Measuring γ -ray dose of terrestrial samples using β - γ Spectrometry

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Background: The existence radionuclides in soil and some building materials produce a β-y radiation field, which in some regions the exposure of these radionuclides to human is high. Materials and Methods: The air-absorbed dose, indoor and outdoor annual effective dose of soil and some building material samples (ceramic, granite, gypsum, etc.). The samples were collected from 35 different regions in the northwest of Iran were calculated. Specific activity of natural radionuclides 226 Ra, 232 Th and 40 K were measured by using a β - γ spectrometer consisted of NaI(TI) and organic scintillators. The analysis of measured spectra was based on maximum likelihood estimation. The calculated data were compared with world's mean values. Results: It is found that the specific activity of 40K ranges from 573.8 to 1392 Bqkg-1, for 232Th ranges from 6 to 54.6 Bqkg-1 and for 226Ra ranges from and 5.1 to 36.2 Bqkg-1. In some regions, the concentrations of natural radionuclides (40K, 232Th, 226Ra) were higher than standard level. Conclusion: Experimental results related to specific activity, indoor and outdoor annual effective doses of ²²⁶Ra. 232Th and 40**K** revealed that radionuclide concentration in soil and some building material samples of some regions of northwest of Iran are of some radiological importance. Iran. J. Radiat. Res., 2011; 8 (4): 237-242

Keywords: Environmental radioactivity, β-γ spectrometry, annual effective dose rate.

INTRODUCTION

Many radionuclides exist naturally in terrestrial soils, rocks and in building materials developed from them. In term of dose, the most important primordial (half-life comparable to the age of the earth) radionuclides are ⁴⁰K, ²³²Th and ²³⁸U. Their absolute and relative concentrations in soil and construction materials can vary dramatically depending on their source ⁽¹⁾. The decay of these radionuclides in soil produces a β-γ radiation field that crosses the soil-air interfaces and produces exposure to humans. Many surveys in order to

estimate their concentration and assessment of the associated dose rate in air have been made in most inhabited regions of the world (2-11) using gamma spectrometry.

The products of uranium and thorium chains are always present in soil and rock. Their gamma radiation causes external exposure with the consequent absorbed doses. The avialable geological data suggests that soils and rocks from the zone of Zonoz County, Caulan mines and Mishu Mountain in the northwest of Iran may have high level naturally occuring radioactivity. The purpose of our research has been to clarify the amount of natural radiation exposure of these regions' population. Gamma spectrometry combined with beta spectrometry, and analyzing the measured spectra applying an algorithm based on maximum likelihood estimation allowed the identification of the potential source of contamination in order to estimate the possible radiological hazards to human health. Considerable attention should be paid to decrease exposure level arising from members of uranium and thorium decay chains and by potassium-40 soils.

MATERIALS AND METHODS

Sample collection and preparation

Samples of soils, rocks and building materials were collected from 35 different regions of Eastern Azerbaijan, Iran. The studied areas, which lie at 38°30′N latitude and 45°45′E longitudes, in figure 1 are shown. Surface soil samples (0–5 cm deep)

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were carefully collected from areas, which were not covered by any type of vegetation and thought to be undisturbed. grounded thoroughly homogenized changed the rock samples into the powder form before the measurements. All soil samples and building materials converted into very small particles and screened with a sieve of about 1mm mesh. The prepared and homogenized samples were weighted and put into Marinelli beakers with the volume of 1 liter and flat vessels 0.1, 0.2 and 0.5 liter volume. Then they were packed into radon-impermeable plastic containers. Each beaker was kept for 4 weeks (>seven halflives of ²²²Rn and ²²⁴Ra) prior to counting in order to ensure that the daughter products of ²²⁶Ra up to ²¹⁰Pb and of ²²⁸Th up to ²⁰⁸Pb achieve equilibrium with their respective parent radionuclides (12).

