

Sequential Extraction and Geochemistry of Heavy Metals in Ayetoro Coastal Sediments, Southwestern Nigeria

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Research Article

Volume 5 Issue 4 Received Date: September 28, 2021 Published Date: November 03, 2021 DOI: 10.23880/jenr-16000261

Abstract

Previous geochemical investigation of Ayetoro area discovered that its coastal sediments are enriched with sulphide mineralization. However, in order to determine the geochemical phases of the heavy metals in the coastal sediments, random sampling method was utilized across 10 locations, at a depth of 40cm using Van grab sampler at a sampling density of 200m interval. Atomic Absorption Spectrophotometer (AAS) Buck Scientific Model 205A was used to analyze nine (9) heavy metal concentrations namely Ni, Zn, Co, Mn, Fe, Pb, Cr, Cd and Cu in the coastal sediments, followed by sequential extraction of the metals, using five fractional phases. The results revealed that the geochemical concentration of the heavy metals as follows: Ni (5.89ppm - 16.82ppm), Zn (2.59ppm - 115.65ppm.), Co (1.22ppm - 22.77ppm), Mn (30.95ppm - 186.49ppm), Fe (6.632ppm -1925.96ppm), Pb (5.17ppm - 55.96ppm), Cr (0.26ppm - 28.06ppm), Cd (0.13ppm -22.23ppm), and Cu (2.26ppm - 41.94ppm) and showed the concentration order as Residual>Reducible>Organic>Exchangeable>Carbonate. Most of the heavy metals in carbonate and exchangeable phase have low concentration except for Cd. This implied that Cd is of low mobility and bioavailability which is very dangerous as its intake by man leads to kidney diseases and causes bones to become weaker. Also, Mobility factor of Cd stood out because of its high concentration in the exchangeable phase compared to other four non-residual phases. The mobility and bioavailability of the heavy metals are in this order: Cd>Co>Ni>Pb>Cr>Mn>Cu>Zn>Fe respectively. The analysis of variance (ANOVA) revealed that the heavy metals are significantly different in all the phases based on their accumulation index in the sediments while majority of the heavy metals lacked the ability to remobilize but can be released into the environment under reducing and oxidizing conditions.

Keywords: Ayetoro; Coastal Sediments; Heavy Metals; Sequential Extraction; Mobility Factor

Introduction

Sediments are considered to be mixture of several components of mineral species and represent important sinks for various pollutants in aquatic systems including heavy metals. It also plays a prominent role in the assessment of heavy metal contamination [1,2]. Total amount of heavy metal concentration in sediment is useful to detect net change.

Never the less, it does not give any sign about the chemical form of each metal in sediment [3]. Ayeku [4] studied the pollution status in the bottom sediment in Awoye, Abereke and Ayetoro Area of Ondo state. He concluded that these metals derived from the upstream rivers from the top soil are products of mechanically weathered rock materials and anthropogenic activities while Ololade [5] agreed that metal speciation should be done to determine the bioavailability of metal in sediment after his own seasonal metal distribution research in Ondo coastal sediment. Heavy metal concentration in the ocean ecosystem is determined by three conditions namely; water, sediments and living organism. Usually, heavy metal exists in lowest concentration in water and reaches considerable concentration in sediments followed by bioaccumulation in living organism. According to Li [6], heavy metals are released from sediments into water bodies and consequently, to living organisms depending on the speciation of metals and other factors such as sediment pH and organic matter [7]. Coastal and marine ecosystems worldwide are continuously bedeviled with pollution, such as eutrophication, acidification, toxic substances, heavy metals, and the likes. Decline in ecosystem productivity, loss of aesthetic beauty of the ocean, impacts on sensitive habitats, impairment on quality of seawater, hazards to human health are some of the consequences of heavy metals accumulation in sediments which makes continuous long term monitoring of heavy metal concentration of coastal sediment extremely important. Therefore, this research attempts to determine the geochemical phases of the heavy metals in the coastal sediments of Ayetoro area using five metal fractional methods, in order to assess the levels of their bioavailability, bioaccumulation and the danger it portends to man and ecology.

Description and Geology of the Study Area

The study area (Ayetoro) is one of the prominent, waterroute nodal communities in the coastal region of Ondo State, Nigeria. Ayetoro was established in 1947 and is inhabited by the Ilaje people, a linguistic subgroup of the Yorubas. The town lies east along the coast from Nigeria's largest city, Lagos. Due to oil exploration activities, the community has lost a considerable portion of coastline to the Atlantic Ocean. Ayetoro which is one of the main settlements in Ilaje local government area lies approximately within latitude 6° 12' 33.786" N and longitude 4° 40' 17.051" E respectively. It is bounded in the north-east by Erunna village, on the southeast by Alagbon Village and North-west by Idi Ogba and south-west by the Atlantic Ocean. The people are extractions of the Ilaje sub-ethnic group of the Yoruba's in south western Nigeria. Ayetoro in its hey-days once had the highest per capital income in the whole of Africa and attracted visitors, tourists and researchers from all over the world. The occupational activities in this area include fishing, canoe making, and lumbering. Others are net making, mat making, launch building, farming and trading. Residents say the ocean incursion has impacted on their livelihoods, particularly since the community's entire mangrove vegetation was destroyed. The Atlantic's surges have also destroyed Ayetoro's marine life; thereby crippling people's fishing businesses, which is the mainstay of the local economy.

