<u>Original Research Article</u> Evaluation of Radiation Health Risk Due to Gamma Exposure From River Water Around Oil Bunking Centre In Rivers State, Nigeria

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ABSTRACT:

Aim: The aim of this study is to assess radiation health risk due to gamma exposure from river water around oil bunkering centers in Rivers state, Nigeria. Study design: This study was purely an experimental work. Place and Duration of Study: Sampling started from the meeting point of Otamiri tributary and Imo River at the Abia/Rivers boundary to over seven kilometer along the Imo River; between July - January, 2017. Methodology: 20 samples of river water were collected along coastal shore of Imo River with pre-washed 1.5 ml Polypropylene bottles. The bottles were rinsed with the water before collection and acidified immediately after collection with few drops of nitric acid. The bottles were sealed tightly with vinyl tapes and kept in the laboratory for 4weeks for secular equilibrium of the radionuclides. The activity concentration of the radionuclides were measured using well calibrated Sodium lodide detector. **Results:** The mean activity concentrations of ²³⁸U, ²³²Th and ⁴⁰K were 2.02±0.02, 3.59±0.21 and 10.43±1.13 BqL⁻¹. The mean annual effective dose estimated for infants, children and adult citizen that ingest river water sampled were 58.64, 0.19 and 0.24 mSvy⁻¹ respectively. The values of annual effective dose for infant and adults exceeded the reference level of 0.26, 0.2 and 0.10 mSvy⁻¹respectively while that for children are within the safe reference level. The estimated fatal cancer risk to adults citizen and the lifetime hereditary effects show that 53 out of 10,000 citizens may suffer some form of cancer fatality and 596 out of 1000,000 citizens may suffer some form of hereditary effect since the values exceeded the USEPA recommended range. Conclusion: The result of this study show that the river water under study have been radiologically impacted by oil bunking activities and may cause significant health risk. Hence few recommendations were made in this work which will help to reduce radiation exposure and possible health impact.

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9 10 Keywords: Radionuclide, lifetime cancer risk, committed dose, hereditary effect, Imo river..

1. Introduction

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12 River water does not exist in a pure form for any appreciable length of time in nature. Even while waterfalls as rain, water picks up small amount of contaminants from the atmosphere and 13 move as it filters through the ground^[1]. Those contaminants may be natural or anthropogenic 14 including biological, chemical, physical and radiological impurities such as industrial and 15 16 commercial solvents, heavy metals, acid salts, and radioactive materials. The major sources of natural radionuclides in water results from weathering and recycling of terrestrial minerals and 17 rocks that give rise to ⁴⁰K, ²³²Th, ²³⁵U and ²³⁸U. The later three decay naturally to produce other 18 19 important radioactive isotopes of elements including radium, radon, polonium and lead ^[2, 3].

Water can pick up radioactive materials as it flows through rocks, soils or cracked cement surrounding of water source thereby contaminating the water sources.

22 In a closed system the progeny of thorium and uranium are present in concentrations determined by the concentration of parent uranium and thorium isotopes and the time since the 23 system became closed to nuclide migration. In nature closed systems rarely exist and 24 25 predictions regarding nuclide concentrations in water bodies invariably include large 26 uncertainties. These nuclides and their decay products are found in ground and spring waters in 27 element specific concentrations dependent on complex hydrogeologic processes and conditions (dissolution, transport and ion-exchange processes as well as redox potentials and pH-28 conditions of the aqueous system). These hydrogeological processes result in non-equilibrium 29 conditions between parent nuclides and their progeny. However, characteristic behaviour in the 30 31 natural environment can provide a basis for assumptions regarding probable behaviour of nuclides used in the radioactivity screening assessment ^[4]. 32

33 In the oxidised zone of the earth's near-surface environment thorium and uranium may both be mobilised, but in different ways. Thorium has an extremely low solubility in natural waters. 34 There is a close correlation of thorium concentration and detrital content of water. Thorium is 35 almost entirely transported in particulate matter and is bound in insoluble resistant minerals or is 36 adsorbed on the surface of clay minerals. Even when thorium is generated in solution by 37 38 radioactive decay of U-234 it rapidly hydrolyses and adsorbs on to the nearest solid surface. Products of radioactive decay in the U and Th series include radon (Rn) gas of which 39 40 three isotopes exist. Rn-222 is the longest-lived and most abundant. Loss of radon will cause disequilibrium between members of a decay chain. Rn-222 has an appreciable solubility in 41 42 water and is often found in concentrations far in excess of the parent nuclide radium, Ra-226. An Rn-222/Ra-226 activity ratio of 450 has been observed in ground waters from central 43 England^[5]. Aeration of water and short half-lives make the contribution of radon negligible in 44 ingestion dose calculations. 45

