Original Research Article

NATURAL RADIOACTIVITY AND RADIOLOGICAL RISK ESTIMATION OF DRINKING WATER FROM OKPOSI AND UBURU SALT LAKE AREA, EBONYI STATE, NIGERIA.

ABSTRACT

Aim: The objectives of this study was to measure the activity concentration of natural radionuclides in different drinking water sources in order to access the associated radiological health risk due to ingestion of such water. Study design: the design of this study is purely experimental. Place and duration: This study was carried out ondrinkingwatersources around Uburu and Okposi salt lakes areas of Ebony state between April and September, 2016. Methodology: sachet waters, borehole water, stream and river waters were collected and chemically treated by adding few drops of nitric acid to each of the samples and then pre-concentrated and kept in a marinelli container for four weeks. The activity concentration of ²²⁶Ra, ²³²Th and ⁴⁰K in all the water samples was measured using the High- Purity Germanium detector. Results: The specific activity concentration of ²²⁶Ra,²³²Th and 40 K ranged from BDL to 3.66± 0.78 Bql⁻¹,BDL to 7.56 ± 0.0.59 Bql⁻¹ and BDL to 23.31 ± 1.65 Bql⁻¹ respectively in sachet water. The activity concentration of ²²⁶Ra, ²³²Th and ⁴⁰K in borehole water ranges from BDL to 5.65±1.25, 0.45±0.09 to 13.00±0.97 and BDL to 26.45±1.83 Bgl⁻¹ respectively. Furthermore the activity concentration of ²²⁶Ra, ²³²Th and ⁴⁰K in stream water ranges from BDL to 2.88±1.32 Bql⁻¹, BDL to 8.93±0.66 Bql⁻¹and BDL to 33.32±2.18 Bql⁻¹ respectively, while that for river water ranges from 0.03 \pm 0.01 to 4.48 \pm 1.13 Bg¹¹, 0.55 \pm 0.10 to 8.60 \pm 0.65Bg¹¹and BDL to 13.85 \pm 0.98 Bql¹respectively. The mean values of annual effective dose obtained for infants, children and adults are within the ICRP and WHO recommended reference values. The life-long cancer risk and hereditary effects due to ingestion of radionuclides by adults show that 16 out of 100,000 may suffer some form of cancer fatality and 9 out of 100,000 may suffer some hereditary effects. Statistical analysis of the data revealed a positively skewed and platokurtic distribution of radionuclides in all the drinking water sources. Conclusion: all the radiological health risk parameters obtained were within their safe values. Therefore, all the sampled drinking water are radiologically safe for domestic use but infant should not be given any of the studied drinking water since the annual total effective dose for infant exceeded the safe value.

Keywords: Radioactivity, High-purity Germanium, Effective dose, Radiological risk, Uburu and Okposi)

1. INTRODUCTION

Humans have always been exposed to natural radiation arising from the earth as well as from outer space. Terrestrial radiations are given out from natural radioactive elements present in varying amounts in all types of water, soil, rocks, food and other environmental media around us [1]. Radiation exposure through drinking water results from naturally occurring radionuclides in drinking water sources, in particular alpha radiation emitting uranium, radium and their progeny including radon [2]. The occurrence of natural radionuclide in drinking water poses a problem of health hazard, when these radionuclide are taken into the body by ingestion. The radionuclide contributing significantly to the ingestion dose via consumption of water is radium. Radium is a naturally occurring isotope found in the earth's crust, a member of the uranium ²³⁸U decay series. The predominant 25 radium isotopes in ground and surface water are ²²⁶Ra, an alpha emitter with half-life of 1600 years 26 and ²²⁸Ra, a beta emitter with a half-life of 5.8 years [2]. Many salts of radium are soluble in water and 27 therefore surface water may be enriched in radium and its descendant radon. ²²⁶Ra is an earth 28 29 alkaline element sharing the metabolic pathways of calcium in the human body. Due to their

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radiotoxicity especially those of ²²⁶Ra, a contamination hazard for humans exists even at low
 concentration levels [3].

32 Ingested radionuclides are absorbed into the blood (International Commission on Radiological 33 Protection, ICRP, 2007) and accumulates in specific tissues that they may damage. Of absorbed 34 uranium, 66% is rapidly eliminated via urine while the rest is distributed and stored in the kidney (12-35 15%), bone (10-15%) and soft tissues (Wrenn et al., 1985). The internal exposure of humans to 36 ionizing radiation is through inhalation and ingestion. When the radioisotope enters the body, it 37 accumulates in the tissue of body organ. The rate of clearance of such radionuclide from the tissue or 38 organ is dependent on the biological half-life. The retention of radioisotope in the tissue or body organ 39 can be expressed by the relationship given by Onoja and Akpa [5] as:

$$A = A_0 e^{-\lambda \varepsilon \tau} \tag{1}$$

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43 Where A is the activity remaining at a time after the depositions of activity A_0 and ε is the effective 44 clearance constant. For practical purposes, the limiting values are reached after about half lives. At 45 this steady state condition, the activity deposited will be equal to the activity eliminated. This defines 46 the maximum concentration of any radionuclide type in drinking water. The level of concentrations of 47 radionuclides according to nature in ground waters are mainly due to uranium and thorium bearing 48 soil androck minerals or with uranium, thorium and radium deposits. Therefore studies has shown that 49 natural radioactivity in water depends on the local geological characteristics of the source, soil or rock 50 [6, 7, 8].

- Natural uranium induces chemical toxicity, especially nephrotoxicity, which is more harmful than 52 53 radiotoxicity; whereas radium and radon are thought to induce solely radiotoxicity. Higher 54 concentration of radioactivity in environmental media can cause exposure risk to the general populace 55 which may lead to radiation related sickness such as leukemia, cancer of bladder, kidney, testis and 56 lungs [4, 9]. Increased concern for the radiological status of drinking water has led to an increased 57 demand for data on water quality. World Health Organization (WHO) [10], recommended reference 58 dose level (RDL) of committed effective dose of 100 µSv from one year consumption of drinking 59 water. Gamma rays can enter the skin and interact with tissues or organs. Uranium and radium found 60 in water and do not emit strong gamma radiation, so showering with that water will not pose any 61 significant risk. However, if this radionuclide are inhaled or ingested through eating and drinking, the 62 emissions can come into direct contact with sensitive tissues or organs in the body [11, 12].
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64 Measurement of natural radioactivity levels in drinking water is relevant in assessing the radiological 65 risk to humans due to water ingestion [10, 11]. Studies of natural radioactivity of bottled water, mineral waters, ground and surface water have been the subject of numerous studies. For instance, the 66 measurement of radium isotopes (²²⁶Ra, ²²⁸Ra), ²²²Rn and ⁴⁰K concentration in bottled water and 67 68 mineral water for Poland, Austria, Romania and Algeria were presented by Nguyen et al.[12], 69 Wallneret al. [13], Elena Botezatuet al. [14]. Studies on natural radioactivity of different brands of commonly sold bottled drinking water in the federal capital Islamabad and Rawalpindi city of Parkistan 70 revealed that activity concentration of ²²⁶Ra, ²³²Th and ⁴⁰K were 11.3±2.4, 5.2±0.4 and 140.9±30.6 71 mBql¹[15]. Activity concentration of ⁴⁰K was measured using high purity germanium (HPGe) detector 72 in some sachet drinking water samples produced in Nigeria. Activity concentration of ⁴⁰K obtained in 73 74 sachet water were within the ICRP safe standard for drinking water. In Nigeria studies related to 75 natural radioactivity monitoring in ground water and surface water has been carried out [16, 17] but no 76 work has been done on sachet water, ground water and surface water from Uburu and Okposi salt 77 lake areas of Ebonyi State.