Journal of Ecology and Natural Resources

There are two distinct geological regions in Ondo State. First, is the region of sedimentary rocks in the south, and secondly, the region of Pre Cambrian-Basement Complex rocks in the north. The sedimentary rocks are mainly of the post Cretaceous sediments and the Cretaceous Abeokuta Formation. The basement complex is made up of mainly the medium-grained gneisses. These rocks are strongly foliated most of the times, occurring as outcrops. The surface of these outcrops is rigorously distorted with alternating bands of dark and light-colored minerals. These bands of lightcolored minerals are essentially feldspar and quartz, while the dark colored bands contain abundant biotite, a small proportion of the state, especially to the northeast, overlies the coarse-grained granites and gneisses, which are poor in dark ferromagnesian minerals. Troughs and undulating low land surfaces cover Ilaje Local Government Area with silt, mud and superficial sedimentary deposits [8]. There are sand formation at the western part of the local government, extending from the Lekki peninsula in Lagos State to Araromi Sea-side and Zion pepe, Mahin and Ugbonla which are in the eastern part of the local government area. This could be the reason why there is sand deposition on the western side like Igbokoda and clay formation in areas like Ayetoro. Crude oil, which is a major source of income in Nigeria, is found in Ilaje Local Government. There are oil wells and fields spreading all over the local government area both onshore and offshore. Oil companies such as Shell, Chevron are believed to be presently harnessing the crude oil found in the area. Apart from petroleum, there are other natural resources and raw materials present in Ilaje land such as glass sand, salt, tar sand, quartz and clay deposits (Figures 1 & 2).



Figure 1: Sampling points of Coastal Sediments.



Figure 2: Geological map of Ondo State (Geological survey of Nigeria).

Materials and Methods

Method of Sampling and Sample collection

The samples were collected using random sampling method from Ayetoro to Eyin More. After removing the overburden, the samples were collected randomly at each station at a depth of 40cm and sample density of 200m intervals. Global Positioning Systems (GPS) G GARMIN eTrex 10 was employed to get the accurate geographical coordinates of each sampling points. A total of ten samples were collected and bagged separately inside the polythene bags and properly labelled to avoid mix up. The samples were air dried and stored in a cool and dry place to avoid contaminations. Observation of geological settings, physical structures and lithology together with human activities were noted and the samples were eventually transmitted to Temsol Consults Laboratory Ltd Shasha, Ibadan for sequential extraction analysis (Table 1).

The field data for the coastal sediment samples is presented in Table 2. Ten coastal sediments were randomly selected from different localities within the study area. Their geographical coordinates as well as their colour and nature of the sediments were taken into consideration. The colour ranges from grey, dark grey, yellowish brown, brown to brownish grey, while the composition of the sediments are clayey and silty in nature.

Station ID	Locations	Latitude	Longitude	Colour	Sediment composition
S1	Ayetoro	6°6′25.76″N	4°46′41.90″E	Dark Grey	Clay
S2	Ero	6°6′48.50″N	4°46′46.50″E	Brownish grey	Clay
S3	Eru-Ona	6°7′3.20″N	4°46′24.30″E	Brown	Silt
S4	Okun Harama	6°7′22.50″N	4°45′52.80″E	Yellowish Brown	Silt
S5	Olootu	6°7′39.60″N	4°45′32.70″E	Grey	Clay
\$6	Niye	6°8′0.10″N	4°44′58.20″E	Dark Grey	Clay
S7	Yaye	6°8′1.30″N	4°44′48.90″E	Dark Grey	Clay
S8	Lepe	6°8′23.30″N	4°44′29.00″E	Grey	Clay
S9	Dogun	6°8′40.78″N	4°44′12.98″E	Grey	Clay
S10	Eyin-more	6°8′55.30″N	4°43′53.00″E	Grey	Clay

Table 1: Field Data and Description of Coastal Sediments.

Laboratory Analysis

Sequential Extraction Procedure

The speciation of toxic metals is important for measuring their bioavailability in the environment, and for evaluating their potential risks to living organisms [9]. In the study, the investigated metals speciation revealed differences in the concentrations that were recorded at each step of the extraction. Using a modified version of Tessier [10] method, the toxic metals were separated into five operationally defined fractions viz. exchangeable (F_1), carbonate-bound (F_2), Fe-Mn oxide-bound (F_4), organic-bound (F_4), and residual (F_5) fractions (F5) [11,12]. The sequential extraction procedures are as follows:

(i) F_1 : 1 g of dried and powdered sediment was extracted at room temperature with 1 M MgCl₂ at pH 7.0 for 1hr with continuous agitation. Then, the mixture was centrifuged. The supernatant obtained on standing was filtered to represent F_1 .

(ii) F_2 : The residue from (i) was leached at 30 °C with 1 M sodium acetate (NaOAc) adjusted to pH 5.0 with acetic acid (HOAc). The mixture was continuously agitated throughout the extraction using a centrifuge. The extracts were decanted to represent F_2 .

(iii) F_3 : 20 ml of 0.04 M hydroxylamine chloride (NH₂OH. HCl) in 25% (v/v) acetic was added to the residue from (ii). The mixture was agitated 96°C for 6 h. Then, the extract was decanted to represent F_3 .

(iv) F_4 : 3ml of 0.02 M HNO₃ and 5ml of 30% H_2O_2 were added to the residue from (iii). The mixture was agitated at 85°C for 5 hours then, 5 ml of 3.2 M NH₄OAc was added and further centrifuging for 30 min before filtering. The supernatant represents F_4 .

