Natural radioactivity levels in soils, rocks and water at a mining concession of Perseus gold mine and surrounding towns in Central Region of Ghana

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Abstract
Levels of naturally occurring radioactive materials prior to processing of gold ore within and around the new eastern concession area of Perseus Mining (Ghana) Limited were carried out to ascertain the baseline radioactivity levels. The study was based on situ measurements of external gamma dose rate at 1 m above ground level as well as laboratory analysis by direct gamma spectrometry to quantify the radionuclides of interest namely, $^{238}$U, $^{232}$Th and $^{40}$K in soil, rock, ore samples and gross alpha/beta analysis in water samples. The average absorbed dose rate in air at 1 m above sampling point using a radiation survey metre was determined to be 0.08 ± 0.02 μGy h$^{-1}$ with a corresponding average annual effective dose calculated to be 0.093 ± 0.028 mSv. The average activity concentrations of $^{238}$U, $^{232}$Th, and $^{40}$K in the soil, rock, and ore samples were 65.1 ± 2.2, 71.8 ± 2.2 and 1168.3 Bq kg$^{-1}$ respectively resulting in an average annual effective dose of 0.91 ± 0.32 mSv. The average Radium equivalent activity value was 257.8 ± 62.4 Bq kg$^{-1}$ in the range of 136.6–340.2 Bq kg$^{-1}$. The average values of external and internal indices were 0.7 ± 0.2 and 0.9 ± 0.2 respectively. The average gross alpha and gross beta activity concentrations in the water samples were determined to be 0.0032 ± 0.0024 and 0.0338 ± 0.0083 Bq l$^{-1}$ respectively. The total annual effective dose from the pathways considered for this study (gamma ray from the soil, rock and ore samples as well as doses determined from the gross alpha/beta activity concentration in water samples) was calculated to be 0.918 mSv. The results obtained in this study shows that the radiation levels are within the natural background radiation levels found in literature and compare well with similar studies for other countries and the total annual effective dose is below the ICRP recommended level of 1 mSv for public exposure control.

Keywords: Natural radioactivity, Gold mine, Gamma spectrometry, Gross alpha, Gross beta, Annual effective

Background
Artificial and natural radioactivities are the two main sources of radiation exposure. Human activities such as mining and mining processes, oil and gas extraction may result in situations where the radioactivity levels of materials that contain natural radionuclides...
are significant enough to warrant regulatory control (UNSCEAR 2000; IAEA 2005). The
two main sources of natural radiation exposure are from terrestrial radionuclides and
cosmogenic radionuclides. They lead to external and internal radiation exposure. This
study focused on levels of radioactivity due terrestrial radionuclides. The radionuclides
of interest are Uranium-238 and Thorium-232 and their decay series nuclides as well as
Potassium-40.

The major sources of external gamma radiation are due to $^{238}\text{U}$, $^{232}\text{Th}$ and their decay
products and $^{40}\text{K}$. $^{238}\text{U}$ and its daughters rather than $^{226}\text{Ra}$ and its daughter products are
responsible for the major fraction of the internal dose received by humans from naturally occurring radionuclides (de Oliveira et al. 2001). Even though the concentrations of
these radionuclides are widely distributed in nature, they have been found to depend
on the local geological conditions and as a result vary from place to place (Xinwei et al.
2006). The specific levels are related to the type of rock from which the soil originates.
Higher radioactivity levels are associated with igneous rocks such as granite and lower
levels with sedimentary rocks. There are exceptions however, as some shales and phos-
phate rocks have relatively high content of radionuclides (Uosif 2007).

Among the natural radionuclides, alpha and beta emitters are considered the most
important with respect to potential internal radiation exposure to humans particularly
through ingestion of food and water (UNSCEAR 2000) as approximately 20 % of Ra iso-
topes and 10–15 % of the Pb isotopes which are decay products of $^{238}\text{U}$ and $^{232}\text{Th}$ con-
sidered for this work reaches the blood stream and are distributed to the whole body and
follows the same metabolism as calcium (UNSCEAR 1988). The percentage distribution
of annual intakes of Uranium and Thorium series radionuclides in diet ranges between
4 and about 96 % (UNSCEAR 2000), as a result accumulation of these radionuclides
through the ingestion have significant health effects such as bone cancer, leukemia and
increase in blood pressure (Tettey-Larbi et al. 2013).

Mining has been identified as one of the potential sources of exposure to naturally
occurring radioactive materials (UNSCEAR 2000). As a result it is necessary for base-
line studies to be carried out prior to the commencement of mining activity and subse-
quently similar studies be conducted to ascertain the levels of these radionuclides being
turned out as a result of the mining activities in the operational phase of the mine.

