Self-absorption Correction Factors: Applying A Simplified Method to Analysis of Lead-210 in Different Environment Samples by Direct Counting of Low-energy Using HPGe Detector

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Abstract. This study aimed to determine the self-absorption correction factors of lead-210 (210Pb) in various Syrian environment samples. Seven sediments, five soils, and four plant samples were analysed by Gamma Spectroscopy using simple and direct analysis method called Cutshall. The method is based on measuring the penetration of gamma which emitted from a standard source, prepared in the laboratory by deposition of QCYB40 Standard Solution on stainless steel disk. The source was placed on top of the studied sample and the reference air sample during the measurement. The purpose was to study the self-absorption inside each sample by calculation of its self-absorption factors without knowing its chemical composition. The self-absorption correction results for the sediment samples SE3, SE6 and SE8 ranged between 36% and 45%, and 34% to 42% for the soil samples S5 and S1, respectively. Also, for the four plant samples, it recorded variance range from 4% to 18%. This is due to the difference in the density of the G4 sample, which appeared to be very low. However, the self-absorption correction factors CF were set for the different environmental samples, and the results show that the density factor of the sample is not the only influent factor in the CF values for low energies measurement, the sample chemical composition (sample matrix) is also more effective in addition to the samples’ particle sizes.

1 Introduction

Radioactive elements uranium and thorium, like most chemical elements, are formed in stars and comprise a small amount of the material that formed in the earth naturally. The radioactive isotopes uranium-238 (238U) and thorium-232 (232Th) have decay times (half-lives) which are comparable with, or larger than, the age of the earth, so they have always been present in the earth’s crust and within the tissues of all living species [1].

Environmental monitoring and analysis of the isotopes’ radioactivity concentration in different environment samples is essential to protecting public and environment from any radioactivity pollution, radiometric anomalies investigation, and also for studying the transportation of radioactive and dating. Lead-210 (210Pb) is a radioactive isotope (22.3 years) produced as a result of the radioactive and dating. Lead

[1161keV, and 0.00013% of Bismuth is decay to Thallium with α emitted. The 210Po decay with α emitted and two different energies, the first is 5.3MeV (99.999%), the second energy is 4.5MeV (0.001%), to reach the stable isotope 206Pb (Figure 1). 210Pb is concentrated in sediments of lakes, rivers, and continental and oceanic water [2], [3]. Its concentration in groundwater reaches 7Bq/L. It also increases in some plant leaf such as lettuce and spinach, where it can reach 6Bq/kg of fresh weight. Eating 200 g of the plant daily leads to an internal dose of 55µSv per year [2], 210Pb is not considered a radiation hazard for human health by itself, but by its daughters’ 210Bi and 210Po and to its chemical toxicity [2].

So, most the radiation dose caused by it is daughters because of its lower energy of β particles (61keV and 15keV), also γ emits (46.5keV) and the lower emission ratio of α-particles.

Fig. 1. 210Pb and its daughters 210Bi and 210Po decay [3]
There are different techniques used to determine the self-absorption correction factors [5] [6] [7]. Since 1983, Cutshall developed an experimental method to determining self-absorption correction factors for low gamma-radiation energies [4]. He assigned the \( ^{210}\text{Pb} \) (46.54 keV) germanium detector (with a beryllium window used for low gamma radiation) to measure the transmittance of gamma rays through sediment samples and by placing the \( ^{210}\text{Pb} \) source on top of the sample while collecting the spectrum, with the same positioning of the source relative to the sample and the detector surface. This method is called the Cutshall method or the permeability method, and it is the method used in this work.

2 Materials and method

2.1 Cutshall Method

Cutshall method depends on the penetration of gamma rays through the sample which we need to set its self-absorption correction factor. Self-absorption correction has many parameters such as source intensity, the isotopic to be assigned and its energy (high or low), sample thickness, density, chemical composition and the weight ratios of each compound or element, the organic content, the humidity coefficient, the homogeneity of the sample and others. Also, there are other parameters that depend on detector specifications, type, dimensions, and source location to the sample and detector during the measurement and spectrum collection. This method requires a radioactive source that emits energy close or similar to the isotope energy. Also, it is required activity of the source higher the studied sample, meaning that the activity of the sample has to be negligible when compared to the source activity. Since the isotope to be determined in this study is \( ^{210}\text{Pb} \), this required preparing the source of \( ^{210}\text{Pb} \), which is placed at the top of the studied sample during the collection of the spectrum and the count value (Count) is recorded under the peak of \( ^{210}\text{Pb} \) and the collection time per (Sec). The count rate (Count Sec-1, CPS) is obtained by dividing the count by the collection time. Consider \( I_\text{s} \) represents the count rate for the studied sample with the source and \( I_\text{air} \) is the count rate for the reference air (empty petri can). Table 1 shows the counting results for the reference air sample. If we consider the attenuation of air is equal to zero, we obtain the correction factor for self-absorption experimentally as the following formula:

\[
P_\text{air,Exp}^s = \frac{\ln \left( \frac{I_\text{s}}{I_\text{air}} \right)}{1} \tag{1}
\]