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Therefore, the aim of this work is to measure the natural radionuclide concentration in drinking water sources around Uburu and Okposi salt lake in order the quantify its associated health risk parameters for different age groups. The result of this work will serve as the base line radiological data of the study area and also add information to the radiological data base in Nigeria.

2. MATERIAL AND METHODS 2.1 STUDY AREA

87 The study area is OkposiOkwu and Uburu town located in Ohaozara LGA and are found in Lower Benue Trough which is the southern portion of Benue Trough; others are Upper Benue and Middle 88 89 Benue Trough. The geology of Lower Benue Trough is associated with tectonic activities that were 90 recorded during the Cenomanian [16]. Lead - zinc - barites mineralization in the Trough is believed to 91 be hydrothermal in origin and it is associated with brine spring [17]. The two towns lie within latitude 92 06° 02' N to 6° 07' N and Longitude 7° 42' 31" E to 7° 51' 37" E. The bedrock of the area is made up 93 of sedimentry rocks belonging to the Asu - River group of Albian age [18, 19,20, 21]. The portable 94 drinking water problem worsened during dry season when water levels and discharge from surface 95 and ground water falls due to the intense drought. OkposiOkwu salt, though believed to be medicinal 96 and relatively expensive than the normal salt and that of Uburu sold in the localities form the bulk of 97 the supply in the local markets. The salt lakes gave Ebonyi State its slogan as the "Salt of the Nation". 98 Figures 1aand 1b shows the map of OkposiOkwu and Uburu salt.

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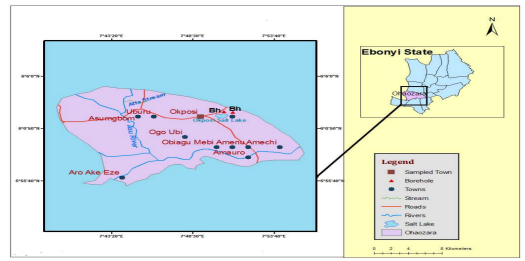
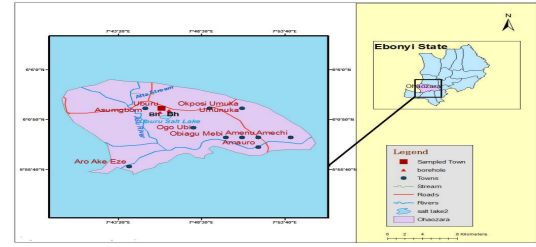


Fig. 1a: Map ofOkposiOkwu salt lake in Ohaozara LGA, Ebonyi state Nigeria
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104 105 Fig. 1b: Map showing Uburu salt lake in Ohaozara LGA, Ebonyi state Nigeria 106

107 **2.2 Sampling and Sample Preparation**

108 In order to measure the natural radioactivity in drinking water sources samples collected from Uburu 109 and Okposi lake environs, a total of thirty- one water samples were randomly collected from borehole 110 water in OkposiOkwu and Uburu, Atta stream and Asu river and also two brand of sachet water mainly distributed in the area were collected for the study.Water samples were collected using 2 litres 111 112 well labeled homogenous plastic containers. All the water samples were acidified withfew drops of 113 concentrated nitric acid (HNO_3) for each 2 litres container to obtain a pH value less than 2 (pH < 2) in 114 order to avoid adsorption of radionuclides on the walls of the container and also to prevent microbial 115 activities.Concentration was carried out by gradual evaporation of each water sample in anoven at a temperatureof 70 °C and 120 ml of the residue was transfer into a 120ml marinellibottlesand were 116 sealed with thick vinyl tapes around their screw necks. These samples were stored for 4 weeks to 117 reach secular equilibrium between ²³⁸U and ²³²Th and their respective progeny[9, 25]. 118

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120 2.3 Experimental Setup

121 The gamma ray spectrometry analyses for the water samples were carried out at the National Institute of Radiation Protection and Research (NIRPR) in University of Ibadan, Ibadan, Nigeria. After the in-122 123 growth period, each water samples was subjected to a low background gamma-ray spectrometer of type; High Purity Germanium (HPGe) P - type detector. The well calibrated, lead shielded HPGe 124 125 detector (with model number, GC8023) manufactured by CANBERRA Industries Inc, with serial 126 number: 9744 has a length and diameter of 69.8 mm and 78 mm respectively. For the water 127 analysis, the detector was connected through a preamplifier (model number: 2002CSL and serial 128 number 13000742), and a PC - based Multichannel Analyzer (MCA). The gamma spectrum peak 129 area and quantification was carried out using Genie 2K and 16K software. HPGe detector used in 130 this work has relatively higher energy resolution with relative efficiency of 80%. The standard source used for calibration was CANBERRA Multi Gamma ray Standard (MGS6M315). The energy and 131 efficiency calibrations of the detector was carried out using 1.33MeV gamma line of ⁶⁰Co resulting to 132 energy resolution of 2.3 KeV Full Width at Half Maximum (FWHM) which is considered adequate to 133 134 distinguish the gamma ray energies of interest in the present study.

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For the purpose of identifying the various radionuclides that may be present in the water samples through the gamma energies they emit, the energy calibration of the detector was performed using standard sources of known radionuclides with well – defined energies. The ²²⁶Ra and ²³²Th (²²⁸Ra) activity concentrations were determined indirectly through their activities of their decay products, while ⁴⁰K content of the water samples was also determined by measuring the 1460.8 KeV gamma rays emitted during the decay of ⁴⁰K. The detection limits of radionuclides²²⁶Ra, ²³²Th and ⁴⁰K are given as 0.03, 0.0013 and 0.002 Bql⁻¹ respectively.

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The background count was determined by counting the empty plastic container volume for 10 hours, thereafter water samples (120 ml) contained in the same container volume were counted in the HPGe detector for a period of 10 hours (36, 000 seconds) each to determine the radionuclides of interest. The net area count under the corresponding photo peaks of each of the radionuclide in the energy spectrum was computed by subtracting counts due to Compton scattering of higher peaks and the background sources from the total area of the peaks. From the measured net counts, the activity concentrations of the radionuclides in the water samples were calculated in Bql^{-1} using equation (2).

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$$A(Bql^{-1}) = \frac{C_n}{\varepsilon_{\gamma}.P_{\gamma}.t_c.V}^2$$

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where C_n is the net peak area at gamma ray energy, ε_{γ} is the efficiency of the detector, P_{γ} is the emission probability of the radionuclides of interest, t_c is the total count time(s) and V is the sample volume in litres.

157 3 Estimation of Radiation Risk parameters

The radiation risk parameters (Annual Effective dose and Excess life time cancer risk) was estimated from the activity concentrations of ²²⁶Ra, ²³²Th and ⁴⁰K in drinking water samples, dose coefficient of each of the nuclides and volume of water intake for 3 age brackets (infants, children and adults). In this work, the water intake rateswasbased on UNSCEAR [26] recommendation of 0.5 I/d and 1.0 I/d for infants (0-1 years) and children (10 years) respectively, and 2 I/d for adults (\geq 17 years) were used for calculations. The annual effective dose due to intake of drinking water sources sampled was computed using the

165 following formula [9, 25].