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Illegal bunkering activities and crude method of refining crude oil along Imo river course has 47 48 introduced a lot of hazardous waste into the water bodies of the area. Recently the entire Rivers state is experiencing massive air pollution (black soot) covering everywhere within Port Harcourt 49 metropolis. Some speculations are pointing towards the illegal refining of crude oil in all those oil 50 bunkering centres which produces some kind of explosions in the process. The inadvertent 51 discharges of petroleum hydrocarbons or petroleum derived wastes streams from oil and gas 52 productions activities are toxic to the coastal waters, soils and sediment near the discharge 53 point^[6]. For human race, water is essential to life as air to breath. Thus, the importance of 54 investigating the levels of radionuclide element in river water is very important ^[1] as river water 55 serves as a major source of drinking water to many communities along Imo River. Estimation of 56 57 radiation dose distribution is vital in assessing the health risk to a population and serves as a 58 reference for documenting changes in environmental radioactivity due to anthropogenic activities^[7]. Hence, the aim of this work is to determine the radiological health risk of the 59 populace from the activity concentration of ²³⁸U, ²³²Th and ⁴⁰K in river water collected from Imo 60

61 River near the bunkering sites in Rivers state. The result will help in assessment of the health 62 impact of oil bunkering activities in Rivers state.

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64 **2.** Materials and Methods

65 2.1 Study Area

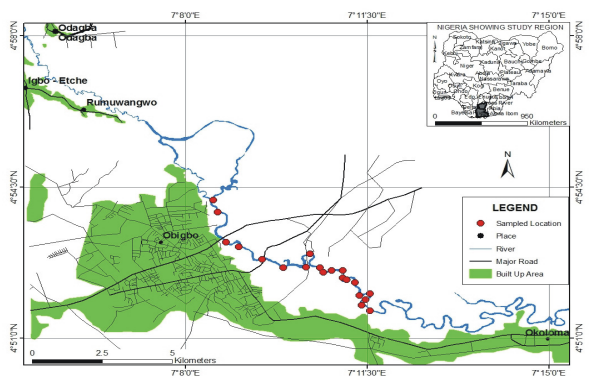
The Imo River is located in the northern part of Rivers State in South eastern Nigeria. The study area is the boundary between Abia State and River State in the Niger Delta region. It lies between longitude 007° 08¹ 11.9¹¹ and 007° 11¹ 35.5¹¹ East and latitudes 04° 54¹ 11.9¹¹ and 04° 51¹ 37.8¹¹ North of equator (Figure 1). It is in southeastern Nigeria and flows 240 km into the Atlantic Ocean. Its estuary is around 40 km wide and the river has an annual discharge of 4 km³ with 26,000 hectares of wetlands. The Imo's tributaries are the Otamiri and Oramirukwa[8]. The Imo River serves as drinking water sources for the surrounding communities.

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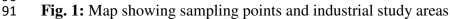
Two geologic formations are covered in the study area, namely: Imo shale and Ameki formations

76 respectively. Imo shale consists of a thick sequence of blue and dark grey shales with occasional bands of clay-ironstones and subordinate sandstones ^[9]. It dips at angles 17° to 25° 77 to the south-west and South^{[10].} It includes three constituent sandstones: the lgbabu, Ebenebe 78 79 and Umuna Sandstones with the last two outcropping in the Imo River Basin. The Umuna 80 sandstone iscomposed of thick sandstone units and minor shales and is generally less than 70m thick. The EbenebeSandstone occurs as a lens in the northwestern extremity of the Imo 81 River Basin. It is similar in lithology to the Umuna sandstone but is relatively thicker with a 82 maximum thickness of 130m^[10]. Ameki Formation (Eocene) consists of sand and sandstones. 83 The lithologic units of the Ameki Formation fall into two general groups ^[11, 12, 13]; an upper grey-84 85 green sandstones and sandy clay and a lower unit with fine to coarse sandstones, and 86 intercalations of calcareous shales and thin shelly limestone. 87