The self-absorption correction factors shown in Tables 2, 3, and 4 were calculated using formula 1. The good thing with the formula, is it does not need to know the standard source activity, and it is sufficient that its activity is higher than the expected activity of the sample. In addition, its results are quick and fairly accurate. By multiplying the measured sample activity by the self-absorption correction factor, we obtain the corrected activity of the sample. The correction ratio is calculated from the following formula [8]:

\[
C_\text{air,Exp}^s = \left( 1 - \frac{1}{P_\text{air,Exp}^s} \right) \times 100 \%
\]

\( P_\text{air,Exp}^s \): represents the experimentally calculated self-absorption correction factor Exp., for the sample, and for the air. \( C_\text{air,Exp}^s \): Represents the correction ratio for self-absorption correction factors experimentally Exp., for sample \( s \), and for air. This formula was used to calculate the self-absorption correction ratio Tables 2, 3, and 4.

2.2 Sample preparation

Samples (sediment, soil, plant) were prepared and filled in a Petri dish, with a thickness of 1.2 cm, and volume of 24.73 cm³.

2.3 Spectrum collection and analysis

The spectrum was collected by a HpGe Spectroscopy system. The spectrum for each sample was collected during 500 sec with the standard \( ^{210}\text{Pb} \) on top of the sample. The previous step was repeated for the different sample’s sediment, soil, plant and air (with empty petri dish). A \( ^{210}\text{Pb} \) peak was recorded for each measurement separately, taking into account the position of the standard source and the sample to each other also, to detector surface center which have to be the same position.

| Table 1. shows the measured permeability results for the reference air sample of the detector (its thickness is 1.2 cm). |
| Sample | Density g/cm² | Sec | COUNT |
|---|---|---|---|
| Air | 1.2923 × 10⁻² | 500 | 2880 |

| Table 2. Correction Factors for sediment |
| Sample ID | g/cm³ | \( F_{\text{Self-Absorption}} \) | \( C_{\text{Self Absorption}} \) % |
|---|---|---|---|
| SE3 | 1.85 | 1.82 | 45 |
| SE4 | 1.41 | 1.52 | 34 |
| SE5 | 1.57 | 1.63 | 39 |
| SE6 | 1.32 | 1.56 | 36 |
| SE7 | 1.44 | 1.60 | 38 |
| SE8 | 1.51 | 1.55 | 36 |
| SE9 | 1.52 | 1.76 | 43 |

| Table 3. Correction Factors for soil samples |
| Sample ID | g/cm³ | \( F_{\text{Self-Absorption}} \) | \( C_{\text{Self Absorption}} \) % |
|---|---|---|---|
| S1 | 1.68 | 1.74 | 42 |
| S2 | 1.65 | 1.71 | 41 |
| S3 | 1.58 | 1.69 | 41 |
| S4 | 1.52 | 1.67 | 40 |
| S5 | 1.50 | 1.52 | 34 |

| Table 4. Correction Factors for plant samples |
| Sample ID | g/cm³ | \( F_{\text{Self-Absorption}} \) | \( C_{\text{Self Absorption}} \) % |
|---|---|---|---|
| G1 | 0.75 | 1.17 | 14 |
| G2 | 0.78 | 1.12 | 11 |
| G3 | 0.79 | 1.22 | 18 |
| G4 | 0.30 | 1.05 | 4 |
3 Results and Discussion

Tables 2 and 3 show that the self-absorption correction ratio ranged between 45% and 36% for the SE3, SE6 and SE8 sediment samples while the correction ratio was less for S5 and S1 soil samples also ranged between 34% and 42%, respectively. It is also a clear relation with most samples’ densities, the samples with higher density recording a higher correction of self-absorption, and opposite for the samples with low density values (Figures 2 and 3).

The SE4, SE8 and SE9 sediment samples were noted variations in CF values in opposite to the soil’s samples except the S4. This variation is due to the different particle sizes in these samples itself and sample matrix.

4 Conclusions

The self-absorption correction factors CF of $^{210}\text{Pb}$ were set for the different environmental samples using Cutshall’s method. The results show the density factor is not the only influent factor in the CF values for low energy measurement, the sample chemical composition (sample matrix) is also more effective in addition to the samples’ particle sizes. The difference in particle size for the sediment and soil samples was an effective factor in calculation of the appearance density of the samples too even after sample grinding.

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