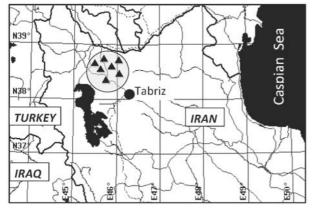


Figure 1. Simplified geology of Northwestern Iran. Locations of soil sampling is shown by solid triangles.

Experimental setup

The $6\text{-}\gamma$ spectrometer which was used $^{(13)}$ consisted of two cylindrical scintillators: a $\Phi128\times8$ mm paraterphenyl-activated polystyrene plastic for beta detection and a $\Phi63\times63$ mm NaI(Tl) for gamma detection. The scintillators were made by ATOMTEX, Belarus. The NaI(Tl) detector had relative efficiencies of < 8.5% for the 662 keV γ -ray line of 137 Cs. To derive maximum detection efficiency, the beta and gamma detectors were both surrounded by 5 cm thick low-activity lead shield. The source-detector geometry has been shown in figure 2. In case of voluminous environmental samples,

the best limit of detection per mass or volume unit was achieved when the sample were filled in Marinelli beaker. The minimum detectable activity (MDA) for the beta-gamma spectrometer system used in the present study was ~2 Bq/kg for ¹³⁷Cs for a counting time of 3 hours and for one liter Marinelli beaker sample ⁽¹⁴⁾.

detectors The were connected to personal computer (which has built in detector HV bias supply, ADC and MCA) for simultaneously beta and gamma ray energy spectrum acquisition. The measured spectra were analyzed by SPTR-ATM program based on maximum likelihood algorithm for activity estimation. One of the measured energy spectrum and assigned gamma line is shown in figure 3. The detector efficiency performed calibration was bv standard point and volume sources.

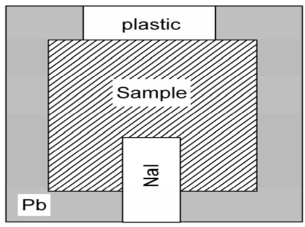


Figure 2. Source-detector geometry. β and γ radiations are emitted from voluminous sample and are detected by plastic and Nal scintillators, respectively.

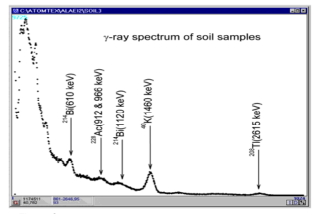


Figure 3. A γ-ray spectrum of a soil sample measured by using Nal(TI) scintillator.

Maximum likelihood activity estimation

For those radionuclides with half life longer than the time period of measurement β-particles or y-rays came from a very large number of independent nuclei within the sample, Poisson statistical model appropriate. For this reason, in the gamma and beta spectrometry of the environmental samples, the Poisson distribution is usually used to analyze the measured spectra. The likelihood maximum (ML)method activity estimation was derived from the condition of the maximum of the logarithm of likelihood function (15). The logarithm of likelihood function for the distribution of counts in the beta or gamma spectrum channels is given as:

$$L^* = \ln L = \sum_{i=1}^{N} \left[-t \sum_{k=1}^{M} A_k f_k(i) + S(i) \ln \left(t \sum_{k=1}^{M} A_k f_k(i) \right) - \ln(S(i)!) \right]$$
 (1)

Where S(i) indicates the beta or gamma experimental spectrum of a mixture of radioactive nuclides, I, is the channel number; N, is the number of channels (=1024); t, is the spectrum accumulation time, M is the number of spectral components; f_k (i) for k = 1, 2, ..., M-1, is the spectrum of the kth nuclide normalized to unit activity and unit time; f_M (i), is the spectrum of a background normalized to unit time; A_k is the unknown activity of the kth nuclide.