Perseus Mining (Ghana) Ltd., is one of the several mining companies in Ghana located
at Ayanfuri, of Ghana. Earlier study by Faanu et al. (2013, 2014), on part of the mine's
concession gave an average activity concentrations for $^{238}\text{U}$, $^{232}\text{Th}$ and $^{40}\text{K}$ as 64.3, 68.4
and 1243.9 Bqkg$^{-1}$ respectively. This study became necessary as a result of operational
expansion to another part of the concession, hence, the need to carried baseline radio-
activity survey in order to establish the levels of radioactivity in the area before com-
 mencement. This will also help to determine whether or not the operational expansion
of the mines to these new concession it operations will have added or raised the levels
of NORMs in and round the study area in a future operation and post operation of the
facility. The new concession is dotted along and within five main towns namely; Gyaman,
Odumkrom, Wampem, Ayanfuri and Nkonya which could be affected as a result of the
mine's operations.

Therefore, the main objective of this study was to measure and assess the baseline radi-
oactivity levels at the new concession of the mine as well as the immediate surroundings
so that reference data could be established before the mine starts processing of the gold ore in the area. The study focused on determination of activity concentration and distribution of naturally occurring radionuclides of U/Th decay series and ⁴⁰K in soil, rock, and ore samples by gamma spectrometry and gross alpha and gross beta activities in water samples. For the gross alpha and gross beta activities in the water samples, particular attention was focused on the major alpha and beta emitting radionuclides in the Uranium and Thorium Series which are of importance to internal irradiation. Lasheen et al. (2008), recognizes ²³⁸U, ²³⁴U, ²³⁰Th, ²²⁶Ra, ²¹⁰Pb and ²¹⁰Po for the Uranium series and ²³²Th, ²²⁸Ra and ²²⁸Th for the Thorium series as the major alpha and beta emitting radionuclides respectively.

**Description and local geology of study area**

The study area is the eastern concession of Perseus Mining (Ghana) Limited and its immediate surroundings up to about 20 km from the plant site. The mine is bounded by two districts namely; the Upper Denkyira East and Amenfi West districts. The goldmine is located at latitude 5°57′21.45″N and longitude 1°54′35.35″W. Within this concession, six gold deposits open pits namely; Chirawewa, Bokiti, Small Fetish, Big Fetish, Esuajah North and Esuajah South will be mined. Surrounding these open pits are the following towns: Gyaman, Odumkrom, Wampem, Ayanfuri and Nkonya.

The gold ore in the area occurs both in classic Ashanti-style sediment of shear zones and with granitic plugs and sills or dykes situated along two or three regional shear structures. In excess of 24 gold occurrences exist in the Ayanfuri property of which granitic intrusive host majority of these and more than 80 % of the known gold resource. While the later deposits formed predominantly ductile regime with generally discontinuous, pinch and swell higher grade gold shoots, the granite hosted occurrences developed in a brittle rock and found to be significantly broader with more evenly distributed, though lower grade gold tenor. Figure 1 is the geological map of the study area with sampling points in red dots.

**Methods**

**Sampling and sample preparation for gamma spectrometry analysis**

A total of 30 samples were randomly collected within selected areas of the new concession and the surrounding communities. They included 14 soil, rock and ore samples and 16 water samples.

In the laboratory, each of the soil, rock and ore samples were air dried on trays for 7 days and then oven dried at a temperature of 105 °C for between 3 and 4 h until all moisture was completely lost. The dried samples were then grinded into fine powder using a stainless steel ball mill and sieved through a 2 mm mesh size and poured into 1 l Marinelli beakers. The Marinelli beakers were hermetically sealed, weighed and stored for 4 weeks, to allow the short-lived daughters of ²³⁸U and ²³²Th decay series to attain equilibrium with their long-lived parent radionuclides (ASTM 1983, 1986). The soil samples were each counted using a sodium iodide detector for a period of 72,000 s (20 h).

For the water samples, 500 ml of each of the water samples was acidified with 1 ml of concentrated HNO₃ and evaporated to near dryness on a hot plate in a fume hood. The residue in the beaker was rinsed with 1 M HNO₃ and evaporated again to near dryness.
The residue was dissolved in minimum amount of 1 M HNO₃ and transferred into a weighed 25 mm stainless steel planchet. The planchet with its content was heated until all moisture had evaporated. It was then stored in a desiccator and allowed to cool and prevented from absorbing moisture. The water samples were each counted using the low background Gas-less Automatic Alpha/Beta counting system (Canberra iMatic™) for a period of 12,000 s (200 min or 3.33 h).

