



Assessment of Radionuclide Concentration and Exhalation Rates in some NORMs and TENORMs of Shivalik Region

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²²⁶Ra, ²³²Th, their decay products ²²²Rn, ²²⁰Rn and ⁴⁰K significantly contribute to the mean dose from natural background radiation. This study reports the concentration of radionuclides in Naturally Occurring Radioactive Materials (NORMS) and Technologically Enhanced Naturally Occurring Radioactive Materials (TENORMS) of the Shivalik region of Punjab and Himachal Pradesh. The activity of radionuclides ²²⁶Ra, ²³²Th and ⁴⁰K in different NORMS (soil, sand, rocks) and TENORMS (tiles, marble, cement) were calculated by Gamma spectrometry method using an HPGe detector. Radon mass exhalation and thoron surface exhalation rates were also determined using a Smart RnDuo monitor for the above samples. From the results, it could be concluded that the radon mass exhalation rate and radium activity were obtained to be maximum for tile no. 2 and minimum for white marble samples. Similarly, the thoron surface exhalation rate and thorium activity occurred maximum in tile no. 2 and minimum for safedi. The radium equivalent activity (Ra_{eq}), gamma activity concentration index (I), internal hazard index (H_{int}) and external hazard index (H_{ext}) were computed using appropriate relations. The values range between 5.858 to 188.944, 0.0403 to 1.377, 0.028 to 0.615 and 0.015 to 0.510 having mean values of 107.256, 0.685, 0.325, and 0.256, respectively.

Keywords: Dose, Gamma spectrometry, Exhalation, Smart RnDuo, hazard index

Introduction

NORMS are present all over the earth's map. This term is used to characterize minerals and other materials, including radionuclides from natural roots, which become a reason for remarkable growth in exposure to employees or people and cannot be ignored from the radiation safety point of view. ²³⁸U, ²³²Th and ⁴⁰K are the three main long-lived radioactive materials in NORM present in the earth's crust. Humans are continuously exposed to this natural radiation every day. Artificial sources ¹³⁷Cs also cause radiation exposure. Due to human actions like energy-yielding, mining, industrial waste, and military functioning like nuclear accidents or weapon examination, these radionuclides may be released into the environment¹. These radionuclides pose a potential hazard to human wellness due to the ejection of ionizing radiation². Another activity, like flying at high altitudes, also increases natural exposure to the ahead limit. Controlling people's exposure to background radiation necessitates learning more about

the classification, characteristics and levels of radionuclides found in rocks and soils. TENORMS include wastes with a small specific activity liberated from the various manufacturing projects, including uranium mining, production of phosphate fertilizers, chemical processing of ores, removal, and cleansing of trace elements, *etc.* In TENORMS, Radionuclides ²³⁸U, ²³²Th and their respective disintegrate products are present in the majority. Other radionuclides ⁴⁰K, ⁸⁷Rb and ¹³⁷Cs are present in levels of background doses.

It is well-known that the distribution of radiation over the globe is not uniform but varies according to factors like the abundance of radioactive elements in the planet's crust³. The level of radionuclides in plants is linearly linked to radionuclide concentration in soil⁴, and it can be further increased by applying phosphate fertilizers which contains radioactivity almost ten times higher than the mean concentration of soil. The manufacturing, shipping, storing and usage of these fertilizers become the causes of increased doses of exposure to humans⁵. The samples of TENORMS are building materials like gypsum,

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granite, marble, cement, different types of wall colors, white cement, etc. These construction materials may also cause increase radioactivity significantly in radiation doses.