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$$H_{ing} (mSvy^{-1}) = \sum_{i=1}^{i=3} DCF_{ing} (i) \times Ai \times I$$
(3)

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Where DCF_{ing} (i) is the dose coefficient of a particular radionuclide in Sv/Bq for a particular age categories (Table 1). A_iis the specific activity concentration of radionuclide in the drinking water sample measured in Bq/I and I, the radionuclide intake in liters per year for each age categories.

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173 The stochastic effects of radiation in adult citizen that takes water from the various drinking water 174 sources was estimated using the international Commission on effects of protection (ICRP) cancer risk 175 methodology [28]. The health risks to members of the public due to exposure to low dose radiation 176 which is regarded as chronic risk of somatic or hereditary effects were also determined. Cancer risk 177 coefficient and hereditary effect coefficient of 5.5×10^{-2} Sv⁻¹ and 0.2×10^{-2} respectively of ICRP report 178 and assumed 70 years lifetime of continuous exposure of the population to low level radiation adopted 179 [9, 25]. According to ICRP methodology and Ndontchueng et al., [9],

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- 181 Fatality Cancer Risk = Total annual Effective Dose (Sv) × Cancer risk factor $(5.5 \times 10^{-2})(4)$
- 182 Lifetime fatality cancer risk to adult = Total annual effective dose $\times 70$ yrs $\times 5.5 \times 10^{-2}$ (5)

183 Severe Hereditary Effects = Total annual Effective Dose (Sv) × Hereditary effect factor (0.2× 10⁻²)

184 <mark>(6)</mark>

(7)

185 Estimated Lifetime hereditary effect in adult = Total annual Effective Dose (Sv) \times 70 yrs \times 0.2 \times 10⁻²

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188Table 1: Effective Dose Coefficients for ingestion of Radionuclides for members of the189public to 70 years of age (ICRP, 2012; Publication 119)

S/N	Radioisotopes	Infant ≤ 1 year	Children 10 years	Adult ≥17 years	
1	²²⁶ Ra	5.7 E-06	8.0E-07	2.8 E-07	
2	²³² Th	1.6 E-06	2.9 E-07	2.3 E-07	
3	⁴⁰ K	5.2 E-05	1.3E-08	6.2 E-09	
Water intake)	0.5 L/day	1.0 L/day	2.0 L/day	

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192 4 RESULTS AND DISCUSSION

193 The activity concentrations of ²²⁶Ra, ²³²Th and ⁴⁰K and total annual effective dose for different age 194 groups measured in water samples collected from different locations near OkposiOkwu and Uburu 195 salt lake area are presented in Table 2 while Table 3 gives the estimated cancer risks and the 196 hereditary effects of adult member of the public.

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198Table 2: Activity Concentration of 226 Ra, 232 Th and 40 K in Water Samples and Annual Effective199Dose for Different Age Categories

<mark>S/N</mark>	Sample	Location	5	oncentration	(Bq/l)	<mark>Total A</mark>	nnual Effec	<mark>tive Dose</mark>
	D		²²⁶ Ra	²³² Th	⁴⁰ K	(µSv/y) Infant	Children	Adult
1	Ubu SA01	$N06^{\circ}02'49$	BDL	0.54±0.11	BDL	<mark>0.432</mark>	0.1566	0.1242
<mark>2</mark>	UbuSA 02	E007 ⁰ 45′20.1 N06 ⁰ 02′ 19.0″	BDL	BDL	BDL			
<mark>3</mark>	UbuSA04	E007 ⁰ 46′07.9″ N06 ⁰ 02′ 17.0″	<mark>2.93±0.69</mark>	0.28±0.03	7.33±0.57	0	0	0
<mark>4</mark>	Ubu SA05	E007 ⁰ 46′ 09.9″ N06 ⁰ 02′ 18.0″	<mark>3.66±0.78</mark>	7.56±0.59	23.31±1.65	<mark>199.16</mark>	<mark>2.616</mark>	<mark>10.86</mark>
<mark>5</mark>	UbuBH01	E007 ⁰ 46′ 05.9″ N06 ⁰ 03′ 12.4″	BDL	2.22±1.35	BDL	<mark>622.54</mark>	<mark>5.726</mark>	<mark>34.43</mark>
<mark>-</mark>	UbuBH	<mark>E007⁰45′ 14.4″</mark> N06 ⁰ 03′ 13.7″	0.97±0.27	1.07±0.46	4.22±0.30	<mark>1.776</mark>	<mark>o.6438</mark>	<mark>1.021</mark>
	02	<mark>E007⁰45′ 23.9″</mark>				<mark>113.34</mark>	<mark>1.196</mark>	<mark>6.268</mark>
<mark>7</mark>	<mark>Ubu</mark> BH04	N06 ⁰ 03′ 13.7″ E007 ⁰ 45′ 24.8″	<mark>1.47±0.31</mark>	<mark>5.58±0.43</mark>	8.20±0.55	<mark>221.85</mark>	<mark>3.007</mark>	<mark>13.56</mark>
<mark>8</mark>	<mark>Ubu</mark> BH05	N06 ⁰ 03′ 13.7″ E007 ⁰ 46′ 22.9″	BDL	<mark>4.45±0.36</mark>	12.61±0.83	<mark>331.42</mark>	<mark>1.618</mark>	<mark>17.68</mark>
<mark>9</mark>	OkpSA 01	N06 ⁰ 02' 02.2" E007 ⁰ 49' 06.5"	BDL	BDL	BDL	0	0 0	<u>0</u>
<mark>10</mark>	OkpSA 02	<mark>N06⁰ 02′ 04.4″</mark>	<mark>1.11±0.62</mark>	0.14±0.03	BDL			
<mark>11</mark>	<mark>Okp SA04</mark>	E007 ⁰ 49′ 15.3″ N06 ⁰ 02′ 04.4″	BDL	<mark>7.06±0.58</mark>	18.49±1.32	3.276	0.9286	0.686
<mark>12</mark>	Okp SA05	E007 ⁰ 48′ 15.2 N06 ⁰ 02′ 04.4″	BDL	<mark>6.98±0.55</mark>	12.24±0.89	<mark>486.38</mark>	2.528	<mark>26.18</mark>
<mark>13</mark>	Okp BH	<mark>E007⁰47′ 14.1</mark> N060 02′ 07.5″	0.86±0.26	0.45±0.09	BDL	<mark>323.82</mark>	<mark>2.342</mark>	<mark>18.39</mark>
<u>14</u>	01 OKPBH04	<mark>E007048′ 4.7″</mark> N060 03′ 02.4″	BDL	2.93±0.25	10.05±0.75	<mark>2.811</mark>	<mark>0.8185</mark>	<mark>0.6886</mark>
15	Okp BH	E007049' 8.5" N060 08' 02.4"	BDL	$\frac{2.93\pm0.23}{4.81\pm0.38}$	17.83±1.22	<mark>263.64</mark>	<mark>1.111</mark>	<mark>13.81</mark>
	<mark>05</mark>	<mark>E007049' 7.5″</mark>				<mark>467.43</mark>	<mark>1.858</mark>	<mark>24.32</mark>
<mark>16</mark>	UgwBH 01	<mark>N060 08' 04.3"</mark> E007049' 3.5"	<mark>4.94±0.99</mark>	<mark>4.14±0.03</mark> 3	<mark>8.91±0.64</mark>	<mark>249.05</mark>	<mark>5.384</mark>	<mark>15.72</mark>
<mark>17</mark>	<mark>Ugw BH02</mark>	<mark>N060 07' 02.6″</mark> E007048' 7.5″	<mark>5.65±1.24</mark>	<mark>13.00±0.9</mark> 7	10.65±0.21	<mark>303.40</mark>	<mark>8.567</mark>	<mark>22.35</mark>
<mark>18</mark>	Ugw BH03	N060 06' 02.3" E007047' 6.5"	BDL	3.50±1.24	12.05±0.84	316.1	<mark>1.328</mark>	16.55
<mark>19</mark>	<mark>Ugw BH04</mark>	N060 08' 07.5" E007048' 6.2"	BDL	11.50±0.8	11.17±0.84	<mark>299.62</mark>	3.625	<u>19.14</u>
<mark>20</mark>	Ugw BH05	<mark>N060 08' 02.4"</mark>	BDL	8.88±0.71	21.30±1.50	560.90	3.129	30.5
<mark>21</mark>	<mark>Ugw BH06</mark>	E007049' 7.5" N060 08' 02.4"	<mark>1.32±0.29</mark>	<mark>5.64±0.44</mark>	<mark>26.45±1.83</mark>			
<mark>22</mark>	AttaST 01	E007049 7.5" N06 ⁰ 01' 56.4"	<mark>1.16±0.57</mark>	BDL	BDL	<mark>695.97</mark>	<mark>3.379</mark>	<mark>36.13</mark>
<mark>23</mark>	Atta ST	E007 ⁰ 48′30.7″ N06 ⁰ 01′ 58.5″	<mark>2.88±1.32</mark>	0.25±0.04	BDL	<mark>0.3306</mark>	<mark>0.928</mark>	<mark>0.6496</mark>
<mark>24</mark>	02 Atta ST04	E007 ⁰ 48′28.2″ N06 ⁰ 01′ 57.6″	BDL	8.93±0.66	11.67±1.02	<mark>8.408</mark>	<mark>2.377</mark>	<mark>1.728</mark>
		E007 ⁰ 47 28.4″				<mark>310.56</mark>	<mark>2.893</mark>	<mark>18.58</mark>
<mark>25</mark>	Atta ST05	N06 ⁰ 01' 57.4" E007 ⁰ 48'27.2"	BDL	5.23±0.41	33.32±2.18	<mark>870.50</mark>	<mark>2.383</mark>	<mark>43.72</mark>
<mark>26</mark>	Atta ST06	N06 ⁰ 01′ 58.5″ E007 ⁰ 48′28.2″	BDL	<mark>4.34±0.34</mark>	<mark>26.94±1.80</mark>	<mark>703.91</mark>	<mark>1.959</mark>	<mark>3.54</mark>

