, giving us our spectrum for the given radioactive source. These spectra, to a certain degree, are unique to the nuclide that decayed to produce them, and thus we should be able to identify an unknown radioactive source purely from its gamma ray spectrum.

## 3 Method

We used the apparatus configured as in Figure 3.1. The detector is an NaI scintillator detector and photomultiplier tube. The preamplifier is an ORTEC model 113. The amplifier is an ORTEC model 572. The MCA is a SPECTECH UCS-30 running with 1024 channels.

The oscilloscope was used to monitor the voltage of the current pulses produced by the amplifier. The MCA has a maximum input voltage of 5V, so the gain on the amplifier was changed in order to keep all pulses of interest below 5V.

The radioactive sources we were able to analyse were the following:  $^{22}\text{Na}$ ,  $^{60}\text{Co}$ ,  $^{109}\text{Cd}$ ,  $^{133}\text{Ba}$ ,  $^{137}\text{Cs}$ ,  $^{152}\text{Eu}$ , and an unknown source. We started by letting the MCA run for a while to collect background data. Then we placed each source in front of the detector and let the MCA collect data for between 10 minutes to an hour on each source.

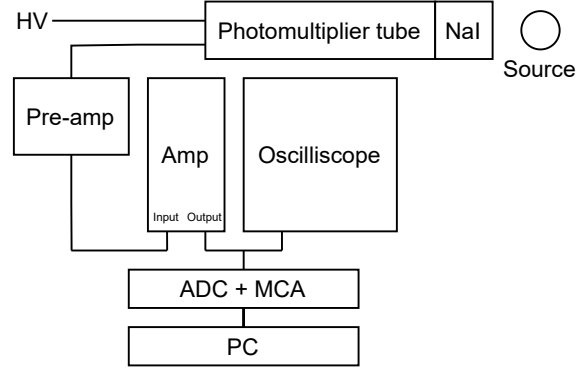


Figure 3.1: Schematic of the set-up used in the experiment.

## 4 Analysis

The data exported by the MCA tells us how many times a current pulse reached the detector within the time spent collecting data, discretised into channels. We normalised these counts with respect to the time spent collecting data, namely the live time as it accounts for the shaping time of the MCA, and then subtracted the background data from them to try to get a cleaner signal. Plotting this data as a histogram we were then able to identify the photopeaks by eye and find an approximate range of channels on which the plot resembles a gaussian distribution. We could then use `scipy.optimize.curve_fit` to fit a gaussian function to the data on those regions in order to approximate a mean for each photopeak. Figure 4.1 shows two of our samples' spectra.

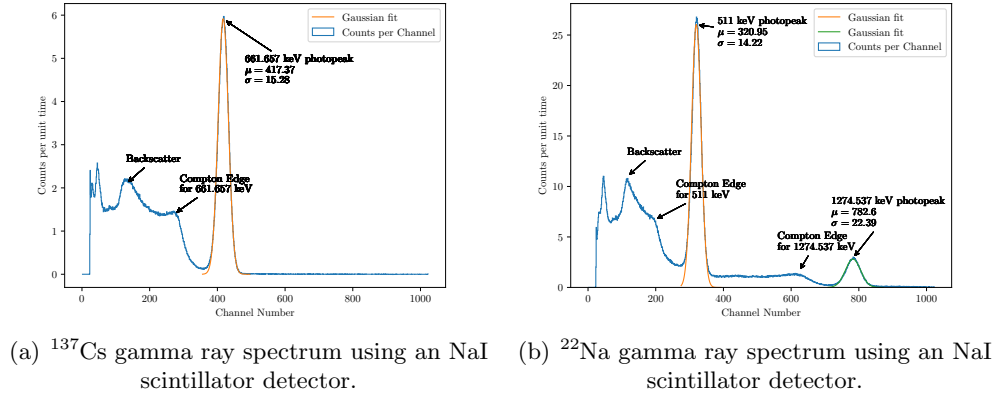
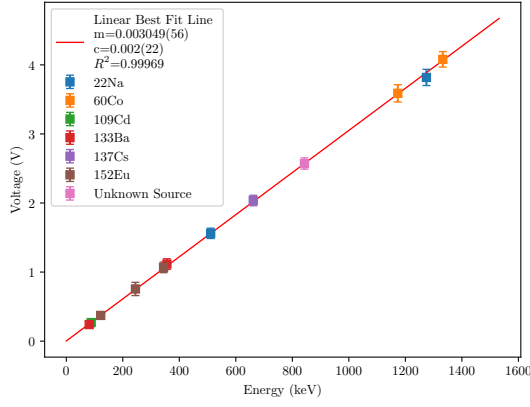


