Treatments of produced water from Niger Delta oil fields using simultaneous mixture of local materials

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Abstract
Produced water (PW) from petroleum reservoirs often contains heavy metals and other contaminants that are harmful to the environment. Most of the commonly used treatment techniques have been reported to be ineffective in reducing some of the contaminants’ concentrations to recommended disposal levels. This study evaluated the effectiveness of four selected bio-adsorbents combined for treating PW from Niger Delta oil fields. In this study, orange peels (I), banana peels (II), sponge gourd (Luffa cylindrica) (III) and palm kernel fibers (IV) were washed with distilled water, sun-dried (24 h) and dried in the oven at 105 ± 5 °C (3 h, I and II), 150 °C (30 min, III) and 80 °C (3 h, IV). They were ground into powder, sieved (150 μ, Group A) and (300 μ, Group B), washed with 0.4 mol/L HNO₃, filtered and rinsed with distilled water. Samples of PW were obtained from fields R, X, and Y in the Niger Delta and analyzed for heavy metals using an atomic absorption spectrophotometer (AAS). Samples were treated in adsorption column over 6 h using the adsorbents simultaneously. Treated samples were analyzed with AAS and characterized. Adsorption of heavy metals were assessed using Langmuir and Freundlich models. Data were analyzed using regression and other statistical methods. For the 150 μ size of sample R, the percentage reductions for the metal concentrations (Pb, Ni, Cd, Cu, Fe, Mg, Cr, Zn, Mn, Ca, Ar, B, Sn and Ba) were found to be 100%, 52.7%, 100%, 100%, 85.87%, 19.48%, 100%, 92.8%, 17.74%, 98.86%, 22.32%, 29.56%, 78.06% and 44.74%, respectively, while the reduction in 300 μ size were 1.52%, 97.2%, 71.4%, 17.1%, 43.8%, 45.6%, 7.04%, 89.6%, 35.4%, 99.6%, 0.0001%, 1.19%, 14.19%, respectively. The finer adsorbents were more effective. Similar results were obtained for PW samples from the other fields. Produced water from Niger Delta oil fields was effectively treated of contaminants using four selected bio-adsorbents mixed simultaneously.

Keywords: Produced water · Heavy metals adsorption · Adsorption isotherms · Bio-adsorbents

Introduction/background of the study
The largest waste stream from oil field operations is produced water. Production across the globe is estimated at over 60 million barrels per day. The volume of oil produced from each petroleum reservoir depends on factors such as geological structure, petrophysics, reservoir drive mechanism, depletion stage, well completion method and production practices (Isehunwa and Onovae 2011).

Produced water management has become very important because of the volume involved, cost of treatment and environmental impact of entrapped chemicals. Many oil producing countries have therefore made laws and guidelines on the quality of produced water to be discharged.

In the early days of crude oil extraction from the subsurface, little or no effort was made in handling produced water. In some situations it was discharged to water bodies with little or no form of treatment, spilled on ground surface, or placed in underground pits to evaporate or soak into the subsurface. As time went on, petroleum engineers realized that water injection into the oil reservoirs could enhance production (Gregory et al. 2011).
Crude oil processing

Crude oil–gas–water mixtures that are extracted from reservoirs are passed through flow lines and manifold system, to a treatment facility called the gas–oil separation system. The first stage in the treatment and processing of the well stream is the separation of the oil, gas and water into separate components. This occurs in a two-phase gas–oil separators when the well stream contains no water or in a three-phase separators when water is involved. Well stream separation carried out in these separators is the main process in a train of oil field processing units. The separators are meant to reduce the excessive pressure as a result of gas associated with the produced well crude and, subsequently, separating the gas from the oil. When water is involved in the well stream, separators as well separate the free water from the crude oil. Once separation is achieved, the entire streams separately undergoes more processing and treatment for further field treatment, as highlighted in Fig. 1.

Crude oil emanating from the separator does not meet the purchaser’s specifications and requirements. The crude oil might still contain about 9% and 14% water that exists as emulsified water. The presence of this emulsified water causes corrosion and scaling problems in transportation and processing operations. Water left in the crude oil is called basic sediments and water (BS&W). A maximum of 1% BS&W is acceptable in most cases. The limit on the salt content of the residual water in crude oil is usually between 10 and 15 PTB (pounds of salt per thousand barrels of crude oil). If these requirements are not met at the end, then more treatment of the crude oil emanating from the separator will be needed. Such treatment includes emulsion treatment/dehydration and desalting processes. After crude oil treatment, it may be a necessary to stabilize the crude oil to optimize the oil recovery and minimize its volatility. Most produced crude oils contain hydrogen sulfide and other sulfur compounds. Sour crude oil typically contains total sulfur content > 0.5% by weight. Sour crude oils have safety as well as corrosion problems. Further treatment called sweetening process will be needed to eliminate hydrogen sulfide or reduce its content to a certain limits (Abdel-Aal et al. 2003).