(v) F_{s} : The final fraction was obtained by digesting the residual from (iv) with 5ml of 25% of HC1 and 5ml HNO₃. for 6hrs at 120°C. The mixture was centrifuged, and the supernatant was obtained as F_{s} .

All reagents used were analytical grade. The supernatant from each extraction was quantitatively transferred into a 25 ml volumetric flask and made up to mark with 1 M HNO₃ before quantifying the trace metals using Atomic Absorption Spectrophotometer (AAS, Buck Scientific Model 205A). All analyses were carried out in triplicates, and reagent blanks were used for quality control.

Metal Analysis

Trace metal concentrations were determined by atomic absorption spectrophotometry Model 210 VGP of the Buck Scientific AAS series with air-acetylene gas mixture as oxidant involving direct aspiration of the aqueous solution into an air-acetylene flame. The following techniques were used for the first four fractions. For the trace metals Cd, Co, Cu, Cr Ni, Pb, and Zn, For the metals present in high concentrations (Fe and Mn) the supernatant solution was diluted (20 to 50 X) with deionized water and the concentrations were obtained directly from appropriate calibration curves prepared with the components of the extraction solution diluted by the same factor. For total or residual trace metal analysis, the solid was digested with a 5:1 mixture of hydrofluoric and perchloric acids. For 1g (dry weight) sample, the sediment was first digested in a platinum crucible with a solution of concentrated $HC10_4$ (2 ml) and HF (10 ml) to near dryness; subsequently, a second addition of $HC10_4$ (1 ml) and HF (10 ml) was made and again the mixture was evaporated to near dryness. Finally, $HC10_{4}$ (1 ml) alone was added and the sample was evaporated until the appearance of white fumes. The residue was dissolved in 12ml NHC1 and diluted to 25 ml. The resulting solution was then analyzed by flame atomic absorption spectrophotometry for trace metals using the standard addition techniques.

Mobility Factor=
$$\frac{F1 + F2}{F1 + F2 + F3 + F4 + F5} * 100$$

Mobility Factor is used to evaluate the potential mobility of and bioavailability of heavy metal. High percentage represents high potential and vice versa

Results and Discussion

Geochemical Phases

Exchangeable Phase

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Table 5 showed the distribution of the maximum, average and minimum concentration of nine heavy metals in the exchangeable phase. The average concentration of the heavy metals is: Mn>Pb>Fe>Ni>Cr>Co>Zn>Cu>Cd. Also, Figures 3 and 4 showed that Mn is predominantly abundant across all sampling points. The following are the range of concentration of heavy metals in this phase: Ni (6.24 - 9.04), Zn (0.07 - 10.64), Co (1.9 - 5.42), Mn (20.16 - 38.72), Fe (2.64 - 18.32), Pb (9.04 - 14.32), Cr (2.72 - 4.85), Cd (1.29 - 1.74), Cu (0.96 - 9.76). Apart from sample point S1, Zn is notably low in all other points. Conc. of Fe in S2, S5 and S10 are notably different from others perhaps because of inflow of some extraneous materials from the land to this particular area.

Toxic Metals	Ni	Zn	Со	Mn	Fe	Pb	Cr	Cd	Cu
Max conc.(ppm)	9.04	10.64	5.42	38.72	18.32	14.32	4.85	1.74	9.76
Average conc.(ppm)	7.52	2.59	3.10	30.95	8.29	12.51	3.38	1.57	2.26
Min conc.(ppm)	6.24	0.07	1.9	20.16	2.64	9.04	2.72	1.29	0.96

Table 2: Maximum, Average and Minimum Concentration of heavy metals in the Exchangeable phase.



Figure 3: Distribution of Heavy Metal in the Exchangeable Phase with Respect to Sampling Points.



Carbonate Phase

Table 3 presents the distribution of the maximum, average and minimum concentration of nine heavy metals in the carbonate phase. Average concentration of the heavy metals are in this order: Mn>Fe > Pb > Ni>Zn > Cu>Cr>Co > Cd. The average concentration of Fe is a little higher than Pb in the carbonate phase unlike the exchangeable phase. Likewise, the average concentration of Zn is higher than Cr

in the carbonate phase. Except for Zn and Mn the average concentration of all the other heavy metals reduce drastically in the carbonate phase. The following are the ranges of concentration of heavy metals in this phase. Ni (3.25 - 6.81), Zn (1.33 - 9.71), Co (0.37 - 1.72), Mn (20.72 - 93.36), Fe (2.08 - 11.68), Pb (4.56 - 7.62), Cr (0.12 - 2.3), Cd (0.54 - 0.94), Cu (1.44 - 6.72) (Figures 5 and 6 respectively).

Heavy Metals	Ni	Zn	Со	Mn	Fe	Pb	Cr	Cd	Cu
Max conc.(ppm)	6.81	9.71	1.72	93.36	11.68	7.62	2.3	0.94	6.72
Ave conc.(ppm)	5.894	3.37	1.222	66.55	6.632	6.278	1.162	0.743	2.854
Min conc.(ppm)	3.25	1.33	0.37	20.72	2.08	4.56	0.12	0.54	1.44

Table 3: Maximum, average and minimum concentration of heavy metals in the carbonate phase.





Figure 6: Bar chart showing the Carbonate Phase with respect to Maximum, average and Minimum of Heavy Metals Concentration.