Instrumentation and calibration
The soil, rock and ore samples measurement were made by direct non-destructive instrumental analysis with a computerized gamma spectrometry system made up of NaI(Tl) detector and measuring assembly. The specifications of the detector system
used for this study are as follows: cylindrical scintillation detector Model 3M3/3-X, Serial number ETI 9305 having a 1.2" × 1.2" end window, manufactured by Saint-Gobain Crystals, USA. The detector system consists of a vertically sealed assembly which includes the NaI(Tl) crystal and is coupled to ORTEC Multichannel Buffers (MCBs) for data acquisition and processing using a MAESTRO®-32 software program. A high voltage supply provides the appropriate bias to the detector system. The conversion gain of the detector is up to 1024 channels. In order to reduce background gamma radiation from the room in which the detector is installed in, a locally fabricated cylindrical lead shield (20 mm thickness) with a fixed bottom and a movable lid to shield the detector. Within the lead shield are also copper, cadmium and plexiglass (3 mm thickness each) to absorb X-rays and other photons that might be produced in the lead. The ambient temperature around the detector varied between 20 and 27 °C during the period of measurement. The identification of individual radionuclides was performed using their gamma ray energies and the quantitative analysis of radionuclides was performed using gamma ray spectrum analysis software, ORTEC MAESTRO-32. Before the analysis of soil/rock-ore samples, energy and efficiency calibration were carried out using liquid mixed standard radionuclide solution supplied by the IAEA with volume and density of 1000 ml and 1.0 g cm⁻³ respectively. The energy and efficiency calibrations were carried out by counting standard radionuclides of known activities with well defined energies in the energy range of 60 to ~2000 keV. The background spectrum was also used to determine the minimum detectable activities of ²³⁸U (0.34 Bq), ²³²Th (0.33 Bq), and ⁴⁰K (1.62 Bq) at the 95 % confidence level. The low background gas-less automatic alpha/beta counting system (Canberra iMatic™) calibrated with alpha (²⁴¹Am) and beta (⁹⁰Sr) standards was used to count the prepared water samples. The system uses a solid state passivated implanted planar silicon (PIPS) detector for alpha and beta detection. The alpha and beta efficiencies were determined to be 36.39 ± 2.1 and 36.61 ± 2.2 % respectively. The background readings of the detector for alpha and beta activity concentrations were respectively 0.04 ± 0.01 and 0.22 ± 0.03 cpm.

Calculation of activity concentration and estimation of doses

**Soil, rock and ore samples**

The activity concentration of ²³⁸U was calculated from the average peak energies of 351.92 keV of ²¹⁴Pb, and 609.31 keV of ²¹⁴Bi. Similarly, the activity concentration of ²³²Th was determined from the average energies of 238 keV of ²¹²Pb and 911.21 keV of ²²⁸Ac. The activity concentration of ⁴⁰K was determined from the energy of 1460.83 keV. The analytical expression used in the calculation of the activity concentrations in Bq kg⁻¹ is as shown in Eq. (1) (Ebaid 2010)

\[
A_{sp} = \frac{N_D}{p \cdot T_c \cdot \eta(E) \cdot m}
\]

where \(N_D\) is the net counts of the radionuclide in the samples, \(p\) is the gamma ray emission probability (gamma ray yield), \(\eta(E)\) is the absolute counting efficiency of the detector system, \(T_c\) is the sample counting time, \(m\) is the mass of the sample (kg).
The external gamma dose rate ($D_{\gamma}$) at 1.0 m above ground for the soil/rock-ore pad-ore pad samples was calculated from the activity concentrations using Eq. (2) (Uosif 2007).

$$D_{\gamma} \text{ (nGy}^{-1} \text{h}) = DCF_K \times A_K + DCF_U \times A_U + DCF_{Th} \times A_{Th}$$

(2)

where $DCF_K$, $DCF_U$, $DCF_{Th}$ are the absorbed dose rate conversion factors for $^{40}$K, $^{238}$U and $^{232}$Th in nGy h$^{-1}$ Bq kg$^{-1}$ and $A_K$, $A_U$ and $A_{Th}$ are the activity concentrations for $^{40}$K, $^{238}$U and $^{232}$Th respectively.

$$DCF_K = 0.0417 \frac{\text{nGy}}{\text{h}} \frac{1}{\text{Bq kg}^{-1}}; \quad DCF_U = 0.462 \frac{\text{nGy}}{\text{h}} \frac{1}{\text{Bq kg}^{-1}}; \quad DCF_{Th} = 0.604 \frac{\text{nGy}}{\text{h}} \frac{1}{\text{Bq kg}^{-1}}$$

The average annual effective dose was calculated from the absorbed dose rate by applying the dose conversion factor of 0.7 Sv Gy$^{-1}$ and an outdoor occupancy factor of 0.2 (UNSCEAR 2000) represented by Eq. (3)

$$E_{\gamma} = D_{\gamma} \times 0.2 \times 8760 \times 0.7$$

(3)

where $E_{\gamma}$ is the average annual effective dose and $D_{\gamma}$ is the absorbed dose rate in air.