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92 2.2 Sample Collection and Preparation

Sampling started from the meeting point of Otamiri tributary and Imo River at the Abia/Rivers 93 boundary to over seven kilometer along the Imo River. In order to measure the specific activity 94 concentration of natural radionuclides in river water, 20 samples of the water were collected. All 95 the water samples were collected with 1.5 litres linear polypropylene bottles which has been 96 carefully washed before sampling. Containers for the samples were washed with a solution of 97 98 detergent and then rinsed with freshly distilled Hydrochloric acid (HCI) to remove an inorganic 99 material that might have stuck to the walls of the container before the samples were collected. About 1.5 L of each water sample were taken and 20 mL of 1 M HNo₃ added immediately ^[14]. 100 101 This is necessary to fix the radioactive elements in the samples. The samples were taken to the National Institute of Radiation Protection and Research (NIRP & R) University of Ibadan for 102 preparation and analysis. The containers were sealed tightly and wrapped with thick vinyl tapes 103 104 around their screw necks. Some 250 mL of each water samples in tightly covered cylindrical containers were stored for 4 weeks to reach secular equilibrium between ²³⁸U and ²³²Th and 105 106 their respective progeny.

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108109 2.3 Gamma Spectroscopy

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111 Activity measurements of radionuclides in river water collected were performed at the National 112 Institute of Radiation Protection and Research Centre (NIRP &R), university of Ibadan with a 113 gamma-ray spectrometry system with a thallium activated $3^{"} \times 3^{"}$ sodium iodide on a NaI(TI)

detector connected to ORTEC 456 amplifier ^[15,16]. The detector in a 100 mm thick lead shield, 114 was connected to a computer program called SAMPO 90 window that matched gamma 115 energies to a library of possible isotopes. Since the accuracy of the quantitative measurements 116 is dependent on the calibration of the spectrometry system and adequate energy, background 117 118 measurement and efficient calibration of the system was made using Cs-137 and Co-60 standard sources from IAEA, Vienna. The analysis was performed using a Canberra S 100 119 computer analyzer. Standard of natural origin were prepared in the same manner as the 120 121 samples, these standards are uranyl nitrate (UO(2), (NO,), 6H,O) 502.18 mol/g, potassium chloride (Kcl) 74.55 mol/g and thorium nitrate (Th $(NO_3)_4$.5H₂O) 570.13 mol/g. One gram of 122 123 each of the standard was taken and dissolved into a 200 mL distilled water to form a standard solution. It is subtle that 1 g of uranyl nitrate contains 0.474 g of uranium which has activity of 124 125 0.0294 Bq/L, also 1 g of potassium chloride contains 0.534 g of potassium which has activity of 0.706 Bg/L and 1g of thorium nitrate contains 0.859 g of thorium with activity of 0.0175 Bg/L^[17]. 126

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Spectrum were accumulated for background for a period of 29000 s at 900 volts to produce strong peaks at gamma emitting energies of 1460 Kev for ⁴⁰K; 63.0 Kev of ²¹⁴Bi and 92.5 Kev of ²²⁸Ac, which were used to estimate the concentration of ²³⁸U and ²³²Th respectively^[16]. The detector was calibrated with cesium-137 and cobalt-60 sources and the energy resolution is 39.5 and 22.2%, respectively ^[18]. The configuration and geometry was maintained throughout the analysis

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135 3. Radiological Risk Estimation

The annual effective dose from ingestion of radionuclide in water samples was estimated on the basis of the mean activity concentration of the radionuclides. This was done for different age categories. Assumptions on the rate of ingestion of water were made. In this work, the rate of water intake rates based on UNSCEAR ^[19] recommendation of 0.5, 1.0 and 2.0 l/d for infants, children and adults (\geq 17 years) respectively, were used for calculations. The conversion factors for ²³⁸U, ²³²Th and ⁴⁰K as reported by ICRP^[20] and presented in Table 3 were used for all the age groups.

143 The total annual effective dose due to ingestion of water was computed using the following 144 formula ^[21, 22](ICRP, 1996, Ndontchueng et al., 2013).