<mark>27</mark>	AsuRv 01	N06 ⁰ 03' 59.4" E007 ⁰ 44'32.1"	0.03±0.01	<mark>0.86±0.44</mark>	BDL	<mark>0.7735</mark>	<mark>0.2734</mark>	<mark>0.4124</mark>
<mark>28</mark>	AsuRv 02	N06 ⁰ 04′ 59.4″ E007 ⁰ 44′ 2.1″	2.0±0.61	0.55±0.10	BDL	<mark>614.0</mark>	<mark>1.76</mark>	<mark>1.373</mark>
<mark>29</mark>	<mark>Asu Rv04</mark>	N06 ⁰ 04' 57.5" E007 ⁰ 44' 33.2	<mark>4.48±1.13</mark>	<mark>4.32±</mark> 0.36	11.20±0.81	<mark>307.42</mark>	<mark>5.128</mark>	<mark>18.38</mark>
<mark>30</mark>	AsuRV 05	N06 ⁰ 04' 56.4" E007 ⁰ 44' 35.2	<mark>2.05±0.46</mark>	<mark>8.60±0.65</mark>	2.76±0.03	<mark>8.4482</mark>	<mark>4.206</mark>	<mark>8.526</mark>
<mark>31</mark>	<mark>Asu Rv06</mark>	N06 ⁰ 04' 59.4" E007 ⁰ 44' 30.9	<mark>2.39±</mark> 0.55	<mark>5.67±0.45</mark>	13.85±0.98	<mark>372.56</mark>	<mark>4.228</mark>	<mark>21.34</mark>

Ubu SA: Uburu sachet water samples, UbuBH:Uburu borehole water samples, Okp SA: OkposiOkwu sachet water samples, Okp BH: OkposiOkwu borehole water samples, Atta ST: Atta stream water samples, AsuRv: Asu River water samples, .BDL = Below Detection Limit

S/N	<mark>Sample</mark> code	GPS	<u>Risks and Hereditary Effect</u> Total Annual Effective Dose (μSvy ⁻¹)		Fatality cancer risk to Adult per year	Lifetime fatality cancer risk	Severe hereditary Effects in Adult per year	Estimate d lifetime hereditar y Effects	
			<mark>Infant</mark>	<mark>Childre</mark> n	<mark>Adult</mark>	× 10 ⁻⁶	<mark>× 10⁻⁵</mark>	<mark>× 10⁻⁸</mark>	<mark>× 10⁻⁶</mark>
1	Ubu SA01	N06 ⁰ 02′ 49 E007 ⁰ 45′20.1	<mark>0.432</mark>	<mark>0.156</mark>	0.124 2	<mark>0.0068</mark>	<mark>0.048</mark>	<mark>0.0248</mark>	<mark>0.0174</mark>
2	UbuSA 02	N06 ⁰ 02′ 19.0″ E007 ⁰ 46′07.9″	0	0	0	<mark>0</mark>	<mark>0</mark>	<mark>0</mark>	<mark>0</mark>
3	UbuSA04	N06 ⁰ 02' 17.0″ E007 ⁰ 46' 09.9″	<mark>199.16</mark>	<mark>2.62</mark>	<mark>10.86</mark>	<mark>0.597</mark>	<mark>4.181</mark>	<mark>2.172</mark>	<mark>1.520</mark>
4	Ubu SA05	N06 ⁰ 02' 18.0″ E007 ⁰ 46' 05.9″	622.5	5.73	34.43	<mark>1.89</mark>	<mark>13.256</mark>	<mark>6.886</mark>	<mark>4.82</mark>
5	UbuBH01	<mark>N06⁰ 03′ 12.4″</mark> E007 ⁰ 45′ 14.4″	<mark>1.78</mark>	<mark>0.64</mark>	<mark>1.021</mark>	<mark>0.056</mark>	<mark>0.393</mark>	<mark>0.204</mark>	<mark>0.143</mark>
<mark>5</mark>	<mark>UbuBH</mark> 02	N06 ⁰ 03′ 13.7″ E007 ⁰ 45′ 23.9″	<mark>113.34</mark>	<mark>1.20</mark>	<mark>6.27</mark>	<mark>0.345</mark>	<mark>2.413</mark>	<mark>1.254</mark>	<mark>0.878</mark>
7	<mark>Ubu</mark> BH04	N06 ⁰ 03′ 13.7″ E007 ⁰ 45′ 24.8″	<mark>221.85</mark>	<mark>3.01</mark>	<mark>13.56</mark>	<mark>0.746</mark>	<mark>5.219</mark>	<mark>2.712</mark>	<mark>1.898</mark>
8	<mark>Ubu</mark> BH05	N06 ⁰ 03′ 13.7″ E007 ⁰ 46′ 22.9″	<mark>331.42</mark>	<mark>1.62</mark>	<mark>17.68</mark>	<mark>0.973</mark>	<mark>6.808</mark>	<mark>3.537</mark>	<mark>2.476</mark>
)	OkpSA 01	N06 ⁰ 02' 02.2″ E007 ⁰ 49' 06.5″	<mark>0</mark>	0	0	<mark>0</mark>	0	0	0
10 1 1	OkpSA 02	N06 ⁰ 02' 04.4" E007 ⁰ 49' 15.3"	<mark>3.28</mark>	<mark>9.29</mark>	<mark>0.69</mark>	0.0377	0.264	0.137	0.0960
11 12	<mark>Okp SA04</mark> Okp SA05	N06 ⁰ 02' 04.4″ E007 ⁰ 48' 15.2 N06 ⁰ 02' 04.4″	<mark>486.39</mark>	<mark>2.53</mark>	<mark>26.18</mark>	1.44	10.077	<mark>5.235</mark>	3.66
12 13	Okp BH	E007 ⁰ 47' 14.1 N060 02' 07.5"	<mark>323.82</mark>	<mark>2.34</mark>	<mark>18.39</mark>	<mark>1.011</mark> 0.028	<mark>7.080</mark> 0.265	<mark>3.678</mark> 0.138	<mark>2.574</mark> 9.640
14	01 OKPBH04	<mark>E007048′ 4.7″</mark> N060 03′ 02.4″	2.81	0.82	0.69	0.759	5.317	2.762	0.0193
15	Okp BH	E007049' 8.5" N060 08' 02.4"	263.64	1.11	13.81	<mark>1.34</mark>	<mark>9.364</mark>	<mark>4.864</mark>	<mark>3.41</mark>
<mark>16</mark>	05 UgwBH 01	E007049' 7.5" N060 08' 04.3" E007049' 3.5"	<mark>467.43</mark> 249.05	<mark>1.86</mark> 5.38	<mark>24.32</mark> 15.72	<mark>0.865</mark>	<mark>6.052</mark>	<mark>3.144</mark>	<mark>2.20</mark>
<mark>17</mark>	Ugw BH02	E007049 3.5" N060 07' 02.6" E007048' 7.5"	249.05 303.40	8.57	22.35	<mark>1.229</mark>	<mark>8.605</mark>	<mark>4.47</mark>	<mark>3.13</mark>
<mark>18</mark>	Ugw BH03	N060 06' 02.3"	316.1	1.33	16.55	<mark>0.910</mark>	<mark>6.373</mark>	<mark>3.31</mark>	<mark>2.317</mark>