For comparative analysis, similar average outdoor external gamma dose rate were determined by taking an average of five measurements of the ambient gamma dose rates at 1 m above the ground of sampling in μGy h$^{-1}$ with a radiation survey meter. The annual effective dose ($E_{\gamma, ext}$) was then estimated from the measured average outdoor external gamma dose rate from the Eq. (4)

$$E_{\gamma, ext} = D_{\gamma, ext} T_{exp} DCF_{ext}$$

(4)

where $D_{\gamma, ext}$ is the average outdoor external gamma dose rate μGy$^{-1}$, $T_{exp}$ is the exposure duration per year, 8760 h and applying an outdoor occupancy factor of 0.2, $DCF_{ext}$ is the effective dose to absorbed dose conversion factor of 0.7 Sv Gy$^{-1}$ for environmental exposure to gamma rays (UNSCEAR 2000).

The radiological risk of NORM in soils in the study area which may be used as building materials was assessed by calculating the radium equivalent activity ($Ra_{eq}$) and the external hazard and internal hazard indices. The $Ra_{eq}$ is a widely used hazard index and it was determined using Eq. (5) (Xinwei et al. 2006):

$$Ra_{eq} = C_{Ra} + 1.43C_{Th} + 0.077C_K$$

(5)

where $C_{Ra}$, $C_{Th}$ and $C_K$ are the activity concentrations of $^{226}$Ra, $^{232}$Th and $^{40}$K respectively. In its application, the definition of $Ra_{eq}$ it is assumed that 370 Bq kg$^{-1}$ of $^{226}$Ra, 259 Bq kg$^{-1}$ of $^{232}$Th and 4810 Bq kg$^{-1}$ of $^{40}$K produce the same gamma ray dose rate. The above criterion only considers the external hazard due to gamma rays in building materials.

Another criterion used to estimate the level of gamma ray radiation associated with natural radionuclides in specific construction materials is defined by the terms External hazard index ($H_{ext}$) and Internal hazard index ($H_{in}$) as shown in Eqs. (6) and (7) (OECD/NEA 1979; Beretka and Mathew 1985; Alam et al. 1999; Higgy et al. 2000)

$$H_{ext} = \frac{C_{Ra}}{370} + \frac{C_{Th}}{259} + \frac{C_K}{4810}$$

(6)
where \( C_{Ra} \), \( C_{Th} \) and \( C_{K} \) are the activity concentrations of \(^{226}\text{Ra},^{232}\text{Th}\) and \(^{40}\text{K}\) respectively. The value of the external hazard index must be less than unity for the external gamma radiation hazard to be considered negligible.

The internal hazard index \( (H_{in}) \) due to Radon and its daughters were calculated from Eq. (7). This is based on the fact that, Radon and its short-lived products are also hazardous to the respiratory organs

\[
H_{in} = \frac{C_{Ra}}{185} + \frac{C_{Th}}{259} + \frac{C_{K}}{4810}
\]

where \( C_{Ra} \), \( C_{Th} \) and \( C_{K} \) are the activity concentrations of \(^{226}\text{Ra},^{232}\text{Th}\) and \(^{40}\text{K}\) respectively. For construction materials to be considered safe for construction of dwellings, the internal hazard index should be less than unity.

**Water samples**

The activity concentration of both gross alpha and gross beta was determined using the expression in Eq. (8) (Tettey-Larbi et al. 2013):

\[
A_{\alpha/\beta} = \frac{\text{Activity}}{W_{Vol}}
\]

where \( A_{\alpha/\beta} \) is the activity concentration of gross alpha or gross beta in Bq l\(^{-1}\) and \( W_{Vol} \) is the volume of the water sample in litres. The activity of alpha or beta in Bq was obtained by subtracting the background activity of both gross alpha and gross beta from the total activity of the sample. The average annual alpha or beta committed effective dose for a particular water sample was determined by averaging the individual annual committed effective doses contributed by the major alpha or beta emitters in the \(^{238}\text{U}\) and \(^{232}\text{Th}\) series of the naturally occurring radionuclides as shown in Eq. (9) (Tettey-Larbi et al. 2013):