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

 CF_{ing} (i) × Ai × I (2)

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

In addition to the estimated annual effective dose, the cancer and hereditary risk due to low dose without any threshold doses known as stochastic effect were estimated using the ICRP cancer risk methodology ^[23]. Radiation risks to members of the public results from exposure to low dose radiation are normally known as chronic risk of somatic or hereditary damage of human tissues, thus much emphasis is always placed on the reduction of these radiological

risks to natural radiation. The nominal lifetime risk coefficient of fatal cancer recommended in the 2007 recommendations of the members of the public is 5.5×10^{-2} Sv⁻¹. For hereditary effects, the detriment-adjusted nominal risk coefficient for the whole population as stated in^[23] for stochastic effects after exposure to low dose rates was estimated at 0.2×10^{-2} Sv⁻¹.

The risk to population was then estimated using the 2007 recommended risk coefficient of ICRP report and assumed 70 years lifetime of continuous exposure of the population to low level radiation. According to ICRP methodology;

161 Cancer Risk = Total annual Effective Dose $(Sv) \times Cancer risk factor (Sv⁻¹)$ (3)

162 Hereditary Effects = Total annual Effective Dose $(Sv) \times$ Hereditary effect factor (Sv^{-1}) (4)

163 The recommended reference levels of the effective dose for infants, children and adults 164 corresponding to one year consumption of drinking water are 0.26, 0.20 and 0.1 mSvy⁻¹ 165 respectively.

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167Table 1: Effective Dose Coefficients (Sv/Bq) for ingestion of Radionuclides for members of168the public to 70 years of age (ICRP, ^[20]; Publication 119)

S/N	Radioisotopes	Infant Children		Adult	
		\leq 1 year	10 years	> 17 years	
1	²³⁸ U	1.4 E-07	6.8 E-08	4.5 E-08	
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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170 **4.0 Results and Discussion**

The activity concentration of ²³⁸U, ²³²Th and ⁴⁰K determined in river water from Imo River and the associated annual effective dose to infant, children and adult population of the communities are presented in Table 2 while the estimated cancer risks and hereditary effects of adult member of the public are shown in Table 3.

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S/N Sample ID		Location	Activity Co	oncentration (Bq/l)	Total Annual Effective Dose (mSv/y)		
		²³⁸ U	²³² Th	⁴⁰ K	Infant	Children	Adult	
1	SW1	Otamiri-Imo River	BDL	BDL	BDL	0.00	0.00	0.00
2	SW 2	NNPC-Alscon	BDL	BDL	3.50±0.27	33.0	0.0167	0.0158
3	SW 3	Obigbo Bridge	BDL	0.85±0.089	20.33±1.50	193.0	0.186	0.235
4	SW 4	Mama Town	BDL	BDL	14.36±1.08	137.0	0.068	0.0650
5	SW 5	Old Imo River	BDL	3.77±0.37	22.11±1.71	212.0	0.504	0.0417
6	SW 6	Imo River Village	1.93±0.50	7.89±0.76	BDL	0.169	0.883	1.388
7	SW 7	Back of Kom-Kom	BDL	BDL	BDL	0.00	0.00	0.00
8	SW 8	Imo River Railway	3.08±084	BDL	BDL	0.079	0.077	0.0604
9	SW 9	NNPC Pipeline	4.36±1.07	BDL	3.84±0.30	13.63	0.034	0.0426
10	SW 10	Imo River	BDL	BDL	13.34±0.95	9.018	0.064	0.061
11	SW 11	Imo River Division 1	BDL	4.34±0.43	BDL	1.271	0.459	0.728
12	SW 12	Imo River Division 3	1.10±0.30	BDL	1.43±0.10	13.63	0.034	0.0426
13	SW 13	Imo River Division 5	BDL	4.12±0.41	BDL	1.206	0.436	0.692
14	SW 14	Imo River Division 7	BDL	BDL	7.99±0.60	76.0	0.038	0.0362
15	SW 15	Imo River Banks 1	1.27±0.34	BDL	2.17±0.17	20.60	0.042	0.0515
16	SW 16	Imo River Banks 2	BDL	4.02±0.40	BDL	1.177	0.425	0.675
17	SW 17	Imo River Banks 3	BDL	BDL	14.98±1.11	142.55	0.071	0.068
18	SW 18	Imo River Banks 4	0.39±0.09	BDL	9.81±0.73	93.40	0.056	0.057
19	SW 19	Imo River Banks 5	BDL	BDL	11.32±0.82	107.72	0.054	0.0512
20	ON 20	Mmiri-Nwayi	BDL	0.50±0.05	BDL	0.146	0.053	0.084
	SW 20	Division 14) Mean	2.02	3.59	10.43	58.64	0.19	0.24
	WHO, 20	08 Standard	10.0	1.0	10.0	0.26	0.20	0.10