Three-phase oil–water–gas separation

Water produced with the oil exists mostly as free water and most times as water-in-oil emulsion. Sometimes, when the water–oil ratio is much, oil-in-water rather than water-in-oil emulsion will form. Free water extracted with the crude oil from the reservoir is defined as the water that will separate
out and settle from the oil by gravity. In separating the emulsified water, treatment via chemicals, electrostatic treatment, as well as heat treatment, or a combination of them will be needed in addition to gravity settling. Apart from water and oil, gas will always be involved and must be separated from the liquid (water and oil). The quantity of gas depends mostly on the producing and separation conditions. When the quantity of gas is small compared to the quantity of liquid, the method applied to separate free water, oil and gas is known as free water knockout. Under this condition, the separation of the water from the crude oil will guide the design of the apparatus. When there is a larger quantity of gas to be separated from the oil and water, the vessel is known as three-phase separator and either the gas volume requirements or the water–oil separation constraints may guide the vessel design. Figure 2 is an example of a gravity-based three-phase separator; highlighting the gravity settling section, oil and emulsion section as well as the water and oil separation outlet at the lower section of the vessel and the gas outlet at the upper section of the separator.

This current work is focused on evaluating the potential of four adsorbents applied simultaneously in treating produced water. The aim of this study was to reduce the concentration of heavy metals and other contaminants from produced water obtained from Niger Delta oil fields. Specific objectives include; design of adsorption column, treatment of produced water samples with selected bio-adsorbents in the adsorption chamber to reduce heavy metals concentration and other organics present. The methodology of the work involved experimental analysis of four selected bio-adsorbents for produced water treatment over three samples of produced water from three different oil fields.

**Literature review**

In this section of the paper, focus was on reviewing existing produced water treatment techniques; in order to identify key challenges and areas of improvement necessitating the current work.

**Emulsion treatment and dehydration of crude oil**

The multiphase stream produced at the wellhead often consists of gas oil and water mixtures. The gaseous and water contents produced at the wellhead must be removed; while the gaseous components are removed via the separators, the produced water in all its form must be removed by a dehydration process. The removal of the water content of the produced fluid mixture at the wellhead depends on the amount of water in the mixture, the salinity as well as the dehydration process used. Field treatment of water can produce oil with minimal water content up to 0.2 and 0.5 of 1%. The water that remains after the oil has undergone a dehydration is classified as part of the bottom sediments and water (BS&W). The process of treatment and facilities should be carefully selected and designed to meet the contract requirement for BS&W. Care should be taken not to exceed the target oil dryness. Discharge of more remnant water than allowed by contract costs more money while generating less income because the volume of oil sold will be based on the contract value of the BS&W (Khaled et al. 2011).

The basic principles for the treatment process includes:

1. The breaking of the emulsion, which could be achieved by either heating, chemical addition or a combination of addition of heat, the addition of chemicals, and the application of electrostatic field
2. Coalescence of tiny and smaller water droplets into larger droplets
3. Settling, by gravity, and removal of free water.

**Oil emulsions**

Oil extraction from the reservoirs hardly takes place without water accompanying the oil to the surface. Salt water is extracted alongside oil in different forms and methods as illustrated in Fig. 3. Apart from free water, emulsified water (water-in-oil emulsion) is the one form that generates much issue in the dehydration of crude oil. Oil emulsion is simply a mixture of oil and water. Generally, an emulsion can be seen as a mixture of two immiscible liquids, of which one is dispersed as droplets in the other (the continuous phase), and
is stabilized by an emulsifying agent. In the oil reservoirs, crude oil and water are formed as the two immiscible phases together. They form water-in-oil emulsion (W/O emulsion), in which water is dispersed as small droplets in the oil. This is identified as type C in Fig. 4. However, as a spike in the water cut comes up, the possibility of forming reverse emulsions (oil-in-water, or O/W emulsion) increases. This is type B in Fig. 4.

Options available for managing produced water

In produced water management, some of the available options to the oil and gas operators include (Igunnu and Chen 2014):

1. Prevent the production of water unto surface facility—using separators downhole which separate water from gas and oil stream and re-inject it into suitable formation.
2. Injection of produced water—this involves re-injection of produced water into the same formation or other suitable formation; it includes the conveyance of the produced water from the production to the injection site and proper purification of the produced water before injection in order to reduce fouling and scaling agents and possible bacteria growth.
3. Produced water discharge—involves treatment of produced water to meet onshore or offshore discharge regulations. Although, some locations do not require treatment before discharge.
4. Re-use of wastewater for oil field operations—involving the treatment of produced water to meet the required quality for drilling and work-over operations.