\[
E_{\text{ing},w}(\alpha/\beta) = \frac{I_{w}}{W_{Vol}} \cdot N_{R}(\alpha/\beta) \sum_{i} A_{\alpha/\beta} \times DCF_{\text{ing}}(\alpha/\beta)
\]

where \( E_{\text{ing},w}(\alpha/\beta) \) is the average gross annual alpha or beta committed effective dose in the water sample, \( A_{\alpha/\beta} \) is the gross alpha or beta activity concentration in the water sample in Bq l\(^{-1}\), \( I_{w} \) is the consumption rate for the intake of the water of 730 l year\(^{-1}\) (WHO 2004), \( W_{Vol} \) is the volume water used for the analysis, \( N_{R}(\alpha/\beta) \) is the number of radionuclides considered as major alpha or major beta emitters in the \(^{238}\text{U}\) and \(^{232}\text{Th}\) series of the naturally occurring radionuclides and \( DCF_{\text{ing}}(\alpha/\beta) \) is the ingestion dose coefficient in Sv/Bq of the natural radionuclides from UNSCEAR report (UNSCEAR 2000).

**Total annual effective dose**

The total annual effective dose \( (E_{T}) \) to members of the public was calculated using ICRP dose calculation method (ICRP 1991, 2007). The analytical expression for the total effective dose is provided in Eq. (10)

\[
E_{T} = E_{\gamma}(U, Th, K) + E_{\text{ing},w}(\alpha/\beta)
\]
where $E_T$ is the total effective dose in Sievert (Sv), $E_\gamma$ (U, Th, K) is the external gamma effective dose from the soil/rock-ore pad samples, $E_{\text{ing}, w}$ ($\alpha/\beta$) is the effective dose from the consumption of water due to gross alpha and gross beta activity concentrations.

### Results and discussion

The list of sample location index with coordinates is given in Table 1 along with remarks for soil, water and ore, respectively.

Table 2 shows the absorbed dose rate measured in air at 1 m above the ground at the soil/rock-ore pad and water sampling points in the study area and its surrounding communities. The measured average absorbed dose rate was determined to be $0.08 \pm 0.02 \, \mu \text{Gy h}^{-1}$ ($80 \pm 20 \, \text{nGy h}^{-1}$). The corresponding average annual effective dose was calculated to be $0.093 \pm 0.028 \, \text{mSv}$ ($93 \pm 28 \, \mu \text{Sv}$) in a range of $0.047–0.142 \, \text{mSv}$ ($49–142 \, \mu \text{Sv}$). According to UNSCEAR (2000) report, the worldwide average absorbed dose rate in air measured from terrestrial gamma radiation is $60 \, \text{nGy h}^{-1}$ ($0.059 \, \mu \text{Gy h}^{-1}$). By comparison, the results of the absorbed dose rates in this study compare well with the range of dose rates values reported for other countries (UNSCEAR 2000) as well as results from similar studies carried out in other mines in Ghana (Darko et al. 2010;
although the average absorbed dose rate of 80 ± 20 nGyh$^{-1}$ measured in air from the study area is above the worldwide average. The reasons for the higher values of the doses for external gamma could be due to difference in geological formations as well as contribution from cosmogenic radionuclides in addition to terrestrial radionuclides as observably in Fig. 2 the pits give the highest average effective dose although the maximum value was recorded around the streams.

Table 3 shows the activity concentrations of $^{238}$U, $^{232}$Th and $^{40}$K in the soil/rock-ore pad samples. The average value of the activity concentrations of $^{238}$U is 65.1 ± 2.2 Bqkg$^{-1}$ in a range of 29.0–97.0 Bqkg$^{-1}$. For $^{232}$Th, the average activity concentration is 71.8 ± 2.2 Bqkg$^{-1}$ in range of 35.0–116.7 Bqkg$^{-1}$ and that of $^{40}$K is 1168.3 ± 15.8 Bqkg$^{-1}$ in a range of 500.0–1795.9 Bqkg$^{-1}$. The results of this study also compare with the previous study that was carried in Perseus Mining (Ghana) Ltd., (Fannu et al. 2013, 2014). As shown in Table 3, the average values of the activity concentrations of $^{238}$U and $^{232}$Th in this study are about two times higher than the world average whilst that of $^{40}$K is about three times higher than values in normal continental soils (UNSCEAR 2000). The high activity concentration of $^{40}$K is because the rock ore of the mine is associated feldspar which belongs to a group of hard crystalline minerals that consist of aluminium silicates of potassium, sodium, calcium or barium. Even though the average values in this study are higher than the worldwide average values, activity concentrations are still far below the exemption values of 1000 Bqkg$^{-1}$ for $^{238}$U and $^{232}$Th and 100,000 Bqkg$^{-1}$ for $^{40}$K in materials that will warrant regulatory control (IAEA 1996).