Reducible Phase

Table 4 presents the distribution of the maximum, average and minimum concentration of nine heavy metals in the Reducible phase. Average concentration of the heavy metals is in this order: Fe> Mn> Zn> Ni> Cu>Pb> Co> Cr> Cd. The concentrations of Fe and Mn have tremendously increased to more than 50% for Mn and 100% for Fe

respectively (Figures 7 and 8). Except for Pb and Cd, the concentration of all other metals increases notably since the previous phase. The following are the ranges of concentration of heavy metal in this phase: Ni (3.66 - 13.6), Zn (10.12 - 90.64), Co (1.87 - 9.12), Mn (60.76 - 463.4), Fe (409.89 - 1222.14), Pb (1.34 - 12.8), Cr (0.83 - 6.48), Cd (0.02 - 0.22), Cu (2.23 - 12.27).

Heavy Metals	Ni	Zn	Со	Mn	Fe	Pb	Cr	Cd	Cu
Max conc.(ppm)	13.6	90.64	9.12	463.4	1222.14	12.8	6.48	0.22	12.27
Ave conc.(ppm)	8.98	35.959	5.093	178.867	735.334	5.165	4.233	0.128	5.99
Min conc.(ppm)	3.66	10.12	1.87	60.76	409.89	1.34	0.83	0.02	2.23

Table 4: Maximum, Average and Minimum Concentration of heavy metals in the Reducible phase.



Figure 7: Distribution of Heavy Metals in Reducible Phase with Respect to Sampling Points.



Organic Phase

The distribution of the maximum, average and minimum concentration of nine heavy metals in organic phase is presented (Table 5). Average concentration of the heavy

metals is in this order: Fe> Cu> Mn> Zn> Ni> Pb> Cr>Cd >Co. The average concentration of Cu over the sampling points in the organic phase increased drastically (Figures 9 and 10).

Heavy Metals	Ni	Zn	Со	Mn	Fe	Pb	Cr	Cd	Cu
Max conc.(ppm)	16.21	37.34	5.62	47.04	628.59	13.6	9.52	0.62	55.21
Ave conc.(ppm)	9.66	27.75	2.809	31.058	424.267	6.776	6.408	0.26	41.939
Min conc.(ppm)	2.69	15.8	0.74	19.46	245.18	1.29	2.71	0.04	21.26

Table 5: Maximum, average and minimum Concentration of heavy metals in the Organic Phase.



Figure 9: Concentration of Heavy Metals in Organic Phase with Respect to Sampling Points.



Residual Phase

Table 6 presents the distribution of the maximum, average and minimum concentration of nine heavy metals in the residual phase. Average concentration of the heavy metals is in this order: Fe> Mn> Zn> Pb> Cu> Cr> Co >Cd Ni. Fe has the highest concentration in this phase (Figures 11and 12).

Toxic Metals	Ni	Zn	Со	Mn	Fe	Pb	Cr	Cd	Cu
Max conc.(ppm)	20.15	127.51	29.65	230.22	4412.61	79.2	40.05	25.1	36.64
Ave conc.(ppm)	16.819	115.647	22.7795	186.498	1925.966	55.963	28.055	22.228	29.434
Min conc.(ppm)	12.66	102.22	2.715	155.65	1000.95	30.3	17.14	17.85	21.16

Table 6: Maximum, Average and Minimum Concentration of Heavy metals in the Residual Phase.





Analysis of Heavy Metals Across all Phases

Nickel (Ni)

The average concentration of Ni over the sampling points and across all phases is between 5.89ppm to 16.82ppm (Table 7). The concentration increases in this order; Resi dual>Organic>Reducible>Exchangeable>Carbonate. Ni is predominantly abundant in the residual and organic phases, except for S7 and S8 (Figures 13 and 14) which have low concentration in the organic phase. This means that it is of mineral origin and do not pose any biological risk. Ni plays an important role in the development of some plants. However, it is not an essential nutrient for human. However, the kidney removes most of the nickel absorbed by humans.

Phases	Max	Ave	Min
Exchangeable	9.04	7.52	6.24
Carbonate	6.81	5.89	3.25
Reducible	13.6	8.98	3.66
Organic	16.21	9.66	2.69
Residual	20.15	16.82	12.66

Table 7: Maximum, Average and Minimum Concentration. Of Ni Over all Phases.



Figure 13: Concentration of (Ni) in Respect to Sampling Point and Phases.



Zinc (Zn)

The average concentration of Zn over the sampling points and across all phases is between 2.59ppm to 115.65ppm (Table 8). The concentration increases in this order; Residual> Reducible>Organic>Carbonate>Exchange able. Zinc is mostly abundant in the residual phase and S1 and S4 have the highest concentrations while it is of high concentration in other phases. For instance, sampling points such as S3, S4, S6, S7, S8 and S10 showed that it is low in the exchangeable phase. Zinc is an important element in the environment. Its deficiency is connected to many diseases such as delayed sexual maturation, infection susceptibility and diarrhea. Zinc is important for over 300 enzymes and 1000 transcription factors, and is stored and transferred in metallothioneins. Excess zinc is however toxic to plants. Fishes cannot tolerate the amount of Zn, as do plants [13]. Excess zinc inhibits calcium uptake in fish, which can be deadly (Figures 15 and 16). In this study, it is discovered that the concentration of Zn exceeded the normal concentration expected to be in sediment hence this necessitates prompt measures to curtail its menace.