5. Used to support other multi-purpose benefits—in some instances, important treatment of produced water is a crucial requirement in order to meet required quality for important benefits such as irrigation, arable land restoration, cattle and animal consumption and drinking water for human consumption.

Water produced during oil and gas production operations constitutes the most prominent waste in the oil industry. The produced water varies both in quality and quantity. Sometimes, the water could be a useful by-product or even a marketable commodity. Produced water is often considered a waste, but the oil industry is beginning to know that this waste water is a potential profit stream. Considering produced water as either a waste or commodity has its cost implication, which needs to be managed in accordance with the specific objectives of each production project and also depending on the region (Fakhrul-Razi et al. 2009). If this is not well addressed, the life of the well could be adversely affected, thereby, leading to huge recoverable reserves being left in the ground. Industry practice on the management of produced water must also protect the environment or the operator could be fined by regulatory authorities. The methodology for managing produced water is a function of the composition of the produced water, the location of the asset in consideration, the volume of reserves and the facilities available in the asset (Gregory et al. 2011).

El-Din et al. (2018) carried out a research work on the potential use of aquatic plant for produced water purification. The aquatic plants used in their research were duckweeds, water hyacinth as well as green algae (Chlorella vulgaris). The plants were prepared and treated before use for the produced water treatment Ahmed et al. (2016); Annadural and Lee (2012). The parameters considered in their research were chemical oxygen demand (COD), biological oxygen demand (BOD). According to their result, COD and BOD were reduced in the produced water by (43% and 42%) by duckweed, 28% and 33% by water hyacinth and 33% and 38% by green algae from the results, duckweeds have higher pollutants removal efficiency than other plants. The limitation of this method is the unavailability of the plants locally within Nigeria.

Aclione et al. (2018), reviewed produced water main treatment technologies. The review was based on the increasing production of oily water and its treatment challenge due to the complexity and volume of water generated, because this undesirable effluent requires treatment before its final disposal in order to meet the legal requirement for disposal in the environment or technical requirements for injection into oil wells. The review highlighted not only for more efficient removal and recovery of oils and other toxic agents, but also financial viability along these technology.

Rasheed et al. (2019) carried out a research on the evaluation of the efficiency of char and biochar for waste water treatment. Char and biochar are carbon rich compounds obtained from pyrolysis. The heavy metals targeted in the waste water were chromium, iron, zinc, nickel, lead and manganese Ahalya et al. (2005); Allen and Brown (2004). From their results, it was concluded that the efficiency of biochar in treating waste water is much better as compared to char as the reduction in heavy metals values were significant with biochar. According to them, biochar could be used as a sorbent for wastewater remediation.

Some other contemporary existing and proposed methods of produced water treatment include the use of membrane technology, the use of novel fibrous coalescence technique, biological treatment in membrane-bioreactors, application of thermal evaporators and the deployment of a novel settling tank for produced water treatment Alzahrani and Mohammad (2014), Dores et al. (2012), Lu et al. (2019) and Shah et al. (2019).

This current study focused on evaluating the potential of four adsorbents applied simultaneously for the treatment of produced water.

Materials and methods

The materials used in this research work were: produced water samples from Niger delta oil fields, Luffa cylindrica (sponge gourds), banana peels, orange peels, palm kernel fiber, chemical reagents, filter paper and glass wares, etc. The equipment used were: milling machine, adsorption column, mechanical shaker, oven, atomic absorption spectrophotometer, weighing balance, pH meter, refrigerator and sieves, etc. The heavy metals analyzed were lead (pb), nickel (Ni), cadmium (Cd), copper (Cu), iron (Fe), magnesium (Mg), chromium (Cr), zinc (Zn), calcium (Ca), Boron (B), Tin (Sn), arsenic (As) and Manganese (Mn) as well as Barium (Ba) and some other contaminants in the produced water samples such as: carbonates, bi-carbonate and acidic components, etc.

Collection of samples and materials

Samples of produced water were collected from four different oil fields in the Niger Delta area. Sample R was collected from Imo River oil field in Rivers State; Sample X was collected from Nembe oil field in Bayelsa State, sample Y was from Kolo creek oil field in Rivers/Bayelsa State. The adsorbents (Banana peels, Orange peels, Luffa cylindrica) were purchased from a local market in Ado-Ekiti, Nigeria.
and the palm-kernel fiber was collected from ABUAD farm in Ado-Ekiti, Ekiti state. Chemicals such as: nitric acid, hydrochloric acid, sodium sulphate, phenolphthalein and sulphric acid as well as other reagents used were of analytical and standard grade.