A comparison of the activity concentrations of the radionuclides in soil and rock-ore pad with exemption levels recommended in the Basic Safety Standards (IAEA 1996) are
Airborn Annual effec/g415ve dose  (mSv/yr)

| Sampling Site         | Min  | Max  | Average ± σ |
|-----------------------|------|------|--------------|
| PITTS                 | 0.04 | 0.16 | 0.09 ± 0.02  |
| SOURROUNDING TOWNS    | 0.04 | 0.12 | 0.08 ± 0.02  |
| SOURROUNDING STREAMS  | 0.04 | 0.12 | 0.08 ± 0.02  |

Fig. 2 Comparison of annual effective doses from different sampling site due to airborne gamma exposure pathways of radiation

Table 3 Average activity concentrations of $^{238}$U, $^{232}$Th and $^{40}$K in soil, rock and ore pad samples in the study area

| Sample location | Activity concentration (Bq kg$^{-1}$) |
|-----------------|--------------------------------------|
|                 | $^{238}$U  | $^{232}$Th | $^{40}$K   |
| P1              | 97.0 ± 2.4 | 70.8 ± 2.4 | 1795.9 ± 17.8 |
| P2              | 72.0 ± 2.3 | 84.2 ± 2.2 | 1445.4 ± 16.3 |
| P3              | 86.8 ± 2.1 | 64.4 ± 2.1 | 1317.2 ± 17.5 |
| P4              | 56.3 ± 2.0 | 61.8 ± 2.2 | 1722.4 ± 14.6 |
| P5              | 62.2 ± 3.8 | 116.7 ± 3.0 | 1445.0 ± 20.4 |
| P6              | 74.3 ± 2.3 | 89.6 ± 2.4 | 1544.1 ± 16.8 |
| Q1              | 62.1 ± 2.1 | 61.6 ± 2.2 | 1470.6 ± 17.2 |
| Q2              | 29.0 ± 1.6 | 35.0 ± 1.6 | 748.3 ± 12.0 |
| S1              | 66.4 ± 1.9 | 81.6 ± 2.1 | 1241.2 ± 15.0 |
| T1              | 69.2 ± 2.6 | 76.1 ± 2.1 | 844.8 ± 14.9 |
| T2              | 70.0 ± 1.9 | 83.7 ± 2.0 | 720.8 ± 14.6 |
| T3              | 57.0 ± 2.1 | 63.9 ± 1.9 | 500.0 ± 13.9 |
| T4              | 61.3 ± 2.1 | 68.7 ± 2.0 | 749.5 ± 14.6 |
| T5              | 48.4 ± 1.9 | 47.1 ± 2.1 | 811.3 ± 15.1 |
| Range           | 29.0–97.0 | 35.0–116.7 | 500.0–1795.9 |
| Average ± σ     | 65.1 ± 2.2 | 71.8 ± 2.2 | 1168.3 ± 15.8 |
| World average   | 35        | 30        | 400        |

σ—Standard deviation
The activity concentrations of the radionuclide in the different types of samples are quite uniform and do not show any significant variation.

The average gamma dose rate and annual effective dose from terrestrial gamma rays calculated from soil/rock-ore pad activity concentrations are shown in Table 4. The average absorbed dose rate was calculated to be 741.6 ± 260.1 nGy h⁻¹ with ranges which are by factors of 6–16 higher than the dose rate measured in air at 1 m above the ground. The average absorbed dose rate due to the soil concentrations is also about 12 times higher than the worldwide average value of 60 nGy h⁻¹ (UNSCEAR 1993, 2000). This difference could be attributed to vast differences in geology and geochemical state of the sampling sites. The corresponding average annual effective dose estimated from the soil concentrations is 0.91 ± 0.32 mSv.

Table 4 also shows the results of the hazard assessment of soil/rock-ore pad with respect to radium equivalent (Ra_eq) activity in Bq kg⁻¹, external hazard (H_ex) and the internal hazard (H_in) indices. Radiological hazard assessment of natural radioactivity in building materials is usually determined from the activity concentrations of 226Ra, 232Th and 40K. 238U is replaced by 226Ra because 98.5 % of the radiological hazard of uranium-series is due to radium and its decay products. The maximum value of Ra_eq in building materials must be <370 Bq kg⁻¹ for the material to be considered safe for use. The external and internal hazard indices must also be less than unity in order to keep the radiation hazard insignificant. This implies that, the average external radiation exposure due to the radioactivity from these radionuclides in materials to be used for constructions must be limited to 1.5 mSv year⁻¹ (OECD/NEA 1979; Beretka and Mathew 1985). The average value of the radium equivalent activity in this study is 257.8 ± 61.1 Bq kg⁻¹ in a range of 136.6–340.3 which is below the recommended limit of 370 Bq kg⁻¹. The calculated