Radon and thoron are substantial donors to radiations' mean natural background dose. They correspond to around half of the total dose received from entirely natural sources of ionizing radiation⁶. These radioisotopes enter into the environment through two processes known as emanation and exhalation. Firstly, escape from the grains forming the medium, a process named emanation, then from the medium's top is called exhalation⁷. The entry of ²²²Rn into the indoor environment happens predominantly due to the exhalation of ²²²Rn from the soil and construction materials. These factors depend upon other parameters like half-life, aggregation rate of these gases in the interior domain, microstructure of materials, diffusion, advection, absorption etc. The time at which half of the ²²²Rn atoms decay is 3.8 days while for ²²⁰Rn is 55.6 sec. Because of this half-life inequality, the dose from thoron and its daughter products is much lesser than radon and its progeny, which is around 10% of the former⁸. Generally, indoor radon concentration is higher than the thoron indoor concentration. However, the latter becomes vital because of the significant accumulation rate of its progeny (²¹²Pb having a half-life of 10.6 h) in breathable air⁹. Some studies^{10, 11} also show that ²²⁰Rn concentrations are justifiably equivalent to ²²²Rn and its progeny in a few areas of lifted radiation's threats.

Kaur¹² reviewed the activity concentration in the states of North India: Uttar Pradesh, Punjab, Rajasthan and Haryana. Out of which, a higher concentration of ²³²Th was observed in some districts of Punjab. In contrast, a higher content of ²²⁶Ra and ⁴⁰K was noticed in some districts of Haryana, which may be due to the excessive adoption of abundant potassium fertilizers for agricultural growth. Mehta¹³ reports the radon mass exhalation rate of Mohali, Punjab which is found to be lower than the mean of the worldwide value 57 mBqKg⁻¹h⁻¹. Semwal¹⁴ publish the exhalation report from Almora, Uttarakhand, India of soil and building materials. Variation of mass exhalation rate observed from 16 to 54 mBq.kg⁻¹h⁻¹, while the area exhalation rate lies between 0.65 and 6.43 Bq.m⁻².s⁻¹. Rani¹⁵ reported the exhalation rate in soil samples of Barnala and Moga districts. The radon mass exhalation rates have been found to vary from 4.56 ± 0.53 to 41.13 ± 1.21 mBqkg⁻¹h⁻¹ with an average value of 24.8 ± 1 mBqkg⁻¹h⁻¹ while thoron

surface exhalation rates lie in the range from 1.50 ± 0.63 to 23.77 ± 1.52 kBqm⁻²h⁻¹ with mean value of 15.62 ± 1.24 kBqm⁻²h⁻¹. Kansal¹⁶ reports the measurements of ²²⁶Ra, ²³²Th and ⁴⁰K and exhalation rates in soil samples collected from some parts of western districts of Haryana, India. The activity concentrations of ²²⁶Ra, ²³²Th and ⁴⁰K range from 13.93 to 142 Bq kg⁻¹, 35.95 to 91.78 Bq kg⁻¹ and 299.57 to 1056.77 Bq kg⁻¹, respectively. The radium equivalent activity (Ra_{eq}) in all the soil samples varies from 92.51 to 287.08 Bq kg⁻¹ with an average of 184.15 Bq kg⁻¹.

Intake of these radionuclides gases and their progenies causes several health hazards. The inadequately ventilated areas result in high population exposure to these gases compared to other areas. Long-term exposure to these gases in houses, buildings and workplaces amplifies the hazard of Lung Cancer¹⁷.

The present study measures the exhalation rate of ²²²Rn and ²²⁰Rn in NORMs (soil, sand, rocks etc.) and TENORMs (common building materials).

1 Study Area

1.1 Punjab

The Indo-Gangetic basin. The region's dominant economic feature is agriculture, which has consequently shaped the foundation of Punjabi culture, where an individual's social standing is often contingent on land ownership. The physiographic setting of the state is characterized by two primary geomorphic entities - the Siwalik foothills in the northeast and the alluvial fill of the Indus drainage basin. The geological units of the Punjab state consist of the Siwalik Supergroup and Quaternary alluvium. The latter includes older and newer alluvium, with a maximum sediment depth of 4500m. The Siwalik Supergroup is further classified into three groups - Lower, Middle, and Upper Siwalik groups. The fertile plains of Punjab are renowned for their high agricultural productivity, accounting for roughly two-thirds of India's annual food grain production. The region predominantly features calcareous soil, including soil and sierozem soil, with alluvial soil generally described as arid and brown. In eastern Punjab, the soil type ranges from loamy to clayey¹⁸.