Cobalt (Co)

The average concentration of Co over the sampling points and across all phases is between 1.22ppm to 22.77ppm respectively (Table 8). The concentration increases in this order; Residual> Reducible>Exchangeable>Carbonate>O rganic. Co is mostly abundant in the Residual phase and relatively low in other phases with S1 having the highest concentration (Figures 17 and 18). Co is always involved in photosynthesis and nitrogen fixation detected in most ocean basins and a limiting micronutrient for phytoplanktons and cyanobacteria. P value of 5.71E-17 in analysis of variance indicated that there is a significant difference from one phase to another.

Phases	Max	Ave	Min
Exchangeable	5.42	3.09	1.9
Carbonate	1.72	1.22	0.37
Reducible	9.12	5.09	1.87
Organic	5.62	2.80	0.74
Residual	29.65	22.77	2.72

Table 8: Maximum, Average and Minimum concentration of Co over all the Phases





Manganese (Mn)

The average concentration of Mn over the sampling points and across all phases is between 30.95ppm to 186.49ppm (Table 9). The concentration increases in this order; Residual> Reducible> Carbonate Organic>Exchangeable. Except for sampling point S8, which has an extremely high concentration in the reducible phase, Mn is evenly abundant in the residual phase followed by the reducible phase. In the aquatic bodies, many enzymatic systems need Mn to function, but in high levels, Mn can become toxic. At level of 500ppm Mn is dangerous to life; therefore Mn in the study area does not pose any threat to lives in the location. 3.2E-10 P value of annova indicates that there is significant difference between the five phases (Figures 19 and 20).

Phases	Max	Ave	Min
Exchangeable	38.72	30.95	20.16
Carbonate	93.36	66.55	20.72
Reducible	463.4	178.86	60.76
Organic	47.04	31.06	19.46
Residual	230.22	186.49	155.65

Table 9: Maximum, Average and Minimum concentration of Mn over all the Phases.





Iron (Fe)

According to Table 10, the average concentration of Fe over the sampling points and across all phases is between 6.632ppm to 1925.96ppm. The concentration increases in this order; Residual> Reducible > Organic>Exchangeable>Carbonate (Figures 21 and 22). Iron is extremely low in the carbonate and exchangeable phase but very high in the residual and also present in the reducible phase. Iron accumulation poses a problem for aerobic organisms because ferric iron is poorly soluble near neutral ph. Iron plays an essential role in marine systems and can act as a limiting nutrient for planktonic activity. Its excess may lead to a decrease in growth rates in phytoplantonic organisms. 5.6E-10 P value of ANOVA indicates that there is significant difference between the five phases.

Phases	Max	Ave	Min
Exchangeable	18.32	8.29	2.64
Carbonate	11.68	6.632	2.08
Reducible	1222.14	735.33	409.89
Organic	628.59	424.27	245.18
Residual	4412.6	1925.96	1000.95

Table 10: Maximum, Average, minimum concentration of Fe over the mineralogical Phases.





Lead (Pb)

Table 11 presents the average concentration of Pb over the sampling points and across all phases to be between 5.17ppm to 55.96ppm. The concentration increases in this order; Residual> Exchangeable> Organic >Carbonate>Reducible (Figures 23 and 24). Pb is more abundant in the Residual phase and has no confirmed safe level of exposure. Even at level considered to pose little or no risk, Pb may cause adverse mental health outcomes. Pb is a highly poisonous metal both inhaled and swallowed, causing great damage to almost every organs and systems in the human body. High level of Pb causes lead poisoning while low levels cause increase susceptibility to terminal diseases. High concentration of lead has the capacity to inhibit photosynthesis. Bioaccumulation of Pb poses a serious hazard to fishes and other sea mammals. From the

results obtained in this study, the concentration is within the residual phase which means that Pb is not bioavailable and will therefore tend not to remobilize. 9.5E-21 P value of ANOVA indicated that there is significant difference between five phases.

Phases	Max	Ave	Min
Exchangeable	14.32	12.51	9.04
Carbonate	7.62	6.28	4.56
Reducible	12.8	5.17	1.34
Organic	13.6	6.78	6.41
Residual	79.2	55.96	30.3







Chromium (Cr)

The result of the average concentration of Cr over the sampling points and across all phases is between 0.26ppm to 28.06ppm (Table 12). The concentration increases in this order; Residual >Reducible>Exchangeable>Carbonate>Orga nic. Cr is more abundant in the residual phase and extremely low in the organic phase (Figures 25 and 26). Chromium is

highly toxic to fish because it is easily absorbed across gills, readily enters blood circulation, and crosses cell membrane and bio concentrate up the food chain. Acute and chronic exposure to chromium affects fish behavior, physiology, reproduction and survival. Toxicity ranges between 50 and 150 ppm [14]. 1.07E-20 P value of ANOVA indicates that there is significant difference between five phases.





Cadmium (Cd)

Table 12 revealed that the average concentration of Cd over the sampling points and across all phases is between 0.13ppm to 22.23ppm. The concentration increases in this order; Residual> Exchangeable> Carbonate>Organic > Reducible. Cd is abundant in the residual and present in an extremely low concentration in other phases (Figures 27and 28) respectively. Cd is considered an environmental pollutant that causes health hazards to living organisms. Anthropogenic sources include fossils fuel combustion and fertilizers. Cd exposure is associated with a large number of illnesses including kidney disease, early atherosclerosis, hypertension and cardiovascular diseases. 5.11E-40 P value of annova indicates that there is significant difference between five phases. Cd in this location must be monitored because it has good percentage of its concentration in the exchangeable phase [15].