![Fig. 3](image)
Hex in the soil/rock-ore pad samples ranged from 0.4 (Big Fetish Pit Ore pad) to 0.9 (soil sample from Chirawewa Pit and Esuajah North and South Pit) and an average value of 0.7 ± 0.2 for the study area. Similarly for the H in, the values ranged from 0.4 (Big Fetish Pit Ore pad) to 1.2 (Chirawewa Pit soil) with an average of 0.9 ± 0.2. For the internal hazard 21.4% of the samples had values exceeding the recommended limit of 1.0 which implies that these materials if used for building purposes could be a source of internal hazard due to Radon and its progeny. While 78.6% of the samples have values either exactly at the recommended limit or below the recommended limit. In general however, the annual effective doses calculated from the various samples are considered insignificant in respect to the annual construction limit of 1.5 mSv year\(^{-1}\).

The activity concentrations of gross-α and gross-β in water samples from the pits, surface water and underground water (bore holes) used in the surrounding communities of the study area and their corresponding committed effective dose are shown in Table 5. Radionuclide concentrations in groundwater depend on the dissolution of minerals from rock aquifers. The activity concentrations of gross-α in the water samples varied in a range of 0.0004 Bq l\(^{-1}\) in surface water at Wampem to 0.0075 Bq l\(^{-1}\) in surface water in Nkonya borehole with a corresponding average annual committed effective dose of 0.0007 ± 0.0005 mSv. For the gross-β, the activity concentrations varied in a range of 0.0104 Bq l\(^{-1}\) for water taken from a stream in Odumkrom to 0.0452 Bq l\(^{-1}\) for water from the raw underground water at the camp site with a correspond average annual committed effective dose of 0.0170 ± 0.0042 mSv. The average committed annual effective dose due to both gross alpha and beta was estimated to be 0.0089 ± 0.0023 mSv. All the water sources had gross-α and gross-β values below the WHO screening levels for drinking water at no further action is required. This indicates that all the water sources

| Sample location | Absorbed dose rate (nGy h\(^{-1}\)) | Radium equivalent activity (Bq kg\(^{-1}\)) | External hazard index | Internal hazard index | Annual effective dose (mSv) |
|-----------------|-----------------------------------|-------------------------------------------|----------------------|----------------------|---------------------------|
| P₁              | 1121.5                            | 336.5                                     | 0.9                  | 1.2                  | 1.38                      |
| P₂              | 914.9                             | 303.7                                     | 0.8                  | 1.0                  | 1.12                      |
| P₃              | 829.0                             | 280.2                                     | 0.8                  | 1.0                  | 1.02                      |
| P₄              | 1071.2                            | 277.4                                     | 0.7                  | 0.9                  | 1.31                      |
| P₅              | 929.3                             | 340.3                                     | 0.9                  | 1.1                  | 1.14                      |
| P₆              | 977.1                             | 321.3                                     | 0.9                  | 1.1                  | 1.20                      |
| Q₁              | 919.3                             | 263.3                                     | 0.7                  | 0.9                  | 1.13                      |
| Q₂              | 469.3                             | 136.6                                     | 0.4                  | 0.4                  | 0.58                      |
| S₁              | 790.2                             | 278.6                                     | 0.8                  | 0.9                  | 0.97                      |
| T₁              | 548.3                             | 243.1                                     | 0.7                  | 0.8                  | 0.67                      |
| T₂              | 477.0                             | 245.2                                     | 0.7                  | 0.9                  | 0.58                      |
| T₃              | 333.9                             | 187.0                                     | 0.5                  | 0.7                  | 0.41                      |
| T₄              | 487.0                             | 217.3                                     | 0.6                  | 0.8                  | 0.06                      |
| T₅              | 513.8                             | 178.2                                     | 0.5                  | 0.6                  | 0.63                      |
| Range           | 333.9–1121.5                      | 136.6–340.3                               | 0.4–0.9              | 0.4–1.2              | 0.41–1.38                 |
| Average ± σ     | 741.6 ± 260.1                     | 257.8 ± 61.1                              | 0.7 ± 0.2            | 0.9 ± 0.2            | 0.91 ± 0.32               |