The estimates of A_k are determined by solving the following system of the M equations deriving from the condition of the maximum of equation 1:

$$\frac{\partial L *}{\partial A_k} = 0$$

$$\sum_{i=1}^{N} \left[\frac{s(i) f_j(i)}{\sum_{k=1}^{M} A_k f_k(i)} - t f_j(i) \right] = 0, \quad j = 1, 2, ..., M$$
(2)

The system of equation 2 is solved by iteration method. In every qth iteration (q=1, 2, ...) one solves the set of the linear equations:

$$\sum_{k=1}^{M} t A_k \sum_{i=1}^{N} \frac{f_k(i) f_j(i)}{D_q(i)} = \sum_{i=1}^{N} \frac{S(i) f_j(i)}{D_q(i)}$$
(3)

$$D_{q}(i) = \begin{cases} \sum_{k=1}^{M} A_{k}^{(q-1)} f_{k}(i) & \text{if } q > 0 \\ S(i) + 1 & \text{if } q = 0 \end{cases}$$
 (4)

Where:

 $A_k^{(q-1)}$ are the estimates of the activities obtained in (q-1) iteration, $D_q(i)$ are the variances and $1/D_q(i)$ are statistical weights of the number of counts in each channel. Such proposed iteration scheme for calculation of ML estimates seems to be very effective, as a rule, 2-3 iterations provide the error of approximation of <1%.

Absorbed dose rates

The gamma dose rates in air were calculated using the results of 40 K, 232 Th and 226 Ra activities and dose coefficients (nGy h 1 per Bq kg ${}^{-1}$) 0.462, 0.604 and 0.0417 given in UNSCEAR (2000) for 226 Ra sub-series, 232 Th series and 40 K respectively and by assuming secular equilibrium between 238 U and 226 Ra ${}^{(16)}$. The absorbed dose is given by:

$$D(nGyh^{-1}) = [(0.462 \times A_{Ra}) + (0.604 \times A_{Th}) + (0.0417 \times A_{k})]$$
 (5)

Where A_{Ra} , A_{Th} and A_{K} are the specific activities of ^{226}Ra , ^{232}Th and ^{40}K , respectively, in Bqkg-1 in the samples under study. We converted the absorbed dose in air into annual effective indoor and outdoor doses received by individuals. Annual estimated average effective dose equivalent received by a member is calculating using a conversion factors of 0.7, 0.8 and 0.9 SvGy-1 for adults, children and infants, respectively (16) which is used to convert the absorbed rate to annual effective dose with an outdoor occupancy of %20 and %80 for indoors (17). The annual effective doses are determined as follow:

eff .ann .Dose =
$$\begin{cases} D (nGyh^{-1}) \times 8760 (h) \times 0.7 (SvGy^{-1}), & Indor \\ D (nGyh^{-1}) \times 8760 (h) \times 0.2 \times 0.7 (SvGy^{-1}), & Outdoor \end{cases}$$

where 0.2 and 8760 h are outdoor occupancy and time factors, respectively as recommended by ⁽¹⁸⁾.

RESULTS AND DISCUSSION

A result summary of natural radionuclide activity concentrations in samples taken from 35 locations is shown in table 1. The specific activities for ²²⁶Ra, ²³²Th and ⁴⁰K are reported in Bqkg⁻¹. The ± values

shown are because of the *s* variation due to counting errors. The specific activity of ²²⁶Ra, ²³²Th and ⁴⁰K in various materials which were studied in the present work, vary from 5 to 36.2, from 6 to 54.6 and from 573.8 to 1392 Bqkg⁻¹, respectively. The world's mean values of ²²⁶Ra, ²³²Th and ⁴⁰K activity concentrations are 32, 45 and 420

Bqkg⁻¹ (13) . A strong positive correlation of r =0.705 was observed between specific activity of ²²⁶Ra and ²³²Th in the collected soil samples. This clearly indicated the presence of significant amounts of monazite and zircon sands in the samples. A weak correlation of r =0.44 was observed between specific activity of ²²⁶Ra and ⁴⁰K in the

Table 1. Specific activities of ²²⁶Ra, ²³²Th, ⁴⁰K of all samples.S2, S3 and S4 are second, third and forth samples of a given site.

