Determination of Absorbed and Effective Dose from Natural Background Radiation around a Nuclear Research Facility

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ABSTRACT

This study presents result of outdoor absorbed dose rate and estimated effective dose from the naturally occurring radionuclides $^{232}$Th and $^{238}$U series $^{40}$K, around a Nuclear Research Reactor at the Centre for Energy Research and Training (CERT), Zaria, Nigeria. A high-resolution in situ $\gamma$-ray spectrometry was used to carry out the study. CERT houses a 30Kw Research Reactor and other neutron and gamma sources for Research and Training. The values of absorbed dose rate in air for $^{232}$Th, $^{238}$U and $^{40}$K range from 8.2±2.5 - 24.5±3.6 nGy h$^{-1}$, 1.9±1.2 - 4.6±2.5 nGy h$^{-1}$ and 12.2 ±5 - 38±6.7 n Gy h$^{-1}$ respectively. The estimated total annual effective dose outdoor for the sites range from 27.3 - 79.9 µSv y$^{-1}$. This showed that radiation exposure level for the public is lower than the recommended value of 1 mSv y$^{-1}$. Hence, the extensive usage of radioactive materials within and around CERT does not appear to have any impact on the radiation burden of the environment.

KEY WORDS: Absorbed dose rate, effective dose, radiation exposure, Global Positioning System (GPS), Centre for Energy Research and Training (CERT), radioactive materials, Naturally Occurring Radionuclides (NORs), terrestrial radionuclides, natural environmental, energy resolution

INTRODUCTION

The presence of Naturally Occurring Radionuclides (NORs) in the Earth's crust is well known. More than twenty primordial nuclides have been identified, most of them exceeding the age of the Earth by several orders of magnitude [1]. The process of nucleosynthesis in stars forms the primordial radionuclides (also called terrestrial background radiation). Some of the primordial NORs, including $^{40}$K , $^{87}$Rb, $^{232}$Th, $^{235}$U and $^{238}$U, have their half-lives roughly in the order of the age of the Earth. Primordial nuclides; $^{232}$Th, $^{235}$U, $^{238}$U are particularly important. Gamma radiation from radionuclides, such as $^{40}$K and the $^{232}$Th and $^{238}$U series and their decay products, represents the main external source of irradiation to the human body. External exposures to gamma radiation outdoors arise from terrestrial radionuclides occurring at trace levels in all rock formations. Therefore, the natural environmental radiation mainly depends on geological and geographical conditions [2]. These have direct effect by modifying the soil composition and natural radioactivity concentration levels; and hence the level of absorbed dose received at a locality. Higher radiation levels are associated with igneous rocks, such as granite and lower levels with sedimentary rocks. There are exceptions, however, as some shale and phosphate rocks have relatively high content of radionuclides [3].

The great interest expressed worldwide for the study of naturally occurring radiation and environmental radioactivity has led to the performance of extensive research in many countries; such as Cyprus, Syria, Greece and Japan. Such investigations can be useful for both assessment of public dose and performance of epidemiological studies, as well as to keep reference-data records, to ascertain possible changes in the environmental radioactivity due to nuclear, industrial and other human activities. The total amount of radioactivity in an environment should be accurately known and kept at a level as Low...
AS Reasonably Achievable (ALARA). Both laboratory and in-situ gamma spectroscopy are often used for such studies [4,5,6,7 and 8]. Environmental radioactivity monitoring is believed to have commenced between late 1959 and early 1960s [9]. Some specific localized studies on radioactivity assessment in Nigerian environment include; radionuclide content of some building materials used in Nigerian dwellings, baseline studies of terrestrial outdoor gamma dose rates levels in Nigeria [10], exposure level around industrial area [11,12].

The principal aim of this work is to develop a systematic procedure using high-resolution portable gamma spectrometry system, primarily for assessment and monitoring, of absorbed dose rates and effective dose in air, in the vicinity of nuclear facility. The present work has been conducted in the Centre for Energy Research and Training (CERT), Zaria, Nigeria and some selected settlements around the facility. CERT is a nuclear energy (radiation) based research centre, thus, dealing with substantial quantities of artificial radioactive materials, such as a nuclear research reactor code named Nigeria Research Reactor-1 NIRR-1; Am-Be isotopic neutron source, neutron generator among others.