Phases	Max	Ave	Min
Exchangeable	1.74	1.57	1.29
Carbonate	0.94	0.74	0.54
Reducible	0.22	0.13	0.02
Organic	0.62	0.26	0.04
Residual	25.1	22.23	17.85

Table 12: Maximum, Average and Minimum Concentrationof Cd over all the Phases.





Cupper (Cu)

The average concentration of Cu over the sampling points and across all phases is between 2.26ppm to 41.94ppm (Table 13). The concentration increases in this order; Organic> Residual > Carbonate> Reducible. Cu is more abundant in the organic phase (Figures 29 and 30). Due to its role in facilitating iron uptake, Cu deficiency can produce anemia-like systems, neutropenia, bone abnormalities and impaired growth. 2.4E-27 P value of annova indicates that there is significant difference between the five phases.

Phases	Max	Ave	Min
Exchangeable	9.76	2.26	0.96
Carbonate	6.72	2.85	1.44
Reducible	12.27	5.99	2.33
Organic	55.21	41.94	21.26
Residual	36.64	29.43	21.16

Table 13: Maximum, Average and Minimum concentrationof Cu over all the Phases.





Mobility Factor across the Metal fractional phases

is presented in Tables 14-22. It can be deduced that Cd has the highest potential to remobilize across all sampling points because of its high mobility factor (Figure 31).

The mobility factors of the various heavy metals analyzed

Sample Points	F1	F2	F3	F4	F5	Mobility Factor(%)
S1	8.72	3.25	3.66	15.6	20.01	3.65
S2	6.56	6.32	6.57	9.39	17.42	7.04
S3	7.84	6.24	6.22	14.41	13.05	6.76
S4	7.28	4.88	10.8	7.67	19	7.21
S5	9.04	6.24	8.81	10.64	17.2	7.38
\$6	7.68	6.32	11.86	16.21	17.05	4.63
S7	6.24	6.32	9.43	2.85	20.15	15.82
S8	6.72	6.81	13.6	2.69	12.66	22.11
S9	8.64	6	12.44	7.66	16.35	9.61
S10	6.48	6.56	6.41	9.48	15.3	7.93

 Table 14: Mobility Factor of Ni.

Sampling Points	F1	F2	F3	F4	F5	Mobility Factor (%)
S1	10.64	2.07	44.26	28.51	127.51	0.34
S2	4.96	9.71	64.02	33.42	115.05	0.37
S3	0.59	5.71	90.64	27.22	107.57	0.21
S4	0.48	1.66	19.34	23.72	127.25	0.07
S5	6.96	2.46	10.12	33.51	102.22	0.27
\$6	0.16	2.46	29.74	27.62	115.71	0.08
S7	0.41	1.74	25.86	15.8	118.55	0.11
S8	0.07	1.33	33.34	28.58	108.58	0.04
S9	1.07	4.85	24.64	37.34	115.09	0.14
S10	0.53	1.71	17.63	21.78	118.94	0.09

Table 15: Mobility Factor of Zn.

Sampling Points	F1	F2	F3	F4	F5	Mobility Factor (%)
S1	2.88	1.46	3.34	1.82	29.35	7.10
S2	2.84	1.68	1.87	2.15	27.25	6.95
S3	2.09	1.26	3.38	2.42	29.65	4.26
S4	5.42	1.72	6.92	2.96	25.25	8.04
S5	5.31	1.34	3.24	4.18	26.75	5.46
S6	1.9	1.26	8.76	4.04	21.08	3.25
S7	2.75	1.29	4.12	0.98	2.715	37.33
S8	2.3	0.95	9.12	3.18	23.18	3.77
S9	2.34	0.89	6.28	5.62	21.41	2.48
S10	3.13	0.37	3.9	0.74	21.16	15.17

Table 16: Mobility Factor of Co.

Sampling Points	F1	F2	F3	F4	F5	Mobility Factor (%)
S1	28.08	76.88	180.21	19.52	205.18	2.44
S2	38.72	20.72	60.76	30.01	175.46	1.10
S3	38.08	76.08	107.55	19.46	175.23	3.14
S4	26.88	53.21	143.81	31.74	155.82	1.54
S5	30.88	68.8	100.26	29.73	210.02	1.54
S6	20.72	68.82	190.62	47.04	205.11	0.90
S7	20.16	76.48	158.86	25.67	155.65	2.27
S8	38.64	93.36	463.4	31.4	171.45	2.20
S9	29.68	63.23	236.6	42.81	230.22	0.91
S10	37.68	67.92	146.6	33.2	180.84	1.68

Table 17: Mobility Factor of Mn.

Sampling Points	F1	F2	F3	F4	F5	Mobility Factor (%)
S1	6.38	11.68	552.04	381.62	1196.8	0.003
S2	12.16	11.22	694.3	481.08	1185.45	0.004
S3	4.16	8.88	409.89	278.84	1484.5	0.003
S4	3.44	5.76	816.28	412.18	1000.95	0.002
S5	18.32	6.08	470.9	628.59	2372.65	0.001
S6	7.44	2.08	1023.8	495.08	1555.3	0.001
S7	2.64	4.46	746.5	245.18	1145.15	0.002
S8	12.24	3.52	1222.14	441.42	4412.61	0.0008
S9	3.24	9.68	718.87	600	3591.2	0.0005
S10	12.88	2.96	698.62	278.68	1315.05	0.004

Table 18: Mobility Factor of Fe.