Sampling sites (kind of samples)	Specific activity (Bqkg ⁻¹)		
	²²⁶ Ra	²³² Th	40 K
Zonoz (soil)	17.7±0.1	37.2±1.1	784.4±7.4
S2	18.8±0.5	39.1±1.7	778.0±5.1
S3	18.5±0.4	38.0±0.9	772.0±6.0
S4	19.0±0.7	37.0±0.8	780.0±6.1
Zonoz (cement)	11.1±0.1	32.9±0.1	610.0±1.8
S2	12.0±0.2	32.0±0.1	614.0±1.9
S3	12.5±0.2	34.0±0.2	600.0±0.4
Zonoz (granite)	13.0±0.3	30.0±0.1	605.0±0.5
Zonoz (brick)	14.5±0.3	35.0±0.3	620.0±1.9
S2	13.8±0.2	36.0±0.2	622.0±2.0
Caulan mine1(soil)	29.8±0.3	49.5±1.4	1206.2±11.1
S2	32.1±0.9	52.8±1.8	1220.0±15.2
Caulan mine1 (gypsum)	34.6±1.0	50.1±0.9	1199.2±10.3
S2	35.5±1.1	54.6±0.9	1200.0±9.8
S3	36.2±1.8	53.8±0.8	1209.1±11.3
Caulan mine2 (soil)	10.7±0.5	11.4±0.1	1317.2±15.5
S2	10.3±0.4	13.5±0.3	1298.0±10.1
S3	10.1±0.5	12.8±0.2	1320.0±14.3
Caulan mine2 (gypsum)	14.2±0.3	18.0±0.5	1392.0±11.6
S2	14.1±0.2	16.5±0.4	1371.0±18.2
S3	16.3±0.4	20.0±0.8	1345.0±13.1
Caulan mine3 (soil)	5.1±0.1	8.2±0.1	573.8±13.6
S2	6.2±0.1	7.2±0.1	592.1±14.8
S3	7.0±0.1	8.0±0.1	602.0±18.1
Caulan mine3 (gypsum)	6.0±0.1	7.5±0.1	608.0±19.8
S2	5.5±0.1	6.0±0.1	599.0±15.0
Mahbobabad village (cement)	9.6±0.5	25.9±0.9	910.2±7.4
S2	8.1±0.3	27.1±1.1	911.1±7.6
Mahbobabad village (brick)	8.7±0.1	30.2±1.3	921.5±8.1
S2	9.9±0.3	33.5±1.8	919.0±1.8
Eyshabad village (cement)	9.0±0.4	41.8±2.1	98.0±6.5
S2	8.4±0.3	44.6±2.5	915.0±8.0
Eyshabad village (brick)	9.8±0.5	40.5±1.9	935.1±8.8
Eyshabad village (granite)	10.0±0.6	39.0±1.8	928.7±6.1
Mishu Mountain (soil)	33.6±2.1	35.1±2.0	960.0±15.0

collected soil samples which indicates that ⁴⁰K concentrations may not be related to the presence of ²²⁶Ra-bearing mineral sands.

The absorbed and annual effective dose rates for different samples were calculated using equations 5 and 6 and are listed in table 2. It was assumed that ¹³⁷Cs, ⁹⁰Sr and

the ²³⁵U decay series can be neglected as they contribute very little to the total dose from environmental background ⁽¹⁹⁻²¹⁾. The minimum and maximum values of absorbed dose and indoor and outdoor annual effective doses which were found, vaied from 33.1 to 99.6 nGyh⁻¹, from 0.191 to 0.611 mSv

Table 2. Radiation absorbed dose and dose equivalents for all samples.