**MATERIALS AND METHODS**

**Gamma spectrometry system**

This is a stand-alone high-resolution spectroscopic system used for the in-situ measurement of the emitted gamma rays in the energy range between 50 keV and 3000 keV. The system consists of a high-purity germanium (HPGe) detector, EG and G ORTEC®, coaxial cylinder of crystal length of 46.8 mm and diameter 55.0 mm, with an efficiency of 23.5% relative to a 7.6 × 7.6 cm² NaI(Tl) crystal. In addition, energy resolution (FWHM) of 2.0 keV was achieved, all for a $^{60}$Co emission point source at 25 cm for 1.33 MeV. This type of detector can sustain warm up when not in use, which is a convenient feature during extended field trips. The detector was mounted on a portable (hand-held) 10 litres liquid nitrogen dewar that features an all attitude capability. Liquid nitrogen was used for cooling the detector during operation. After filling the portable dewar with liquid nitrogen, it requires a 6 hour cool-down time before becoming operational with 24 h nominal holding time. The detector assembly was mounted on a 1 m tripod with the crystal end cap facing down towards the ground. This orientation maximizes the flux that will be intercepted and registered by the detector [13]. The detector unit was connected to a battery powered EG and G ORTEC® “Normad Plus” portable computer based spectroscopy system. High voltage and preamplifier power were supplied to the detector by the system. Advanced Multi-Channel Analyzer (MCA) emulsion software (MAEASTRO-32) was used for data acquisition, storage display and analysis of the acquired gamma-spectra.

**Field measurement and analysis**

The study site is located at 007°38.523’– 007°40.822 E and 11°07.83’ – 11°09.790’ N within the Zaria sheet 102. The number of monitoring points (MPs) includes; seven locations within the Centre for Energy Research and Training (CERT), Ahmadu Bello University, Zaria and three other locations at selected settlements around CERT within 2Km radius. The chosen sites were undisturbed with little or no surface features and modest vegetation. These sites were marked-out using Global Positioning System (GPS).

The field measurement of terrestrial gamma radiation was based on the assumption that there exist laterally uniform distribution of natural radionuclides in the environment and that the vertical contribution from the soil is limited to the first horizon ($≈ 10-30$ cm). Measurements were performed over flat terrain, which allows source geometry to be represented as an infinite half-space; that is $2\pi$ geometry in terms of solid angle subtended by the source.

Measurements of spectra in the field were made for a period of 5000s. However, series of random short readings of about 600s were first taken, to ensure that, there is approximate uniformity, consequently, the desired counting statistics. The measuring system was routinely checked with $^{137}$Cs
Based on two-point energy calibration as set for the operation, prominent peaks were identified and the appropriate regions of interest were set up. The set energy bands define the peaks where the left and right channel markers are representative of the Compton continuum. Detector’s specific calibration factors (measured efficiency) determined in an earlier experiment [14], were applied to convert net peak count rate to activity concentration. Only peaks with reasonable gamma-ray emission probabilities were considered.

Since natural radioactive nuclides are assumed to be uniformly distributed in the ground, the dose rates (nGy h⁻¹) at ~1 m above the ground were calculated by the following formula:

\[ D = A_E \times C_F \]  (1)

Where:
- \( A_E \) = The activity concentration measured in Bq Kg⁻¹,
- \( C_F \) = The conversion factor (absorbed dose rate in air per unit activity per unit of soil mass, in units of nGy h⁻¹ per Bq Kg⁻¹)

In the present study, the considered Dose Rate Conversion Factors (DRCF) for the \(^{232}\text{Th}\) and \(^{238}\text{U}\) series and \(^{40}\text{K}\) in the dose rate calculations were those determined by Saito et al. [14] which had been used extensively for all similar calculations in the UNSCEAR [15] report. They were multiplied by the measured activity concentration in order to deduce the dose rate due to the entire series. It should be pointed out here that, using this calculation, the dose rate for the \(^{232}\text{Th}\) and \(^{238}\text{U}\) series is the average of the respective radiological concentrations multiplied by the conversion factors corresponding to each series. The total dose rate for each of the measured samples is the sum of the dose rate contributed by both series of \(^{232}\text{Th}\) and \(^{238}\text{U}\) and by \(^{40}\text{K}\).

Finally, the effective doses were estimated using Eq. 2. The effective dose outdoor \( H_E \) in unit of Sv y⁻¹ is given by:

\[ H_E = DTF \]  (2)

Where:
- \( D \) = The calculated dose rate (nGy h⁻¹) \( \times (24 \text{ h} \times 365.25 \text{d}) \)
- \( T \) = The occupancy factor (0.2)
- \( F \) = Conversion coefficient (0.7 Sv Gy⁻¹ \( \times 10^{33} \))

Taking into account the conversion coefficient from absorbed dose in air to effective dose and the outdoor occupancy factor, a value of 0.7 Sv Gy⁻¹ was used for the conversion coefficient from absorbed dose in air to effective dose received by adults and 0.2 for the outdoor occupancy factor UNSCEAR [15].