σ—Standard deviation

\(H_{ex}\) in the soil/rock-ore pad samples ranged from 0.4 (Big Fetish Pit Ore pad) to 0.9 (soil sample from Chirawewa Pit and Esuajah North and South Pit) and an average value of 0.7 ± 0.2 for the study area. Similarly for the \(H_{in}\), the values ranged from 0.4 (Big Fetish Pit Ore pad) to 1.2 (Chirawewa Pit soil) with an average of 0.9 ± 0.2. For the internal hazard 21.4% of the samples had values exceeding the recommended limit of 1.0 which implies that these materials if used for building purposes could be a source of internal hazard due to Radon and its progeny. While 78.6% of the samples have values either exactly at the recommended limit or below the recommended limit. In general however, the annual effective doses calculated from the various samples are considered insignificant in respect to the annual construction limit of 1.5 mSv year\(^{-1}\).
in the study area which are designated for drinking and domestic purposes do not have significant natural radioactivity.

Figure 4 shows the comparison of the results of the annual effective doses due to $^{238}\text{U}$, $^{232}\text{Th}$ and $^{40}\text{K}$ calculated from the airborne gamma radiation and soil/rock-ore pad samples. Figure 4 shows that the annual effective doses due to $^{238}\text{U}$, $^{232}\text{Th}$ and $^{40}\text{K}$ in the soil/rock-ore pads were higher than that due to the external gamma dose rates. The reasons for the higher values of the doses for soil/rock-ore pads could be due to contribution from the high concentration of terrestrial radionuclides especially $^{40}\text{K}$ since is a purely gamma emitter. It also compares the total annual effective dose to the recommended limit of 1 mSv for the public.

**Conclusions**

The study considered public exposure in the mining environment due to two exposure pathways; namely direct external gamma ray exposure from natural radioactivity concentrations in soil/rock-ore pads radioactivity due to $^{238}\text{U}$, $^{232}\text{Th}$ and $^{40}\text{K}$ and internal exposure due to natural radioactivity by accessing the activity concentrations of gross-α and gross-β in water samples from the pits, surface water and underground water (bore holes) used in the surrounding communities of the study area.
The average activity concentrations of $^{238}\text{U}$, $^{232}\text{Th}$ and $^{40}\text{K}$ in the soil and rock-ore pad samples were calculated to be $65.1 \pm 2.2$, $71.8 \pm 2.2$ and $1168.3 \pm 15.8 \text{Bqkg}^{-1}$ respectively. The results in this study compared well with studies carried out in Ghana and other countries and with the worldwide average activity concentrations (UNSCEAR 2000; Darko et al. 2010; Faanu et al. 2010, 2013, 2014; Huang et al. 2014). The average annual effective doses estimated from direct external gamma-ray exposure from natural radioactivity at 1 m above sampling area and that due to activity concentrations of $^{238}\text{U}$, $^{232}\text{Th}$ and $^{40}\text{K}$ in the soil and rock-ore pad samples were $0.093 \pm 0.028$ and $0.91 \pm 0.32 \text{mSv}$ respectively. In order to assess whether the soil/rock-ore pad in the study area could be a source of public radiation exposure if used for building purposes, the following hazard indices were determined; radium equivalent ($\text{Ra}_{\text{eq}}$) activity in Bq/kg, external ($H_{\text{ex}}$) and the internal hazard ($H_{\text{in}}$). The average value of the radium equivalent activity in this study is $257.8 \pm 61.1 \text{Bqkg}^{-1}$ which is below the recommended limit of $370 \text{Bqkg}^{-1}$. The mean external and internal indices were $0.7 \pm 0.2$ and $0.9 \pm 0.2$ respectively where also below the recommended limit of 1. Therefore the annual effective doses calculated from the various samples are considered insignificant in respect to the annual construction limit of 1.5 mSv year$^{-1}$.

In general the results in this study are comparable to similar studies carried out previously at the mines and in other mines in Ghana (Darko et al. 2010; Faanu et al. 2010, 2013, 2014) as well as in other countries (UNSCEAR 2000; Huang et al. 2014). It also indicates insignificant levels of the natural radionuclides in the study area and therefore radiation exposure to workers as well as members of the public is not of any significant hazard and the level risk is considered generally insignificant. Finally it also implies that
previous mining activities had not imparted negatively in terms of radiological hazard to the communities in this area.

Authors' contributions
OKA, LT‑L and HL carried out the sample collection, field experimental works and prepared the manuscripts. CK, PAA, AOE, ADI, BA, RK and LA assisted in the samples preparation, counting and analysis. AF received the funds and coordinated the results as well as final proof reading the manuscript with EOD, GE‑R, DOK and ARA. All authors read and approved the final manuscript.

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Competing interests
The authors declare that they have no competing interests.

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