Table 2: Activity Concentration of ²³⁸U, ²³²Th and ⁴⁰K in Water Samples and Total Annual Effective Dose for Different Age Categories

S/N	N Sample ID	Total Annual Effective Dose (mSv/y)			Fatality cancer risk to Adult per year	Lifetime fatality cancer risk	Severe hereditary Effects in Adult per/y	Estimated lifetime hereditary Effects
		Infant	Children	Adult	× 10 ⁻⁶	× 10 ⁻⁴	× 10 ⁻⁷	× 10 ⁻⁶
1	SW1	0.00	0.00	0.00	0.00	0.00	0.00	0.00
2	SW 2	33.0	0.017	0.016	0.87	0.61	0.32	2.22
3	SW 3	193.0	0.186	0.235	12.93	9.05	4.70	32.90
4	SW 4	137.0	0.068	0.065	3.57	2.50	1.30	9.09
5	SW 5	212.0	0.504	0.042	2.29	1.60	0.83	5.83
6	SW 6	0.169	0.883	1.388	76.35	53.44	27.76	194.30
7	SW 7	0.00	0.00	0.000	0.00	0.00	0.00	0.00
8	SW 8	0.079	0.077	0.060	3.32	2.32	1.21	8.45
9	SW 9	13.63	0.034	0.043	2.34	1.64	0.85	5.96
10	SW 10	9.018	0.064	0.061	3.35	2.34	1.22	8.522
11	SW 11	1.271	0.459	0.728	40.06	28.04	1.46	101.98
12	SW 12	13.63	0.034	0.043	2.34	1.64	0.85	596.4
13	SW 13	1.206	0.436	0.692	38.03	26.60	13.83	96.80
14	SW 14	76.0	0.038	0.036	1.99	1.39	0.72	5.06
15	SW 15	20.60	0.042	0.052	2.84	1.98	1.03	7.22
16	SW 16	1.177	0.425	0.675	37.12	25.98	1.35	9.45
17	SW 17	142.55	0.071	0.068	3.73	2.61	1.35	9.48
18	SW 18	93.40	0.056	0.057	3.15	2.20	1.14	8.01
19	SW 19	107.72	0.054	0.051	2.82	1.97	1.03	7.17
20	CW 20	0.146	0.053	0.084	4.62	3.23	1.68	1.18
	SW 20 Mean	58.64	0.19	0.24	13.43	9.40	3.39	61.67

182 Table 3: Estimated Cancer Risks and Hereditary Effects of Adult Member of the Public

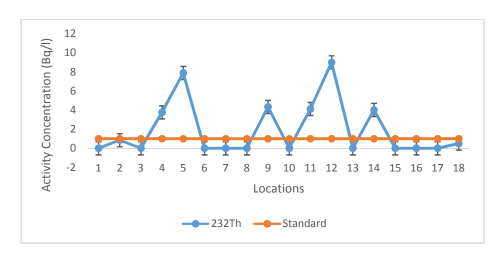
184 **4.1** Specific Activity Concentration of ²³⁸U, ²³²Th and ⁴⁰K in River waters

The specific activity concentration of ²³⁸U, ²³²Th and ⁴⁰K in the river watersamples are shown in 186 Table 2 and ranges from BDL to $4.36 \pm 1.07 \text{ BqL}^{-1}$ with an average value of $2.02 \pm 0.02 \text{BqL}^{-1}$, 187 BDL to 7.89 ± 0.76 Bql⁻¹ with an average value of 3.59 BqL⁻¹ and BDL to 22.11 ± 1.71 BqL⁻¹ with 188 an average value of 10.43 BqL⁻¹. The result clearly show that ²³⁸U are sparsely distributed along 189 the coastal shore. This could be due to high mobility of uranium-238 in river water. Uranium -190 191 238 were below detectable limit in most of the locations along the shore. This is in agreement with the fact that uranium in natural environment are variable in uranium content, depending 192 mainly on factors such as contact time with uranium bearing rocks, uranium content of the 193 contact rock, amounts of evaporation and availability of complexing ions. The ability of 194 uranium to undergo inorganic dissociation and re-precipitation is probably the most important 195 process in the natural environment to cause disequilibrium between the nuclides in the decay 196 chains. The large variation of uranium observed in this work could be due to PH values which 197 cause precipitation of uranium from the solution along the flow direction^[4]. 198