	WHO, 2004;	IAEA, 2000	<mark>0.26</mark>	<mark>0.20</mark>	<mark>0.10</mark>		**		
			<mark>13.0076</mark>	<mark>7.610E-05</mark>	<mark>472.30</mark>	<mark>2.59E-05</mark>	<mark>18.185E-</mark> 04	<mark>9.447E-07</mark>	<mark>6.613E-05</mark>
<mark>31</mark>	<mark>Asu Rv06</mark>	<mark>N06⁰ 04' 59.4″</mark> E007 ⁰ 44' 30.9	<mark>372.56</mark>	<mark>4.23</mark>	<mark>21.34</mark>	<mark>1.174</mark>	<mark>8.216</mark>	<mark>4.268</mark>	<mark>2.988</mark>
		E007 ⁰ 44′ 35.2	<mark>84.45</mark>	<mark>4.21</mark>	<mark>8.53</mark>	0.469	<mark>3.283</mark>		1.194
<mark>30</mark>	AsuRV 05	<mark>E007⁰44′ 33.2</mark> N06 ⁰ 04′ 56.4″	<mark>307.42</mark>	<mark>5.13</mark>	<mark>18.38</mark>			1.705	
<mark>29</mark>	Asu Rv04	N06 ⁰ 04' 57.5"				1.011	<mark>7.078</mark>	<mark>3.677</mark>	<mark>2.574</mark>
<mark>28</mark>	AsuRv 02	N06 ⁰ 04' 59.4" E007 ⁰ 44' 2.1"	<mark>6.14</mark>	1.76	<mark>1.37</mark>	<mark>0.076</mark>	<mark>0.529</mark>	<mark>0.275</mark>	<mark>0.192</mark>
<mark>27</mark>	AsuRv 01	N06 ⁰ 03′ 59.4″ E007 ⁰ 44′32.1″	<mark>0.77</mark>	<mark>0.27</mark>	<mark>0.41</mark>	<mark>0.023</mark>	<mark>0.158</mark>	<mark>0.0083</mark>	<mark>0.058</mark>
		E007 ⁰ 48'28.2"	<mark>703.91</mark>	<mark>1.96</mark>	<mark>35.4</mark>	<mark>1.947</mark>	<mark>13.63</mark>	<mark>7.08</mark>	<mark>4.956</mark>
<mark>26</mark>	Atta ST06	E007 ⁰ 48′27.2″ N06 ⁰ 01′ 58.5″	<mark>870.50</mark>	<mark>2.38</mark>	<mark>43.72</mark>				
<mark>25</mark>	Atta ST05	<mark>N06⁰ 01′ 57.4″</mark>				<mark>2.405</mark>	<mark>16.833</mark>	<mark>8.745</mark>	<mark>6.12</mark>
<mark>24</mark>	Atta ST04	N06 ⁰ 01′ 57.6″ E007 ⁰ 47 28.4″	<mark>310.56</mark>	<mark>2.89</mark>	<mark>18.58</mark>	<mark>1.022</mark>	<mark>7.153</mark>	<mark>3.716</mark>	<mark>2.601</mark>
<mark>23</mark>	<mark>Atta ST</mark> 02	E007 ⁰ 48'28.2"	<mark>8.41</mark>	<mark>2.38</mark>	<mark>1.73</mark>	<mark>0.095</mark>	<mark>0.665</mark>	<mark>0.346</mark>	<mark>0.242</mark>
		<mark>E007⁰48'30.7″</mark> N06 ⁰ 01' 58.5″	<mark>3.31</mark>	<mark>0.93</mark>	<mark>0.65</mark>				
<mark>22</mark>	AttaST 01	E007049 7.5″ N06 ⁰ 01′ 56.4″				0.036	0.250	0.130	0.091
<mark>21</mark>	Ugw BH06	<mark>N060 08' 02.4</mark> "	<mark>695.97</mark>	<mark>3.38</mark>	36.13	<mark>1.987</mark>	<mark>13.911</mark>	<mark>7.226</mark>	<mark>5.058</mark>
<mark>20</mark>	<mark>Ugw BH05</mark>	N060 08' 02.4" E007049' 7.5"	<mark>560.90</mark>	<mark>3.13</mark>	<mark>30.5</mark>	<mark>1.677</mark>	<mark>11.741</mark>	<mark>6.099</mark>	<mark>4.270</mark>
		<mark>E007048' 6.2″</mark>	<mark>299.62</mark>	<mark>3.63</mark>	<mark>19.14</mark>	<mark>1.053</mark>	<mark>7.369</mark>	<mark>3.828</mark>	<mark>2.679</mark>
<mark>19</mark>	Ugw BH04	E007047' 6.5" N060 08' 07.5"				1 052	7.200	2.020	2 670