1.2 Himachal Pradesh

Himachal Pradesh is a state located in the northern part of India, positioned within the Western Himalayas. Being one of the thirteen mountain states,

it is renowned for its challenging terrain, featuring numerous peaks and vast river systems. The geology of Himachal Pradesh is primarily characterized by Precambrian rocks that were assembled and deformed during the collision between India and Asia, followed by the Himalayan orogeny. The state's elevation ranges from 320m to 6975m, with rock materials largely sourced from the Indian craton, and their ages ranging from the Paleoproterozoic era to the present day. It is generally agreed that the Indian craton collided with Asia around 50-60 million years ago, causing extensive thrusting and folding of rock sequences during the collision. The area has also undergone substantial orographic precipitation, glaciation, and rapid erosion, shaping its overall landscape. Himachal Pradesh's exposed geological formations range from the Proterozoic to the Quaternary period, representing a classic geological sequence. The undifferentiated Proterozoic rocks are predominantly confined to the Lesser Himalaya and are represented by the Jutogh and Vaikrita groups. The Jutogh Group comprises formations such as Panjerli, Manal, Bhotli, Khirki, Taradevi, Kanda, Naura, Badrol, Rohru, Chirgaon, and Jaknoti, consisting of a thick sequence of carbonaceous phyllite, quartzite, and slate¹⁸.

2 Materials and Methodology

2.1 Sampling Procedure

The materials were categorized into two classes—NORMS and TENORMS. In the category of TENORMS, 15 samples of building materials such as gypsum, granite, limestone, bricks, marble, white cement, interlock tiles, cement and tiles were aggregated from various vendors. For the collection of NORMS, different types of soils, sand, and rocks of various sizes were assembled from the river named 'Sutluj' collected. The samples must be free from other impurities. The materials, like bricks, tiles, marbles etc. were in their original size. Firstly, they were broken using a heavy pestle into coarse parts. Afterwards, all the samples were grinded using a grinder and sieved through 100-micron mesh to keep the uniformity of the grains. The mass of each sample varied from 1.5 kg to 2 kg. The collected soil samples were appropriately labelled. The samples were kept in an oven to dry at 105°C for 48 hours and sieved. Before the radiometric analysis, each sample was sealed in an air-tight Marinelli beaker to prohibit liberation of ²²²Rn and ²²⁰Rn from the samples and stored for four weeks to obtain equilibrium between

²²⁶Ra, ²³²Th and their decay products. To determine the rate of exhalation from each location, dried samples of mass approximately 1 kg were taken in a polyethylene bag.

2.2 Smart RnDuo Monitor

SMART RnDuo Monitor (Alpha-Scintillometer GBH2002 with Lucas cell), designed and calibrated by BARC (Bhabha Atomic Research Centre), Mumbai, India¹⁹, is adopted for the quantification of ²²²Rn mass exhalation (J_{rm}) and ²²²Rn areal exhalation (J_{ts}) rate in compiled samples. The ZnS:Ag screen works like a scintillation material having a volume of 150 cc. The principle is defined as the spotting of alpha radiations produced from radon decay and their progenies within the walls of detector volume detected by a scintillation screen coated alongside ZnS:Ag. The sensitivity factor of the monitor is 1.2 counts per hour ($Bq \cdot m^{-3}$), and the detection limit lies between 8 $Bq \cdot m^{-3}$ and 10 $MBq \cdot m^{-3}$.

A sample of known weight was placed into an exhalation chamber to measure J_{rm} . The experimental setup shown in Fig. 1 shows that measurement was done in diffusion mode and detector probe should be coupled with an exhalation chamber. The exhalation chamber consists of a cylindrical accumulator of stainless steel with an inner diameter of 30 cm and a height of 5 cm. The time cycle for measurement was 60 minutes, and a minimum of 18 cycles was required for measurement. The formula used for radon concentration $C(t)$ at time T is²¹

$$C(t) = \frac{J_{rm}M}{V\lambda_r} (1 - e^{-\lambda_r T}) + C_0 e^{-\lambda_r T} \quad \dots(1)$$

Where, C_0 is the radon concentration ($Bq \cdot m^{-3}$), M is the weight of the prepared sample (kg) placed in the chamber, λ_r is the effective radon decay constant, and T is the time (h) of measurement.