Sampling Points	F1	F2	F3	F4	F5	Mobility Factor(%)
S1	12.88	6.56	5.49	5.22	79.2	4.43
S2	14.01	6.12	8.44	13.6	71.15	2.02
S3	14.16	4.56	6.25	7.02	61.6	4.09
S4	12.96	5.76	1.56	2.68	66.4	9.44
S5	14.32	6.32	1.88	10.42	30.3	6.10
S6	12.8	6.08	7.21	6.46	41.1	6.47
S7	11.76	7.36	1.34	4.64	59.85	6.41
S8	9.04	6.56	3.81	4.82	41.18	7.15
S9	11.84	5.84	2.87	11.61	47.85	3.06
S10	11.36	7.62	12.8	1.29	61	17.18

Table 19: Mobility Factor of Pb.

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Sampling Points	F1	F2	F3	F4	F5	Mobility Factor(%)
S1	4.85	1.34	4.76	5.84	22.05	4.43
S2	4.12	2.3	2.14	9.36	28.54	2.32
S3	3.08	0.96	6.24	5.12	38.35	1.95
S4	2.96	2.24	5.26	4.5	26.41	4.02
S5	3.66	1.54	6.48	8.63	32.56	1.77
S6	3.11	1.17	4.44	6.58	29.95	2.07
S7	3.22	0.51	5.68	2.71	40.05	3.16
S8	3.04	1.12	3.94	7.14	17.14	3.18
S9	2.72	0.32	2.56	9.52	18.05	1.71
S10	3.06	0.12	0.83	4.68	27.45	2.40

Table 20: Mobility factor of Cr.

Sampling Points	F1	F2	F3	F4	F5	Mobility Factor (%)
S1	1.55	0.75	0.12	0.11	17.85	52.46
S2	1.74	0.66	0.15	0.25	23.35	28.61
S3	1.58	0.54	0.16	0.16	24.19	34.46
S4	1.62	0.65	0.13	0.12	19.34	48.08
S5	1.66	0.61	0.19	0.48	23.94	16.27
S6	1.52	0.73	0.22	0.62	23.13	13.38
S7	1.52	0.94	0.11	0.18	20.25	39.58
S8	1.29	0.85	0.14	0.36	20.36	22.26
S9	1.67	0.82	0.02	0.28	25.1	26.10
S10	1.55	0.88	0.04	0.04	24.77	70.21

 Table 21: Mobility factor of Cd.

Sampling Points	F1	F2	F3	F4	F5	Mobility Factor (%)
S1	1.36	2.64	6.29	21.26	30.43	0.60
S2	9.76	6.72	5.81	47.83	36.64	0.92
S3	2.48	2.41	12.27	32.49	32.04	0.46
S4	0.96	3.44	2.23	30.88	24.82	0.56
S5	1.28	1.44	2.26	34.64	31.18	0.25
S6	1.36	2.48	5.81	45.62	21.16	0.39
S7	1.36	2.08	11.4	54.4	26.34	0.24
S8	1.22	2.85	4.2	55.21	30.36	0.24
S9	1.36	2.56	4.21	50.86	30.95	0.25
S10	1.52	1.92	5.42	46.2	30.42	0.24

Table 22: Mobility factor of Cu.

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Statistical Analysis

Analysis of Variance (ANNOVA)

The analysis of variance using single factor for the geochemical phases between and within the geochemical

groups are presented in Tables 23-31 using SPSS IBM 15. The results showed that there are no significant variations between the metals except cadmium but there are significant variations between and within the groups of geochemical speciation.

	SUMMARY									
Groups	Count	Sum	Average	Variance						
Exchangeble Phase	10	75.2	7.52	1.052444						
Carbonate Phase	10	58.94	5.894	1.121849						
Reducible Phase	10	89.8	8.98	10.38747						
Organic Phase	10	96.6	9.66	22.74793						
Residual Phase	10	168.19	16.819	6.736054						
ANOVA										
Source of Variation	SS	df	MS	F	P-value	F crit				
Between Groups	704.1037	4	176.0259	20.93267	8.39E-10	2.578739				
Within Groups	378.4117	45	8.40915							
Total	1082.515	49								

Table 23: Ni.

	SUMMARY										
Groups	Count	Sum	Average	Variance							
Exchangeable Phase	10	25.87	2.587	13.50333							
Carbonate Phase	10	33.7	3.37	7.078444							
Reducible Phase	10	359.59	35.959	600.1978							
Organic Phase	10	277.5	27.75	39.28258							
Residual Phase	10	1156.47	115.647	65.64078							
ANOVA											
Source of Variation	SS	df	MS	F	P-value	F crit					
Between Groups	85872.09	4	21468.02	147.9119	2.82E-25	2.578739					
Within Groups	6531.326	45	145.1406								
Total	92403.41	49									

Table 24: Zn.

SUMMARY								
Groups	Count	Sum	Average	Variance				
Exchangeable Phase	10	30.96	3.096	1.575938				
Carbonate Phase	10	12.22	1.222	0.160662				
Reducible Phase	10	50.93	5.093	6.289912				
Organic Phase	10	28.09	2.809	2.294099				
Residual Phase	10	227.795	22.7795	60.26222				
ANOVA								
Source of Variation	SS	Df	MS	F	P-value	F crit		
Between Groups	3188.203	4	797.0506	56.46207	5.71E-17	2.578739		
Within Groups	635.2455	45	14.11657					
Total	3823.448	49						

Table 25: Co.