From Table 2, the specific activity concentration of ²²⁶Ra,²³²Th and ⁴⁰K ranged from BDL to 3.66± 0.78 210 Bql⁻¹, BDL to 7.56±0.59 Bql⁻¹ and BDL to 23.31 ±1.65 Bql⁻¹ respectively in sachet water. The average 211 activity concentration of²²⁶Ra and ²³²Th in sachet water produced in OkposiOkwu and Uburu are 212 213 found to be higher than 0.02 Bql⁻¹ and 0.03 Bql⁻¹ in mineral bottled water produced in Cameroon [25] except for ⁴⁰K.The activity concentration of ²²⁶Ra, ²³²Th and ⁴⁰K in borehole water ranges from BDL to 214 215 5.65± 1.24, 0.45±0.09 to 13.00±0.97 and BDL to 26.45± 1.83 Bql⁻¹ respectively. Furthermore the activity concentration of ²²⁶Ra,²³²Th and ⁴⁰K in stream water ranges from BDL to 2.88±1.32, BDL to 216 8.93±0.66 Bql⁻¹ and BDL to 33.32± 2.18 Bql⁻¹ respectively, while that for river water ranges from 217 0.03±0.01 to 4.48±1.13 Bql⁻¹, 0.55±0.10 to 8.60±0.65 Bql⁻¹and BDL to 13.85 ± 0.98 Bql⁻¹ respectively. 218 The variations in activity concentrations of these radionuclides are due to the variations in the 219 chemical composition of local geological formations and the aquifer geochemistry from where the 220 drinking water originate. The presence of the salt lake might have contributed to increase 221 concentration of ²²⁶Ra, ²³²Th and ⁴⁰K in stream water near the salt lakes. 222 223

The highest activity concentration value of ²²⁶Ra (5.65±1.24) was recorded in borehole water 224 (UgwBH02) which could be due to infiltration from the salt lakes and other activities in the area. The 225 mean activity concentrations of ²²Ra, ²³²Th and ⁴⁰K obtained in all the different water resources were 226 1.24, 4.17 and 9.82 Bql⁻¹ respectively which is slightly higher than the reference value of 1.0 and 0.1 227 Bql⁻¹ recommended by WHO [6] except for ⁴⁰K which is within the safe value. The average results of 228 both OkposiOkwu and Uburu borehole water samples were lower than the Tap water results 229 230 measured by Ononugbo et al., [29] at Ogba/Egbema/Ndoni LGA of Rivers State in oil producing 231 communities, Niger Delta Region of Nigeria due to different geological composition the areas and the 232 oil producing activities in Onelga. However, the obtained results were higher than the results of Osman et al., [30] who studied natural radioactivity levels of ground waters of Kuhliate and Miri Bara 233 in Kadugli, Saudi. The variation in the results is traceable to their local geology and geochemistry of 234 235 the aquifer as well as the environmental management practices. The levels of gamma radiation in 236 ground water sources could directly be associated with the mineralogical compositions and activity

concentrations of radionuclide in aquifer bedrock and the age of the ground water in the aquifer. The
 result of this study also show that the activity concentration of ²²⁶Rais higher in Atta stream than Asu
 river while activity concentration of ²³²Th in Asu river is higher than that of Atta Stream and also high
 than the result obtained by Jibiri*et al.*,[31] from Obafemi – Owode area in Abeokuta, Nigeria.

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242 The annual effective dose due to ingestion of drinking watersourcessampled was estimated for three 243 different age groups: Infants, children and adults using equation 3 and presented in Table 2. The calculated annual effective dose for different age groups that drinks sachet water ranges from 0.03 to 244 0.20 mSvy⁻¹ for infants, 0.029 to 0.04 mSvy⁻¹ for children and from 0.038 to 0.036 mSvy⁻¹ for adult 245 respectively. In borehole water, it ranges from 0.028 mSvy⁻¹ to 0.695 mSvy⁻¹ in infants, 0.008 to 0.009 246 mSvy⁻¹ for children and 0.0069 to 0.036 mSvy⁻¹ in adult. For stream water, total effective dose ranges 247 248 from 0.0033 to 0.871 mSvy⁻¹, in infant, 0.009 to 0.034 in children and 0.0065 to 0.044 mSvy⁻¹ in adult. 249 The total annual effective dose calculated from activity concentration of radionuclides in river water samples for infant, children and adult ranges from, 0.0077 to 0.37 mSvy⁻¹, 0.0027 to 0.051 mSvy⁻¹ and 250 251 0.0041 to 0.021 mSvy⁻¹ respectively. It can be observed that the radiation dose received by infants is 252 relatively higher than that received by children and adults. The total annual effective dose obtained 253 for infant, children and adults are higher than that obtained by Yussufet al., [1] but are within the result 254 obtained by Ajayiet al., (2009) and WHO[6] and UNSCEAR [26] reference levels of the effective dose 255 for infants, children and adult due to one year continuous ingestion of various drinking water of 0.26, 0.20 and 0.10 mSvy⁻¹ respectively. The effective doses obtained are higher than the reference values 256 257 for infants, children and adults that consume stream water and river water and from the radiation 258 protection point of view, life-long ingestion of these sampled drinking waters may cause significant 259 radiological health risk. It is observed that infants have higher radiation risk than children and adult 260 since the total effective dose calculated for infants exceeded the WHO reference value in all drinking 261 water sources studied.

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Fatality cancer risk and severe hereditary effects of ingestion of the sampled drinking water sources 263 264 was determined and presented in Table3. The result showed that cancer risk for adults varies from 0.0068×10^{-6} to 2.41 $\times 10^{-6}$ and the lifetime hereditary effects varies between 0.046 $\times 10^{-5}$ and 16.83 \times 265 10^{-5} . Whereas the hereditary effect to adult per year calculated varied from 0.025 x 10^{-8} to 8.75x 10^{-8} 266 and the lifetime hereditary effectshereditary effect in adult varies from 0.017 \times 10⁻⁶ to 9.64 \times 10⁻⁶. This 267 268 implies that 16 out of 100,000 may likely suffer some form of cancer fatality and the result also shows that 9 out of 100,000 may suffer some hereditary effects. The United States Environmental protection 269 Agency (USEPA) recommended acceptable cancer fatality risk limit of 1.0×10^{-6} to 1.0×10^{-4} (i.e. 1) 270 271 person out of 1 million to 10,000 persons suffering from some form of cancer fatality) [9, 34].

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The result of the lifetime cancerrisk and the lifetime hereditary effect obtained in this study are in agreement with that obtained by Ndontchueng*et al.*[9] in mineral waters in Cameroon. The obtained results are within the acceptable risk factor as recommended by USEPA [34].

277 5. STATISTICS

Statistical analysis of the measured activity concentration of ²²⁶Ra, ²³²Th and ⁴⁰K in water samples are 278 279 Presented in Table 4 while the histograms are presented in figure 3. When the standard deviation is 280 higher than the mean value, it shows low degree of uniformity and vice versa. In this present study, standard deviation values of activity concentrations of ²²⁶Ra, ²³²Th and ⁴⁰K are higher than the mean 281 282 values indicating low degree of uniformity. Skewnessrefers to asymmetric nature of the shape of 283 frequency distribution. Skewed distribution could either be positively or negatively skewed [34]. From Table 4, the skewness of the activity concentration of ²²⁶Ra, ²³²Th and ⁴⁰K are positive which shows 284 that their distributions are asymmetric. 285

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Table 4: DESCRIPTIVE STATISTICS