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The activity concentration of ⁴⁰K is highest at the old Imo River basin due to illegal oil and gas 200 bunkering activities that releases its wastes into the river. The activity concentration of ²³²Th in 201 river water was relatively higher than that of ²³⁸U because thorium is very insoluble^[24]. The 202 activity concentration of ²³²Th and ⁴⁰Kare slightly higher than the reference levels of 1.0 and 203 10.0 BqL⁻¹ while that for ²³⁸U is within the reference levels. The results obtained in this work was 204 compared with other works done in a similar environment within this country and other countries 205 of the world as presented in Table 4. Figures 2 and 3 shows the comparison of the activity 206 concentration of ²³⁸U, ²³²Th and ⁴⁰K in river water with standard value prescribed by ICRP, ^[20]. 207 It shows that ²³⁸U activity concentrations are lower than the standard value in all the locations 208 while about six locations, activity concentration of ²³²Th exceeded the standard value. The ICRP 209 ^[20] and WHO,^[25] regulations for drinking water quality does not include a listing for ⁴⁰K but 210 specifies that the maximum allowable concentration limit for beta and photon emitters should 211 correspond to a committed effective dose of 1.0 mSvy⁻¹ from annual intake at the rate of two 212 liters' of drinking water per day^[26]. 213





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Fig.2:Comparison of activity concentration of ²³²Th with ICRP, 2012 Standard

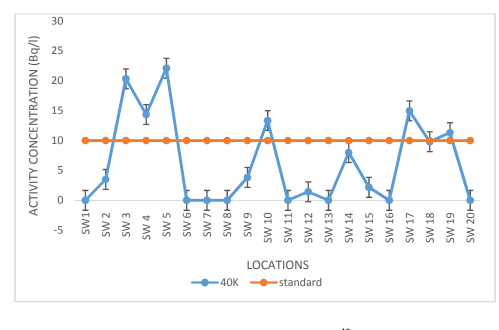
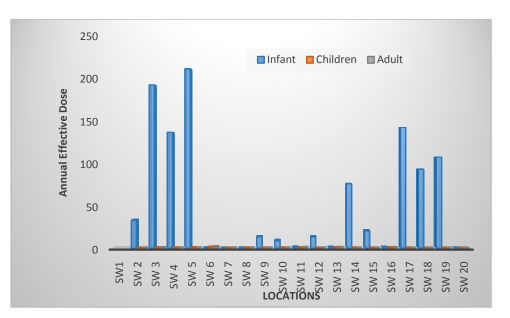


Fig.3: Comparison of activity concentration of ⁴⁰K with ICRP, 2012 Standard



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Fig. 4: Variations of total annual Effective dose for different age groups

The annual effective dose due to ingestion of the river water sampled was estimated for three 226 227 different age groups: Infants, children and adults. The calculated total annual effective dose for different age groups as shown in Table 2 range from 0.00 to212.0 mSvy⁻¹ for infants, 0.00 to 228 0.883 mSvy^{-1} for children and from 0.00 to 1.388 mSvy $^{-1}$ for adult with average values of 58.64, 229 0.19 and 0.24 mSvy⁻¹ respectively. It can be observed from Figure 5 that the radiation dose 230 received by infants is relatively higher than that received by children and adults. The WHO^[25] 231 and UNSCEAR^[19] reference levels of the effective dose for infants, children and adult due to 232 one year continuous ingestion of various drinking water are 0.26,0.20 and 0.10 mSvy⁻¹ 233 respectively. The effective doses obtained were higher than the reference values for infants, and 234 adults that consume river water but that for children are within the reference level of 0.2 mSvy 235 236 ¹.From the radiation protection point of view, life-long ingestion of these sampled river waters may cause significant radiological health problems. 237