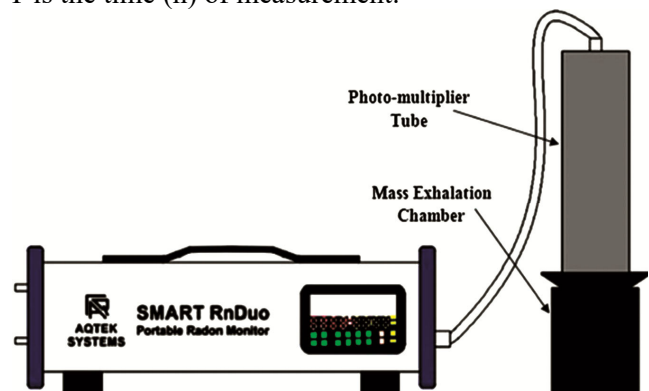


Fig. 1 — Experimental setup of Smart RnDuo for measurement of ²²²Rn exhalation²⁰.

The sampling was done in flow mode for the measurement of J_{ts} , where two cylindrical pipes of diameter 0.5 cm were used through the chamber's top to function as the inlet and outlet of airflow for the determination of thoron concentration. The exhalation chamber is linked to the pump inlet of the monitor. A closed-circuit loop formed as the detection box was linked to the Smart RnDuo monitor via the airflow pump. Four cycles of time 15 min were used for analysis. Pump should be kept on for an initial 5 min to determine the background of the thoron. The J_{ts} ($\text{Bqm}^{-2}\text{h}^{-1}$) of thoron in closed loop²² is given by Eq. (2):

$$J_{ts} = \frac{C_{eq} \times V \times \lambda}{A} \quad \dots(2)$$

Where, C_{eq} (Bq.m^{-3}) accounts for the mean of observations of thoron gas concentration, V (m^3), is

Effective volume, λ is the decay constant (0.012464 s^{-1} for ^{222}Rn), and A is the sample's surface area (m^2). The measurement of every sample was done 2-3 times to increase the precision and accuracy found by the inventor of the instrument.

2.3 Gamma Spectrometric analysis

A 42% relative efficiency n-type low background HPGe coaxial cylindrical gamma spectrometer with a detector made by CANBERRA, USA, was used for radioactivity measurements. Lead was used as a shielding material with fixed bottom and a purity of

99.9%. The movable cover was used to reduce background radiation in the spectrum. The detector was linked with 16K MCA and ^{60}Co emission for data acquirement. The period of 50,000 sec was enough for data collection to yield better statistics. The net sample count rate was derived after subtracting the corresponding background rate at each energy peak. In the same way, an empty Marinelli beaker was also computed for the determination of background radiation. The calibration of energy and efficiency of the spectrometer was calculated by adopting Marinelli calibration sources, including ^{210}Pb , ^{241}Am , ^{109}Cd , ^{57}Co , ^{113}Sn , ^{85}Sr , ^{137}Cs

3 Results and Discussion

Table 1 shows that the activity concentrations of ^{226}Ra , ^{232}Th , and ^{40}K obtained from samples vary from $4.5 \pm 0.4 \text{ Bq.kg}^{-1}$ to $41.7 \pm 1.1 \text{ Bq.kg}^{-1}$, 0.1 Bq.kg^{-1} to 89.75 Bq.kg^{-1} , and 1.4 Bq.kg^{-1} to $508 \pm 11 \text{ Bq.kg}^{-1}$ respectively. These values lie within the range of radionuclides given by UNSCEAR (United Nations Scientific Committee on the Effects of Atomic Radiation)²². The values of the concentrations of radionuclides ^{226}Ra , ^{232}Th and ^{40}K and Radium equivalent activity (R_{ea}), Gamma activity concentration index (I), internal hazard index (H_{int}) and external hazard index (H_{ext}) are given in Table 1.