Statistics							
		Ra226	Th232	K40	AEDEinfant	AEDEchildren	AEDEadult
N	Valid	31	31	31	31	30	31
	Missing	32	32	32	32	33	32
Mean		1.2226	4.1742	9.8242	279.3492	2.5151	13.7631
Std. E	Error of Mean	.29768	.65303	1.69532	45.34643	.35395	2.20893
Media	an	.0300	4.3200	10.0500	299.6200	2.3595	13.8100
Mode		.00	.00	.00	.00	.00	.00
Std. [Deviation	1.65743	3.63594	9.43915	252.47822	1.93866	12.29880
Varia	nce	2.747	13.220	89.098	63745.249	3.758	151.261
Skew	ness	1.332	.625	.754	.565	1.172	.614
Std. E	Error of Skewness	.421	.421	.421	.421	.427	.421
Kurto	sis	.870	298	123	518	1.913	342
Std. E	Error of Kurtosis	.821	.821	.821	.821	.833	.821
Rang	e	5.65	13.00	33.32	870.50	8.57	43.72
Minim	num	.00	.00	.00	.00	.00	.00
Maxir	num	5.65	13.00	33.32	870.50	8.57	43.72
Sum		37.90	129.40	304.55	8659.83	75.45	426.66

Skewness is the extent to which the data are not symmetrical. From Table 4 and figure 2, the obtained data are positive skewed or right skewed data since the tail of the distribution points to the rightand its skewness value greater than zero. It means that the frequency of positive returns exceeds that of negative returns resulting in the distribution displaying a fat right tail or positive skewness. Kurtosis is a measure of peakedness of the distribution curve. Kurtosis indicates the extent to which the values of the variables fall above or below the mean and manifests itself as a fat tail. With the exception of ²²⁶Ra and AEDE children, all the other data had negative kurtosis. The negative value of kurtosis indicates less peaked than normal curve and is called platykurtic [34]. This indicates that returns veryhigh above or below the mean occurred very frequently and the distribution exhibits high kurtosis. It has a flattened shape.²²⁶Ra and AEDE_{children} showed a positive kurtosis. This implies that there are lesser returns above or below the mean and the frequency of occurrences increases around the mean and the distribution shows low kurtosis, in order words, it is leptokurtic. This distribution has high peak.

In order to determine the mutual relationships and strength of association between pairs of variables,
 correlation between them were drawn using SPSS 16.0 software as shown in Table 5. Low positive
 correlation was observed between ²²⁶Ra and ⁴⁰K and ²³²Th and ⁴⁰K. This is due to the fact that ²²⁶Ra

and ²³²Th comes natural decay series whereas ⁴⁰K, though a naturally occurring radionuclide is not
 part of any such decay series. This indicates that ⁴⁰K concentrations may not be related with the
 presence of ²³²Th and ²²⁶Ra bearing minerals. Weak negative correlation coefficient was observed
 between ²²⁶Ra and ²³²Th shows that their sources in the environment differs.



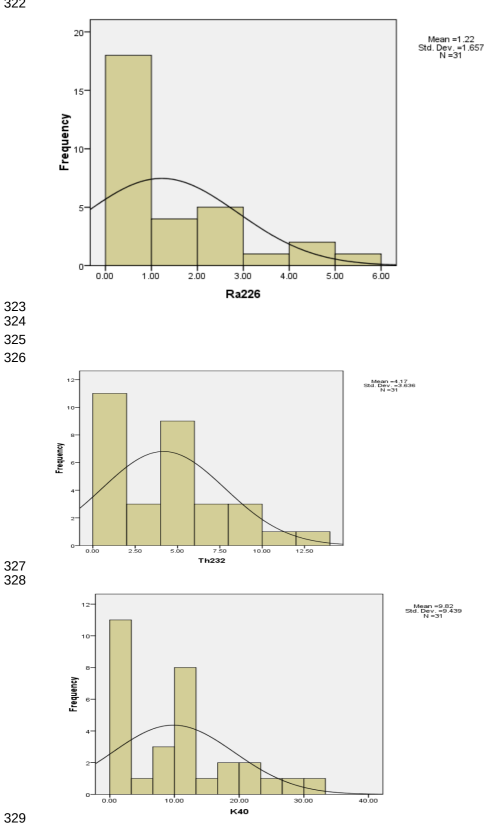


Fig.2: Frequency distributions of ²²⁶Ra, ²³²Th and ⁴⁰K in drinking water sources

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- 336

Table 5: Pearson's Correlation Coefficient Analysis 337

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			Correlation	IS	<u>.</u>		
		Ra226	Th232	K40	AEDEinfant	AEDEchildren	AEDEadult
Ra226	Pearson Correlation	1	.187	046	.009	.780**	.122
	Sig. (2-tailed)		.314	.808	.960	.000	.514
Th232	Pearson Correlation	.187	1	.557**	.464**	.744**	.677**
	Sig. (2-tailed)	.314		.001	.008	.000	.000
K40	Pearson Correlation	046	.557**	1	.902**	.381 [*]	.868**
	Sig. (2-tailed)	.808	.001		.000	.038	.000
AEDEinfant	Pearson Correlation	.009	.464**	.902**	1	.354	.778**
	Sig. (2-tailed)	.960	.008	.000		.055	.000
AEDEchildren	Pearson Correlation	.780**	.744**	.381 [*]	.354	1	.551**
	Sig. (2-tailed)	.000	.000	.038	.055		.002
AEDEadult	Pearson Correlation	.122	.677**	.868**	.778**	.551**	1
	Sig. (2-tailed)	.514	.000	.000	.000	.002	
	Ν	31	31	31	31	30	31

**. Correlation is significant at the 0.01 level (2-tailed).

*. Correlation is significant at the 0.05 level (2-tailed).

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6. CONCLUSION 340

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The natural radioactivity level of ²²⁶Ra, ²³²Th and ⁴⁰K have been estimated in various water resources 342 of Uburu and Okposi salt lake area of Ebonyi state using high purity Germanium based gamma 343 spectroscopy. This study showed slight elevation of activity concentration of ²²⁵Ra, ²³²Th and ⁴⁰K in 344 all the water samples. ²³²Th contributed the largest activity concentration and ⁴⁰K the least compared 345 to their respective reference values. 346

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348 The use of water samples that have been investigated in this study show much lower internal exposure than the WHO and ICRP reference limits of 0.10 mSvy⁻¹ and 1.0 mSvy⁻¹ respectively for 349 children and adult but slightly higher in infants. Fatality risk and hereditary risk analysis showed that 350 351 only 16 out of 100,000 adult persons exposed to these water studied might develop cancer in their 352 lifetime and 9 out of 100,000 may suffer some hereditary effects. Therefore all the water sources sampled are safe to be used by children and adult humans either as drinking or daily routine activities 353 354 but not suitable for infants. This study provided a data base on environmental radioactivity burden of the water resources of the study area. 355