	Annual effective dose (mSv)		
Sampling sites (kind of samples)	Absorbed dose (nGyh ⁻¹)	Indoor	Outdoor
Zonoz (soil)	63.4±1.0	0.388±0.006	0.077±0.001
S2	64.7±1.8	0.397±0.011	0.079±0.002
S3	63.7±0.9	0.391±0.005	0.078±0.001
S4	63.7±1.1	0.391±0.006	0.078±0.001
Zonoz (cement)	50.4±0.2	0.309±0.001	0.062±0.0002
S2	50.5±0.2	0.310±0.001	0.062±0.0002
S3	51.3±0.2	0.315±0.001	0.063±0.0002
Zonoz (granite)	49.4±0.2	0.303±0.001	0.061±0.0002
Zonoz (brick)	53.7±0.4	0.329±0.002	0.065±0.0004
S2	54.1±0.3	0.332±0.002	0.066±0.0003
Caulan mine1(soil)	93.9±1.5	0.576±0.009	0.115±0.001
S2	97.6±2.1	0.598±0.013	0.119±0.002
Caulan mine1 (gypsum)	96.3±1.4	0.591±0.009	0.118±0.001
S2	99.4±1.5	0.609±0.009	0.122±0.001
S3	99.6±1.8	0.611±0.011	0.122±0.002
Caulan mine2 (soil)	66.8±0.9	0.410±0.006	0.082±0.001
S2	67.1±0.8	0.411±0.004	0.082 ± 0.0009
S3	67.4±0.9	0.413±0.006	0.083±0.001
Caulan mine2 (gypsum)	75.5±0.9	0.463±0.006	0.093±0.001
S2	73.6±1.1	0.451±0.006	0.090±0.001
S3	75.7±1.2	0.464±0.007	0.092±0.001
Caulan mine3 (soil)	31.2±0.7	0.191±0.004	0.038 ± 0.0008
S2	31.9±0.7	0.196±0.004	0.039 ± 0.0008
S3	33.2±0.8	0.203±0.005	0.041±0.0009
Caulan mine3 (gypsum)	32.7±0.9	0.201±0.005	0.040 ± 0.001
S2	31.1±0.7	0.191±0.004	0.038 ± 0.0008
Mahbobabad village (cement)	58.0±1.1	0.356±0.006	0.071±0.001
S2	58.1±1.1	0.356±0.006	0.071±0.001
Mahbobabad village (brick)	60.7±1.2	0.372±0.007	0.074±0.001
S2	63.1±1.3	0.386±0.007	0.077±0.002
Eyshabad village (cement)	33.5±1.7	0.205±0.010	0.041±0.002
S2	60.9±2.0	0.373±0.012	0.075±0.002
Eyshabad village (brick)	68.0±1.8	0.416±0.011	0.083 ± 0.002
Eyshabad village (granite)	66.9±1.6	0.410±0.009	0.082 ± 0.002
Mishu Mountain (soil)	76.8±2.8	0.471 ± 0.017	0.094 ± 0.003

and 0.038 to 0.122 mSv, respectively.

It should be noted that the measured specific activity values for 226Ra and 232Th were analogous to the world values, which were determined for soil and building materials. However, specific activity of 40K in all of samples specially Caulan mine2 with 1392 Bqkg-1 was much higher than the world values. Hence, these samples pose some health hazard for the occupants residing in these regions and for those using these building materials. The external world wide average annual effective dose for adults is 0.07 mSv (19). According to these results, it seems that the radionuclides concentrations in the examined samples are some radiological importance. natural radioactivity levels in the building construction materials and soils are to a small degree higher than the acceptable limit. Consequently, permanent settlers of these regions are exposed to an increased level of natural ionizing radiation from these samples.

CONCLUSION

Measurement results presented in this work confirm that radiation exposure and attributed risk could be reduced by careful choice of building materials during construction. The experiments also showed that the spectrometer system had a better discrimination performance for low energy beta and gamma sources.

Since radionuclides are also present in water, milk, fruits, meat and so forth, therefore the tracing of these radionuclides should be carried out for agriculture products from the northwest of Iran.

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