Table 1 — Activity of ^{226}Ra , ^{232}Th and ^{40}K in Bq.kg^{-1} , R_{ea} , I, H_{int} and H_{ext} for different samples

Sr. No.	Sample	^{226}Ra (Bq.kg^{-1})	^{232}Th (Bq.kg^{-1})	^{40}K (Bq.kg^{-1})	R_{ea}	I	H_{int}	H_{ext}
1	Tile 2	41.7 ± 1.1	89.7 ± 2.1	357.51 ± 91	177.19	1.2713	0.591	0.478
2	Tile 3	15.4 ± 0.8	68.2 ± 2.7	319.74 ± 19	142.546	1.031	0.441	0.384
3	Rocks	18.1 ± 1.2	27.7 ± 1.6	142.11 ± 4.6	200.903	1.447	0.648	0.543
4	Putty	15.4 ± 0.8	3.8 ± 0.6	35.3 ± 4.1	23.56	0.164	0.105	0.636
5	Tile 1	31.6 ± 1.1	78.8 ± 2.6	274 ± 9	187.882	1.336	0.614	0.507
6	Cement 2	39.3 ± 1.2	66.9 ± 2.4	356.9 ± 10.8	162.448	1.168	0.544	0.438
7	Sand	31.6 ± 0.9	58.4 ± 1.8	508 ± 11	154.228	1.133	0.502	0.416
8	Plaster of Paris	6.5 ± 0.6	1.8 ± 0.5	52.8 ± 4.6	13.139	0.096	0.0531	0.035
9	White marble	4.5 ± 0.4	0.6 ± 0.3	6.5 ± 1.5	5.858	0.0403	0.0279	0.016
10	Safedi	15.9 ± 1.1	<0.1	<1.4	16.151	0.108	0.086	0.043
11	Samosam	26.8 ± 1.0	<0.1	9.4 ± 3.3	<25.167	0.169	0.134	0.068
12	Cement 1	41.6 ± 1.2	60.6 ± 2.2	317 ± 10	152.667	1.094	0.525	0.412
13	Limestone	9.2 ± 0.7	6.9 ± 0.9	<1.4	<19.174	0.131	0.0766	0.051
14	White cement	26.8 ± 0.9	16.4 ± 1.1	74.8 ± 5.2	56.012	0.392	0.223	0.152
15	Gypsum	20.8 ± 0.8	31.8 ± 1.5	394 ± 11	96.612	0.719	0.317	0.261
12	Cement 1	41.6 ± 1.2	60.6 ± 2.2	317 ± 10	152.667	1.094	0.525	0.412
13	Limestone	9.2 ± 0.7	6.9 ± 0.9	<1.4	<19.174	0.131	0.0766	0.051
14	White cement	26.8 ± 0.9	16.4 ± 1.1	74.8 ± 5.2	56.012	0.392	0.223	0.152
15	Gypsum	20.8 ± 0.8	31.8 ± 1.5	394 ± 11	96.612	0.719	0.317	0.261

3.1 Radium Equivalent Activity (Ra_{ea})

Ra_{ea} may be computed from the relation given below²³:

$$Ra_{ea} = 1.43A_c(Th) + A_c(Ra) + 0.077A_c(K)$$

where A_c(Th) is the activity concentration of ²³²Th in Bq.kg⁻¹, A_c(Ra) is the activity concentration of ²²⁶Ra in Bq.kg⁻¹, A_c(K) is the activity concentration of ⁴⁰K in Bq.kg⁻¹.

3.2 Internal Hazard Index (H_{int})

The internal exposure to radon and its progeny is assessed by H_{int}, which is defined by the following relation:

$$H_{int} = (A_c(Ra)/185) + (A_c(Th)/259) + (A_c(K)/4810)$$

3.3 External Hazard Index (H_{ext})

The H_{ext} is interpreted as follows

$$H_{ext} = (A_c(Ra)/370) + (A_c(Th)/259) + (A_c(K)/4810)$$

3.4 Exhalation Rate

Table 2 lists the J_{rm} in mBq.kg⁻¹.h⁻¹ and J_{ts} in Bq.m⁻².h⁻¹ for different samples. The J_{rm} varied from 4.5 to 21.55 mBq.kg⁻¹.h⁻¹ while J_{ts} extends from 42.44 to 2376.91 Bq.m⁻².h⁻¹. The average values were 11.287 mBq.kg⁻¹.h⁻¹ and 1128.246 Bq.m⁻².h⁻¹ for mass exhalation and surface exhalation rates, respectively. The distribution of radon and thoron exhalation rates in shown by graph given in Fig. 2 & 3. The highest value of J_{rm} and J_{ts} was observed for tile 2. The minimum value of J_{rm} for the white marble sample, while the lowest area exhalation rate was found for safedi. Fig. 4 shows the correlation (R²=0.85) between ²³²Th and ²²⁰Rn surface exhalation rate, and Fig. 5 shows the correlation (R²=0.81) plot between

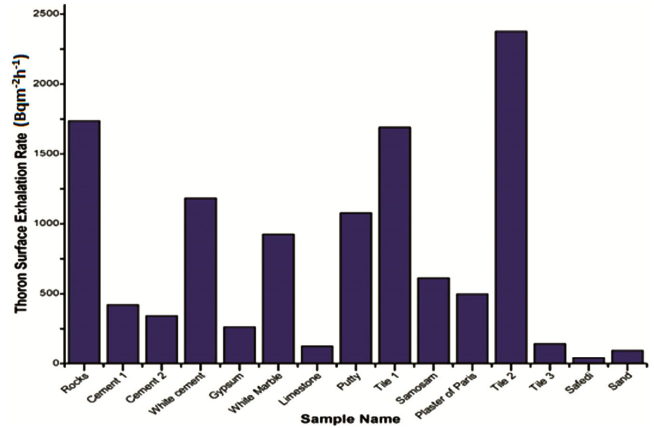


Fig. 2 — Bar distribution of thoron surface exhalation rate (J_{ts}) with samples.

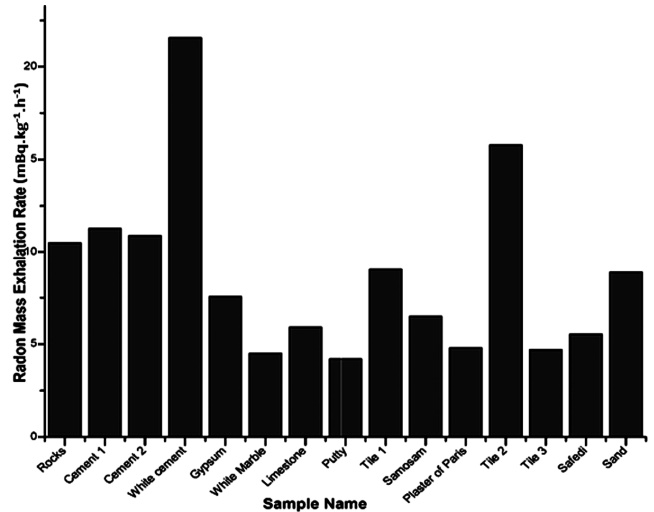


Fig. 3 — Bar distribution of radon mass exhalation rate (J_{rm}) with samples

Table 2 — J_{rm} and J_{ts} for the samples

Sr. No.	Sample Name	J _{rm} (mBqkg ⁻¹ .h ⁻¹)	J _{ts} (Bqm ⁻² .h ⁻¹)
1.	Rocks	10.5	1736.91
2.	Cement 1	11.27	421.12
3.	Cement 2	10.89	341.66
4.	White cement	21.55	1183.35
5.	Gypsum	7.55	262.55
6.	White Marble	4.5	925.71
7.	Limestone	5.91	125.12
8.	Putty	4.2	1078.12
9.	Tile 1	9.03	1691.16
10.	Samosam	6.5	612.3
11.	Plaster of Paris	4.8	498.13
12.	Tile 2	15.77	2376.91
13.	Tile 3	4.71	142.55
14.	Safedi	5.53	42.44
15.	Sand	8.87	93.66