Figure 5 captures the *Luffa cylindrica* and palm kernel fiber samples, which were used for the treatment of the produced water samples obtained from the oil field.

In Fig. 6, banana peels and raw orange peels also used for the treatment of the produced water samples are also captured.

Figure 7 captures some of the produced water samples and filtrates from adsorption process, as samples for the experiment conducted.

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**Fig. 5** *Luffa cylindrica* and palm kernel fiber

**Fig. 6** Banana peels and raw orange peels

**Fig. 7** Produced water samples and filtrates from adsorption process
Preparation of banana peels, orange peels, *Luffa cylindrica*, palm kernel fiber and the experimental procedure

The adsorbents were prepared by washing them thoroughly with distilled water to remove any dirt that may be stuck on it. They were then cut into pieces, for easy handling; suns dried (24 h during hammer time period), and were finally dried in an oven at 105 °C for 3 h for banana and orange peels, 150 °C for 30 min for *Luffa cylindrica*, and 80 °C for 3 h for palm kernel fibre.

Simultaneous mixture of the adsorbents in the adsorption column

After drying, the adsorbents were crushed using the milling machine and were sieved with the mechanical shaker using mesh size of 150 and 300 μ. The sieved samples were treated separately with 200 mL of 0.4 mol/L HNO₃ for 24 h. After which they were filtered and rinsed with distilled water until the filtrate was near neutral. 2-Gram each of the 150 μ particle size of the four (4) adsorbents was measured to sum up to 8-g in total. The adsorbents were mixed together simultaneously at a ratio of 1:1:1:1 within the adsorption column. 250 mL of sample R (produced water) was flown through the adsorption column containing the adsorbents to allow adsorption to take place. The filtrate was collected at an interval of 2 h and was subjected to analysis using AAS to determine the concentration of the heavy metals listed above. The mixture of the adsorbents on the column was replaced with a fresh one and sample X and Y was flown in that order through the adsorption column and the filtrate of each was collected at an interval of 2 h as well and analyzed. The process was repeated all over again using a particle size of 300 μ.

Figure 8 shows the adsorption column, where the four (4) adsorbents were mixed together to achieve simultaneous mixture as key part of the experimental work.

Results and discussion

The results presented in Tables 1, 2, 3 and 4 were obtained from the simultaneous mixture of the adsorbents used for the treatment of produced water samples (R, X, Y) in the adsorption column. The results from each sample were for both 150 and 300 μ particle size. The plots in Fig. 9 extracted from Table 1 were more of linear form showing that the more the contact time, the more the produced water sample loses its toxic metals. At 8 h with 150 μ particles size, lead (Pb), cadmium (Cd), cupper (Cu) and chromium (Cr) were all completely removed from the produced water sample. This means that Pb, Cd, Cu and Cr were reduced to zero from their raw values of 0.132, 0.014,
0.076 and 0.071–0.00 mg/L each. Other metals (nickel, iron, magnesium zinc, manganese, calcium, arsenic, boron and tin barium etc.) were as well reduced to the barest minimum beyond the regulatory limits as set by the regulatory bodies. The result obtained with the 300 μ particle size as captured in Fig. 10 were equally good as the metals were all affected positively with respect to the reduction in their concentrations on the produced water. The metals were all reduced to allowable limits though none of them could be removed completely as can be seen when compared with 150 μ particle size. This goes to show that in the adsorption process, contact time and particle size matter a lot as they are among the determining factors for effective and efficient adsorption process. The more the surface area of the adsorbents and the more the contact time, the better the adsorption process.

The plots (Fig. 11 from Table 2) are all linear with negative slopes; it shows that the more the contact time between the adsorbents and the produced water sample, the more the metal concentrations in the produced water go down. Lead (Pb) and copper (Cu) were completely adsorbed from the sample after 8 h of treatment while other metals were positively affected but not completely removed. The metals removal depends on the activeness of the adsorbents. The more the availability of the surface of the adsorbents (adsorption sites) as a result of the large adsorption surface, the more the metals are adsorbed.