RESULTS AND DISCUSSION

The measured absorbed gamma dose rates in air at 1 m above ground level due to activity concentrations of \(^{232}\text{Th}\) and \(^{238}\text{U}\) series and \(^{40}\text{K}\) are presented in Table 1. These values range from 8-25 nGy h⁻¹, 2-5 nGy h⁻¹, 12-38 nGy h⁻¹ respectively. The total absorbed dose rates range from 22-65 nGy h⁻¹ with a mean of 46.2±12.5 nGy h⁻¹. According to UNSCEAR [15],

![Fig. 1: \(^{232}\text{Th}, \ 238\text{U}, \ 40\text{K}, \ \text{percentage contribution to the total dose rate in air outdoor for the ten monitoring points}](image)

\[ \text{Fig. 1: } ^{232}\text{Th}, \ 238\text{U}, \ 40\text{K}, \ \text{percentage contribution to the total dose rate in air outdoor for the ten monitoring points} \]
Fig. 2: Estimated effective dose rate due to $^{238}\text{Th}$, $^{238}\text{U}$, $^{40}\text{K}$, the error bar represent the uncertainty

The corresponding worldwide average values range from 18-93 nGy h$^{-1}$ and the population-weighted values gave an average absorbed dose rate in air outdoors from terrestrial gamma radiation as 60 nGy h$^{-1}$. This shows that the mean value obtained by this study is less than population-weighted worldwide average.

Of all the locations MP004 and MP011 appear to have the highest dose rate due to $^{232}\text{Th}$, also MP004 exhibits the highest dose rate of $^{238}\text{U}$, while the highest dose rate of 38 nGy h$^{-1}$ due to $^{40}\text{K}$ was recorded at MP012. The contributions in percent of $^{232}\text{Th}$ and $^{238}\text{U}$ series and $^{40}\text{K}$ at each measurement location are presented in Fig. 1. These contributions to absorbed gamma dose rate of the $^{232}\text{Th}$ and $^{238}\text{U}$ series and $^{40}\text{K}$ range from 29.8% at MP001 to 46.3% at MP011, 4.3% at MP012 to 9.0% at MP111 and 45.9% at MP011 to 62.5% at MP001 with a mean contributions of 39.3, 7.3 and 57.3% respectively. The estimated outdoor effective dose according to equation 2 for the sites as presented in Fig. 2 range from 27.3-79.9 µSv y$^{-1}$ with a mean of 0.57 mSv y$^{-1}$ which is significantly lower than the world-wide average exposure of 0.7 mSv y$^{-1}$ [15].

Table 1: Global Positioning System (GPS) coordinates and the measured absorbed gamma dose rates in air at 1 m above ground level due to activity concentrations of $^{232}\text{Th}$ and $^{238}\text{U}$ series and $^{40}\text{K}$ for the monitoring points

<table>
<thead>
<tr>
<th>Location</th>
<th>$^{232}\text{Th}$ series</th>
<th>$^{238}\text{U}$ series</th>
<th>$^{40}\text{K}$ Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Site ID</td>
<td>(nGy.h$^{-1}$)</td>
<td>(nGy.h$^{-1}$)</td>
<td>(nGy.h$^{-1}$)</td>
</tr>
<tr>
<td>MP001</td>
<td>11°08.439' N</td>
<td>07°39.840' E</td>
<td>11.2±2.9</td>
</tr>
<tr>
<td>MP003</td>
<td>11°08.476' N</td>
<td>07°39.901' E</td>
<td>19.8±2.8</td>
</tr>
<tr>
<td>MP004</td>
<td>11°08.497' N</td>
<td>07°39.851' E</td>
<td>24.5±3.6</td>
</tr>
<tr>
<td>MP011</td>
<td>11°08.415' N</td>
<td>07°39.776' E</td>
<td>24.4±5.5</td>
</tr>
<tr>
<td>MP012</td>
<td>11°08.358' N</td>
<td>07°39.912' E</td>
<td>24.2±4.6</td>
</tr>
<tr>
<td>MP013</td>
<td>11°08.505' N</td>
<td>07°39.963' E</td>
<td>17±5.7</td>
</tr>
<tr>
<td>MP014</td>
<td>11°08.549' N</td>
<td>07°39.823' E</td>
<td>16.3±3.8</td>
</tr>
<tr>
<td>MP111</td>
<td>11°07.830' N</td>
<td>07°39.840' E</td>
<td>16.3±3.8</td>
</tr>
<tr>
<td>MP112</td>
<td>11°08.749' N</td>
<td>07°40.822' E</td>
<td>8.2±2.5</td>
</tr>
</tbody>
</table>

Thus, the exposure level for the public is within the recommended value of 1 mSv y$^{-1}$ (IAEA Safety Series No. 115-I). The highest and lowest effective doses of 79.9 and 27.3 µSv y$^{-1}$ were registered at MP012 and MP113 respectively.
CONCLUSION
In this study portable gamma ray spectroscopy was exploited to develop a systematic procedure for rapid and effective determination of outdoor absorbed and effective dose in a vicinity of nuclear facility. The in situ results obtained by the present work would serve as a baseline data for assessment and operational monitoring of outdoor exposure around Centre for Energy Research and Training (CERT). Values obtained were lower than the population-weighted worldwide average.

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REFERENCES