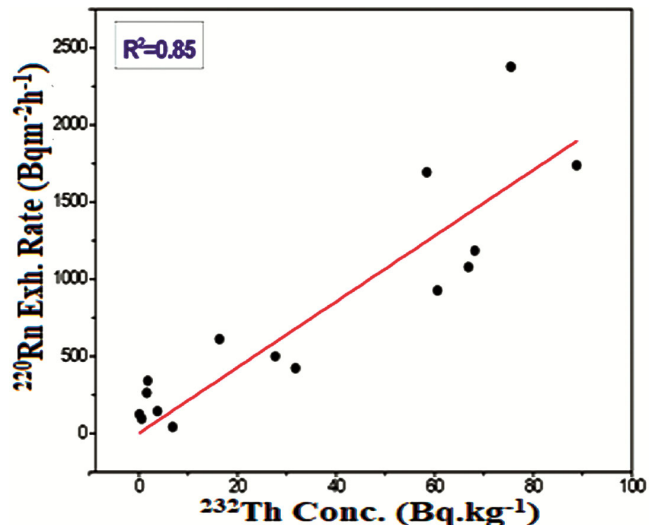


Fig. 4 — Correlation plot between the activity of ²³²Th and ²²⁰Rn exhalation

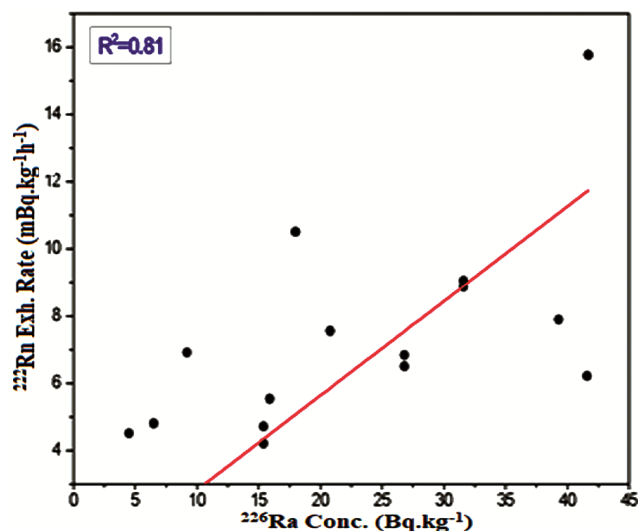


Fig. 5 — Correlation plot between the activity of ^{226}Ra and ^{222}Rn exhalation Rate

the activity of ^{226}Ra and ^{222}Rn mass exhalation rate of the samples. J_{m} in the present study is below the worldwide average ($57.6 \text{ Bqm}^{-2}\text{h}^{-1}$). Hence residents do not possess any health hazards.

4 Conclusions

In this study, the activity of ^{226}Ra , ^{232}Th and K^{40} gamma spectrometry techniques using high resolution in some NORMs and TENORMs were measured by HPGe gamma spectrometer. Smart RnDuo monitor was used to measure the J_{m} and J_{is} . The correlations between the exhalation rates of these gases and their parent nuclide concentrations (radium/thorium) were examined. Finally, Ra_{ea} , I , H_{int} and H_{ext} were calculated. For ^{220}Rn , the J_{is} ranged from 42.44 to 2376.91 $\text{Bqm}^{-2}\text{h}^{-1}$, as shown in Fig. 2. In the case of ^{222}Rn , the J_{m} varied from 4.5 to 21.55 $\text{mBq kg}^{-1}\text{h}^{-1}$ shown in Fig. 3. The activity of ^{226}Ra was found to vary between $4.5 \pm 0.4 \text{ Bq.kg}^{-1}$ and $41.7 \pm 1.1 \text{ Bq.kg}^{-1}$. The activity of ^{232}Th was found to lie between 89.75 Bq.kg^{-1} and 0.1 Bq.kg^{-1} . The activity of ^{40}K was higher in the sand ($508 \pm 11 \text{ Bq.kg}^{-1}$) and minimum in limestone ($< 1.4 \text{ Bq.kg}^{-1}$). The radiological parameters Ra_{ea} and H_{ext} were also within the recommended limits. The activity values attained from the current study show that the natural background radiations are in resemblance with other studies.

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