SUMMARY								
Groups	Count	Sum	Average	Variance				
Exchangeable Phase	10	309.52	30.952	51.45948				
Carbonate Phase	10	665.5	66.55	368.4856				
Reducible Phase	10	1788.67	178.867	12479.66				
Organic Phase	10	310.58	31.058	77.58173				
Residual Phase	10	1864.98	186.498	617.8134				
ANOVA								
Source of Variation	SS	df	MS	F	P-value	F crit		
Between Groups	243340.5	4	60835.14	22.37409	3.20E-10	2.578739		
Within Groups	122355	45	2719					
Total	365695.5	49						

Table 26: Mn.

SUMMARY							
Groups	Count	Sum	Average	Variance			
Exchangeable Phase	10	82.9	8.29	28.29149			
Carbonate Phase	10	66.32	6.632	12.28304			
Reducible Phase	10	7353.34	735.334	59956.47			
Organic Phase	10	4242.67	424.267	17505.33			
Residual Phase	10	19259.66	1925.966	1378534			
ANOVA							
Source of Variation	SS	df	MS	F	P-value	F crit	
Between Groups	25075699	4	6268925	21.52736	5.60E-10	2.578739	
Within Groups	13104330	45	291207.3				
Total	38180029	49					

Table 27: Fe.

SUMMARY								
Groups	Count	Sum	Average	Variance				
Exchangeable Phase	10	309.52	30.952	51.45948				
Carbonate Phase	10	665.5	66.55	368.4856				
Reducible Phase	10	1788.67	178.867	12479.66				
Organic Phase	10	310.58	31.058	77.58173				
Residual Phase	10	1864.98	186.498	617.8134				
ANOVA								
Source of Variation	SS	df	MS	F	P-value	F crit		
Between Groups	243340.5	4	60835.14	22.37409	3.20E-10	2.578739		
Within Groups	122355	45	2719					
Total	365695.5	49						

Table 28: Pb.

SUMMARY								
Groups	Count	Sum	Average	Variance				
Exchangeable Phase	10	33.82	3.382	0.422818				
Carbonate Phase	10	11.62	1.162	0.543796				
Reducible Phase	10	42.33	4.233	3.488001				
Organic Phase	10	64.08	6.408	5.13164				
Residual Phase	10	280.55	28.055	59.15085				
ANOVA								
Source of Variation	SS	df	MS	F	P-value	F crit		
Between Groups	4849.124	4	1212.281	88.18244	1.07E-20	2.578739		
Within Groups	618.6339	45	13.74742					
Total	5467.758	49						

Table 29: Cr.

SUMMARY								
Groups	Count	Sum	Average	Variance				
Exchangeable Phase	10	15.7	1.57	0.014867				
Carbonate Phase	10	7.43	0.743	0.016623				
Reducible Phase	10	1.28	0.128	0.003751				
Organic Phase	10	2.6	0.26	0.032822				
Residual Phase	10	222.28	22.228	6.496929				
ANOVA								
Source of Variation	SS	df	MS	F	P-value	F crit		
Between Groups	3728.939	4	932.2348	710.0045	5.11E-40	2.578739		
Within Groups	59.08493	45	1.312998					
Total	3788.024	49						

Table 30: Cd.

SUMMARY								
Groups	Count	Sum	Average	Variance				
Exchangeable Phase	10	22.66	2.266	7.090849				
Carbonate Phase	10	28.54	2.854	2.136604				
Reducible Phase	10	59.9	5.99	11.51213				
Organic Phase	10	2.6	0.26	0.032822				
Residual Phase	10	294.34	29.434	18.46745				
ANOVA								
Source of Variation	SS	df	MS	F	P-value	F crit		
Between Groups	5825.948	4	1456.487	185.5877	2.40E-27	2.578739		
Within Groups	353.1587	45	7.847972					
Total	6179.107	49						

Table 31: Cu.

Conclusion

This research has provided background information on the geochemical speciation of the heavy metals in the coastal sediments of Ayetoro area [16]. The geochemical phases include: exchangeable, carbonate, reducible, organic and residual. Previous studies carried out in the area had earlier confirmed sulphide mineralization in the study area. However, most of the elements in each of the phases do not show much variation in concentration with respect to each sampling points. For instance, what is observed in a particular phase is almost in the same range throughout the 10 sampling points except a few cases of different geochemical background and bio-availability. This observation confirmed that, the heavy metals are evenly distributed across the sampling points. The exchangeable and Carbonate phases both naturally have high degree of mobility and bioavailability which confirmed that mobility of metals like Cd that had high concentration in those phases were expectedly high and has the tendency to migrate through all the phases unhindered [17]. Cd is a dangerous heavy metal that has devastating effects on the human body when consumed such as causing kidney problems and renal failures in most cases. It can also cause leukemia. The major sources of pollution of this metal are from indiscriminant dumping of wastes from industries, sewage disposals and from organic leachates. Also, heavy metal bound to the reducible phase are easily released into the environment under reducing chemical conditions

while those in the oxidizable fraction are easily released into the environment under oxidizing conditions whereas, heavy metals bounded in the residual fraction are not easily released into the environment because the metals are firmly bounded within the crystal structure of the mineral comprising the sediment, hence they have low mobility and not biochemically available to influence mineralization but can concentrate to form ore deposits.

Acknowledgements

I want to acknowledge the contributions of my final year research students from the Department of Marine Science and Technology, The Federal University of Technology Akure, Ondo State, Nigeria for their field and technical assistance during the sampling and gathering of vital information towards the successful execution of this research. I am very grateful.to you all.

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