Figure 4.1: Gamma ray spectra for 2 of the radioactive sources to demonstrate the process of finding  $\mu$  and  $\sigma$ .

As our goal was to identify the unknown source, we wanted to be able to accurately estimate the energy associated with its photopeak. In order to do this we needed to find a relation between MCA channel number and photon energy. To do this we found the approximate voltage of the centroid value of each clearly identifiable peak (voltage is associated with channel number by a scaling factor of 5/1024) and then found the expected energy of that peak using the Nudat2 database [2] and could then correlate the two with a linear least squares regression.



	Channel	Expected Energy [keV]
<sup>22</sup> Na	320(15)	511
	782(24)	1274.537(7)
<sup>60</sup> Co	735(26)	1173.228(3)
	835(23)	1332.494(4)
<sup>109</sup> Cd	55.2(59)	88.0336(10)
<sup>133</sup> Ba	49.1(82)	88.9979(11)
	228(16)	356.0129(7)
<sup>137</sup> Cs	417(16)	661.657(3)
<sup>152</sup> Eu	76(10)	121.7817(3)
	155(20)	244.6974(8)
	219(16)	344.2785(12)

Figure 4.2: Correlating detected voltage with gamma ray energy

We then had our calibration equation:

$$E_\gamma = \frac{V - 0.002}{0.003049}$$

$$u(E_\gamma) = E_\gamma \sqrt{\left( \frac{\sqrt{u(V)^2 + 0.022^2}}{V - 0.002} \right)^2 + \left( \frac{0.000056}{0.003049} \right)^2}$$

where  $E_\gamma$  has units keV and  $V$  is the detected voltage. Using this equation we were able to determine an energy for the photopeak in Figure 4.3:  $E_\gamma = 843(31)$  keV.

When it comes to identifying which nuclide it is that produced this spectrum, it's not a particularly simple job. We began by looking at the gamma ray table on <http://atom.kaeri.re.kr/> [3], setting the bounds on energy to be the upper and lower bounds of our 68% confidence interval and setting the minimum half life to 10 days. This gave us 19 results. Looking at all the results with a non-negligible intensity, 14 in total, we noticed that only one of the options had a single emitted gamma ray from ground state, and no 511 keV annihilation gamma rays of significant intensity. This option was <sup>54</sup>Mn, whose only gamma ray emission of significant intensity is of energy 834.848(3) keV with intensity 99.9760(10)% [2].

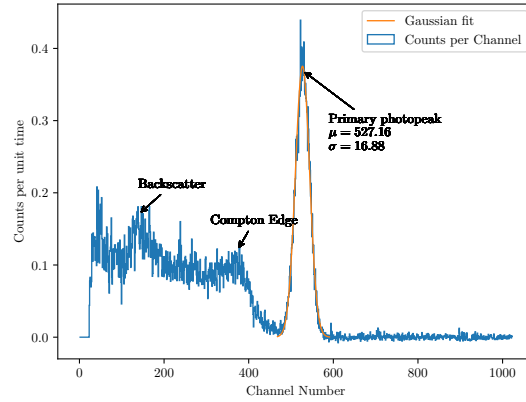


Figure 4.3: The unknown sample's gamma ray spectrum

## References

- [1] Gordon Gilmore. (2008). *Practical Gamma-ray Spectrometry*. 2. Chichester: Wiley
- [2] Nudat 2. Nndc.bnl.gov. <https://www.nndc.bnl.gov/nudat2/chartNuc.jsp>
- [3] Nuclear Data Center at KAERI. Atom.kaeri.re.kr. <http://atom.kaeri.re.kr/>