In order to determine the radiation risk due to ingestion of ²³⁸U, ²³²Th and ⁴⁰K in river water 238 sampled, ICRP^[20] methodology was adopted in the study and the results are shown in Table 3. 239 The results of the cancer and non-cancer risk components were evaluated from the estimated 240 total annual effective dose of the various age groups. The result of the estimated fatal cancer risk 241 to adult per year in each of the stream water sampled ranged from 0.87×10^{-6} to 76.35×10^{-6} 242 with the associated lifetime fatality cancer risk of 0.61×10^{-4} to 53.44×10^{-4} . The estimated 243 hereditary effect to adult per year varied from 0.32×10^{-7} to 27.76×10^{-7} with its associated 244 lifetime hereditary effect in adult of 1.18×10^{-6} to 596.40×10^{-6} . This means that in terms of the 245 lifetime fatality cancer risk to adult approximately 53 out of 10,000 may suffer some form of 246 cancer fatality and for the lifetime hereditary effect approximately 596 out of 1000,000 may 247 suffer some hereditary effects. The negligible cancer fatality risk value recommended by 248

USEPA is in the range of 1.0×10^{-6} to 1.0×10^{-4} (ie 1 person out of 1 million to 10,000 persons suffering from some form of cancer fatality is considered trivial).

Comparing the estimated results of the lifetime fatality cancer risk in the present study with the acceptable risk factor, it can be seen that all estimated results of the lifetime fatality risk in adult member of the Nigerian population due to ingestion of radionuclide in the studied stream water are higher than the range of acceptable risk value recommended by USEPA.

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Table 4: Comparison of activity concentration of ²³⁸U, ²³²Th and ⁴⁰K in water samples of Imo River Rivers State Nigeria and other studies in different parts of the world.

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Samples	Country	²³⁸ U	²³² Th	⁴⁰ K	References
		(Bq l ⁻¹)	(Bq l ⁻¹)	(Bq I ⁻¹)	
Stream OD W (Nigeria)	Nigeria	0.59	1.8	27.7	[27]
Stream OW	Nigeria	4.62	4.06	42.57	[27]
Stream water Well OD	Nigeria Nigeria	9.044±3.11 3.16	2.28±0.57 2.38	100.37±23.47 235.64	[1] [27]
Mineral bottled water	Cameron	0.022	0.035	0.107	[22]
Portable water Borehole water	Nigeria Nigeria	0.000833 0.49	0.00005039 0.30	0.4191 7.40	[26] [28]
Stream water	Nigeria	2.02	3.59	10.43	This study

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261 **4.2 Statistical Analysis**

Basic statistics with statistical software package SPSS version 11.0 for windows was used to demonstrate the distribution and behavior of the measured radionuclide in stream watersand presented in Table 5. The statistical parameters determined includes the range (minimummaximum), arithmetic mean (AM), arithmetic standard deviation (SD), median, mode, skewness, kurtosis and the type of frequency distribution for the three radionuclides for all the water samples.

The frequency distribution curves of 238 U, 232 Th and 40 K are shown in Figure 5. From Table 5, all the radiological parameters have positive skewness which shows that 238 U, 232 Th and 40 K have asymmetric distribution and only 40 K has a negative kurtosis indicating relatively flat distribution. Pearson's correlation analysis was also carried out to ascertain if there are mutual relationship between the pairs of variables by calculating their linear correlation coefficient R^2 . It

is important to note that a positive correlation among variables indicates similar source andbehavior in the given environment.

Results of the Pearson correlation coefficient among all the three studied radionuclide and the associated radiological parameters are presented in Table 6. From Table 6, it can be observed that positive correlation exists among the three radionuclides and all the radiological parameters except 238 U having a negative correlation with AEDE_{children} and AEDE_{adult} indicating that uranium did not contribute to gamma emission on children and adult. Strong correlation were observed between 232 Th and 40 K while 238 U is weakly correlated with 232 Th and 40 K.

The strong positive correlation between ²³²Th and ⁴⁰K shows that their origin and behavior in the coastal environment are the same while weak positive relationship between ²³⁸U and the other two indicates that they may have the same origin but their behavior in the river environment differs. All the three radionuclides have strong positive correlation coefficient with the radiological parameters except for Uranium-238 that showed negative correlation with AEDEchildren and AEDEadult. This means that two of the radionuclide only contributed significantly to gamma-ray emission at the sampling points.