356 357 358 359 REFERENCES 360 361 Yussuf, N.M., Hossain, I. and Wagiran, H. Natural radioactivity in drinking and mineral water [1] 362 in Johor Bahru , Malaysia. Scientific Research and Essays 7(9):1070-1075, 2012. 363 Guogang J., Giancario, T., Leandro M. Concentrations of ²³⁸U, ²³⁵U, ²³²Th, ²³⁰Th, ²²⁸Th, ²²⁴Ra, ²¹⁰Po and ²¹²Pb in drinking water in Italy. Reconciling safety standard based on measurement 364 [2] 365 of gross alpha and beta. J. Environ. Rad., 100: 941-949, (2009). 366 367 368 Onoja, R.A., Akpa, T.C. Gross Alpha and Beta radioactivity in drinking water from some [3] 369 towns in North Western Nigeria. Nigerian Journal of Physics, 20 (1) pp: 44-48, (2008). 370 371 AjayiO.S. ,Achuka Radioactivity in drilled and dug well drinking water of Ogun State, South-[4] 372 Western Nigeria and consequent dose estimates. Rad. Prot. Dos., 135 (1) 54-63.(2009). 373 374 Avwiri, G.O , Onunugbo, C.P.,, Egieya, J.M. Evaluation of natural radionuclide content in [5] 375 surface and ground water and excess lifetime cancer risk due to gamma radioactivity. 376 Academic Research International, 4(6): 636 – 647, 2013 377 378 [6] World Health Organization,. Guidelines for Drinking Water Quality. Third Edition 379 Incorporating the first and second Addenda, Volume 1, Recommendations; WHO Geneva pp 380 1 - 200, 2008381 382 Yussuf, N.M. Hossani, I. and Wagiran Natural radioactivity in drinking and mineral water in [7] JahorBahru (Malaysia). Scientific Research and Essays vol.7 (9) pp; 1070 -1075. Academic 383 384 Journals, 2012. Guogang J., Giancario, T. Estimation of Radiation Doses to members of the public from 385 [8] intakes of some important naturally occurring radionuclides. J. Environ. Rad., 76:654-72, 386 387 2007. 388 389 [9] Ndontechueng, M.M., Suno A. Nguelem , E.J.M. Beyala, J.F. and Kryezie, D. (2013). Preliminary study of Natural Radioactivity and Radiological risk assessment in some mineral 390 391 bottled water produced in Cameroon. International Journal of science and Technology 392 3(3):271-276. 393 World Health Organisation (2008). [10] 394 395 Amrani D. Natural radioactivity in Algerian bottled water. Journal of Radio analytical and [11] 396 Nuclear Chemistry, Vol. 252(3):597-600, 2002. 397 Nguyen Dinh Chau, Barbara Michalec. Natural radioactivity in bottled natural spring mineral 398 [12] 399 and Therapeutic waters in Poland. Journal of Radio analytical and Nuclear Chemistry, 279 91): 121-129.2009. 400 401 Wallner G., G. Steininger. Radium isotopes and ²²²Rn in Austrian drinking waters. Journal of 402 [13] 403 Radioanalytical and Nuclear Chemistry, Vol. 274, No.3:511-516.2007. [14] 404 Elena, Botezatu Olga Iacob, Angela Aflorei, GarofitaElisei, Olga Capilanu. Natural 405 radioactivity of some mineral waters and population doses. Journal of preventive medicine 406 9(3):3-14.2001. 407 [15] 408 Ononugbo, C.P. and Tutumeni. Natural Radioactivity and radiation dose estimation in 409 various water samples in Abua/Odua Area, Rivers state, physical science Journals 11, (4) 1-410 12,2016. 411 412 [16] Avwiri, G.O., Ononugbo, C.P., Nwokeoji, I.E. Radiation hazard indices and excess lifetime 413 cancer risk in soil, sediment and water around Mini - Okoro/Oginigba Creek Port Harcourt,

- 414 Rivers State, Nigeria. Comprehensive Journal of Environmental and Earth Sciences, 3(1):
 415 38 50.2014.
 416
- 417 [17] Fatoye, F.B., and Gideon, Y.B. Geology and mineral resources of the Lower Benue Trough,
 418 Nigeria. Advances in Applied Science Research, 4(6): 21 28.2013.

427

435

447

468

- Fatoye, F.B., Ibitomi, M.A., Omada, J.I. Lead zinc barites mineralization in Benue Trough,
 Nigeria: Their geology, occurrence and economic prospective. Advances in Applied *Science Research*, 5(2): 86 92. 2014.
- 424 [19] Egboka, B.C.E., and Uma, K.O., Hydro chemical contaminant transport and tectonic effects
 425 in the Okposi Uburu salt lake area of Imo State, Nigeria. *Hydrological Science Journal*,
 426 31(2): 205 221,1986.
- 428 [20] Okoyeh , E.I, &Egboka, B.C.E, Evaluation of hydro chemical parameters of Okposi and
 429 Uburu salt lakes, Nigeria. *International Journal of Scientific and Engineering Research*, 4(6):
 430 1 7.2013.
- 432 [21] Obasi, P.N.I., andAkudinobi, B.E.B. Hydrochemical evaluation of water resources of the
 433 Ohaozara area of Ebonyi State, Southeastern, Nigeria. *Journal of Natural Science*434 *Research*, 3(3): 75 50. 2013.
- 436 [22] Okoye, E.I., Akpan A.E., Egboka, B.C.E., Okolo, M.C., andOkeke, H.C. Geophysical delineation of sub surface fracture associated with Okposi Uburu salt lake, Southeastern, Nigeria. *International Research Journal of Environmental Sciences*, 4(2):1 6.2015.
 439
- 440 [23] Umunnakwe, J.E. and Aharanwa, B.C. Assessment of water quality and heavy metals levels 441 of fish species in Ogutalake, Imo State Nigeria. *Journal of Natural Sciences Research*, 4(8): 442 103 – 112. 2014.
- Taher, A.G. andSolimara, A.A. Heavy metal concentration in surficial sediments for Wadi El Natrun Saline Lakes, Egypt. *International Journal of Salt Lake Research,* 8 (1): 74 92.
 1999.
- 448[25]Tug, G.N. and Duman, F. Heavy metals accumulation in soils around a salt lake in Turkey.449Pak. J. Bot., 42(4): 2327 2333. 2010.
- 450 [26] Ndontchueng, M.M., Simo, A., Njuelem, E.J.M., Beyala, J.F., Kryeziu, D. Preliminary study
 451 of natural radioactivity and radiological risk assessment in some mineral bottled water
 452 produced in Cameroon. International Journal of Science and Technology, 3(5): 271 –
 453 276.2013.
- 455 [27] UNSCEAR,. Sources and effect of ionizing radiation. Unscear, 2008 report to the general
 456 Assembly with scientific Annesvol II.2008
 457
- International Commission on Radiological Protection, ICRP. Age dependent doses to members of the public from intake of radionuclides: Part 5. Compilation of Ingestion and Inhalation dose co – efficient, ICRP Publication 72, Pergamon Press, Oxford.(1996).
- 462 [29] International Commission of Radiological Protection (ICRP). 2006 recommendations of the
 463 (ICRP) 103 Publication. Pegammon Press,2007
 464
- Isinkaye, M.O. and Emelue, H.U. Natural radioactivity measurements and evaluation of radiological hazards in sediment of Oguta Lake, South East Nigeria. *Journal of Radiation Research and Applied Sciences,* volume 8, issue 3: 459 469, 2015.
- 469 [31] Onoja, R.A., Daniel, J.A. Sunday, O. Physical parameters a total radioactivity concentration in some borehole water. *Archives of Applied Science Research*, 5(3): 211 219,2013.
- 472 [32] Jibiri, N.N., Chijioke, M.N., George, O.A. Radionuclide contents and physicochemical water 473 quality indicators in stream, well and borehole water sources in high radiation area of

474 475 476		Abeokuta, Southwestern Nigeria. Journal of Water resources and Protection, 2: 291 – 297,2010.
477 478 479 480	[33]	International Commission of Radiological Protection (ICRP). Compendium of Dose Coefficients based on ICRP publication 60. ICRP publication 119. Ann. ICRP 4(Suppl.), 2012)
481 482 483	[34]	US-EPA .2012 Edition of the Drinking Water Standards and Health Advisories. EPA 822-S- 12-001. Washington DC,2012.
484 485 486 487	[35]	Grupta S.P. "Statistical methods" Sultan Chand & sons, Educational publishers, New Delhi . 2001.