	U-238	Th-232	K-40	AEDEInfant	AEDEChildren	AEDEAdult
N Valid	18	18	18	18	18	18
missing	0	0	0	0	0	0
Mean	.6739	1.9161	6.9544	58.6442	.1945	.2442
Std. Error of Mean	.29716	.68676	1.77320	16.90533	.05726	.08878
Median	.1800 ^a	.3636 ^a	3.6700 ^a	18.2767 ^a	.0660 ^a	.0607 ^a
Mode	.00	.00	.00	13.63	.03	.04
Std. Deviation	1.26076	2.91367	7.52305	71.72324	.24293	.37666
Variance	1.590	8.489	56.596	5144.224	.059	.142
Skewness	2.067	1.431	.746	1.027	1.679	2.087
Std. Error of	500	500	500	500	500	500
Skewness	.536	.536	.536	.536	.536	.536
Kurtosis	3.795	1.065	741	261	2.420	4.132
Std. Error of Kurtosis	1.038	1.038	1.038	1.038	1.038	1.038
Range	4.36	9.00	22.11	211.92	.87	1.37
Minimum	.00	.00	.00	.08	.02	.02
Maximum	4.36	9.00	22.11	212.00	.88	1.39
Sum	12.13	34.49	125.18	1055.60	3.50	4.40

Table 5: Descriptive statistics of radiological parameters

a. Calculated from grouped data.

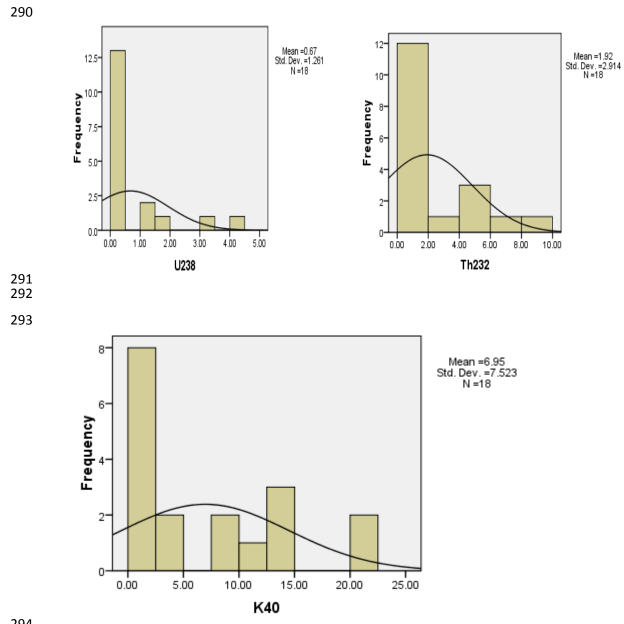




Fig. 5 : Frequency Distribution of ²³⁸U, ²³²Th and ⁴⁰K in stream water

303 Table 6: Pearson Correlations of measured parameters

304

	238U	232Th	40K	AEDEinfant	AEDEchild	AEDEAdult
238U	1					
232Th	-0.12834	1				
40K	-0.36877	-0.18064	1			
AEDEInfant	-0.37325	-0.10098	0.923147	1		
			-	-		
AEDEchildren	-0.03014	0.680037	0.152366	0.0826631	1	
				-		
AEDEAdult	0.027668	0.642057	-0.42099	0.3598336	0.89566772	1

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306 Conclusion

307 The activity concentration of ²³²Th and ⁴⁰K measured in river water collected from the meeting

point of Otamiri tributary and Imo River at the Abia/Rivers boundary to over seven kilometer

along the Imo River exceeded the reference level of 1.0 and 10.0 Bql⁻¹ while the activity

310 concentration

of²³⁸U measured are within the reference level of 10.0 Bql⁻¹. The meantotal annual effective dose

determined for infant, children and adult population that drink river water from the Imo River are

58.64, 0.19 and 0.24 mSvy⁻¹ respectively. AEDE estimated for infant are 94% higher than the reference

Level of 0.26 mSvy^{-1} and also higher than that for children and adult.

317

The estimated fatal cancer risk to adult per year and the lifetime hereditary effect shows that 53 out of

10,000 population may suffer some form of cancer fatality and approximately 596 out of
 1000,000 might

Suffer some hereditary effects. Statistically all the radionuclide showed positive skewness and kurtosis except ²³⁸U. Pearson correlation of the radionuclides and all the radiological parameters showed positive correlation between ²³²Th and ⁴⁰K which indicate same origin and behavior in the coastal environment. 238U showed negative correlation with the radiological parameters which shows that uranium -238 did not contribute to gamma emission and probably had a different origin.

328

The result of this study showed that the activity of oil bunkering along the creeks, river shore has impacted negatively on the river water which in turn might lead to health challenges of the population that depend on such river for their daily living.

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