From the plots in Fig. 12 for 300 μ size, very good results were also obtained just like that of 150 μ particle size. The concentration of almost all the metals investigated were reduced to the barest minimum as required by the regulatory bodies.

| Table 2 | Results of the concentrations (mg/L) of metals in sample X (Nembe sample) for simultaneous mixture |
|---------|----------------------------------------------------------------------------------|
| Sample X | pb Ni Cd Cu Fe Mg Cr Zn Mn Ca Ar B Sn Ba |
| Raw     | 0.078 0.026 0.007 0.07 0.783 5.106 0.121 0.163 0.071 390 5.544 2.151 0.144 0.045 |
| After treatment with adsorbents using 150 μ particles size |
| 2 h     | 0.068 0.024 0.006 0.058 0.422 4.722 0.099 0.161 0.067 200 5.297 2.011 0.141 0.042 |
| 4 h     | 0.054 0.023 0.005 0.003 0.233 4.281 0.097 0.143 0.055 120 5.024 1.722 0.122 0.035 |
| 6 h     | 0.007 0.022 0.005 0.001 0.007 3.445 0.092 0.089 0.049 44 4.188 1.323 0.077 0.031 |
| 8 h     | ND 0.022 0.004 ND 0.005 3.256 0.082 0.059 0.036 4.8 3.612 1.121 0.034 0.021 |
| After treatment with adsorbents using 300 μ particle size |
| 2 h     | 0.074 0.026 0.007 0.068 0.544 4.721 0.121 0.161 0.069 250 5.542 2.151 0.142 0.043 |
| 4 h     | 0.058 0.024 0.006 0.052 0.038 3.223 0.098 0.098 0.057 120 5.412 2.010 0.133 0.043 |
| 6 h     | 0.033 0.023 0.006 0.041 0.023 2.101 0.094 0.095 0.053 70.52 4.231 2.005 0.127 0.041 |
| 8 h     | 0.023 0.023 0.005 0.024 0.016 1.401 0.091 0.095 0.039 1.41 3.822 1.994 0.127 0.035 |

| Table 3 | Results of the concentrations (mg/L) of the metals in sample Y (Kolo creek sample) for simultaneous mixture |
|---------|----------------------------------------------------------------------------------|
| Sample Y | pb Ni Cd Cu Fe Mg Cr Zn Mn Ca Ar B Sn Ba |
| Raw     | 0.192 0.231 ND 0.038 0.805 5.578 0.397 0.174 0.081 505 4.522 1.921 1.74 0.091 |
| After treatment with adsorbents using 150 μ particles size |
| 2 h     | 0.188 0.189 ND 0.035 0.651 4.981 0.281 0.172 0.074 308 3.221 1.722 1.505 0.072 |
| 4 h     | 0.185 0.164 ND 0.033 0.362 3.554 0.179 0.158 0.058 150 2.792 1.441 1.449 0.058 |
| 6 h     | 0.171 0.124 ND 0.021 0.244 2.482 0.155 0.132 0.023 55.88 2.204 1.202 1.322 0.051 |
| 8 h     | 0.171 0.118 ND ND 0.172 1.326 ND 0.112 0.019 1.11 1.678 1.142 1.228 0.045 |
| After treatment with adsorbents using 300 μ particle size |
| 2 h     | 0.184 0.231 ND 0.037 0.755 5.499 0.395 0.173 0.075 300 4.442 1.898 1.722 0.085 |
| 4 h     | 0.066 0.188 ND 0.035 0.621 5.355 0.382 0.165 0.071 160 4.311 1.852 1.667 0.065 |
| 6 h     | 0.043 0.094 ND 0.031 0.433 3.221 0.341 0.122 0.055 40 3.311 1.733 1.552 0.032 |
| 8 h     | 0.008 0.028 ND ND 0.291 0.244 0.332 ND ND ND 3.201 1.733 1.499 ND |

| Table 4 | Analysis of Pb (lead) using Langmuir and Freundlich models for simultaneous mixture of the adsorbents using sample R |
|---------|----------------------------------------------------------------------------------|
| t (h)   | $C_e$ (mg/L) | $q_e$ (g/L) | $C_e/q_e$ (g/L) | Log $C_e$ (mg/L) | Log $q_e$ (g/L) |
| 2       | 0.112 | 0.000625 | 179.200 | −0.9508 | −3.2041 |
| 4       | 0.027 | 0.00328 | 8.231 | −1.5686 | −2.4841 |
| 6       | 0.011 | 0.00378 | 2.910 | −1.9586 | −2.4225 |
| 8       | 0.000 | 0.00413 | 0.000 | 0.000 | −2.3840 |
body though better results were obtained with 150 \( \mu \) size because of larger surface area.

The plots in Fig. 13 extracted from Table 3 were all linear with negative slopes; it shows that the more the contact time between the adsorbents and the produced water sample, the more the metal concentrations in the produced water drop. Copper (Cu) was found to be completely removed from the sample after 8 h of treatment. Based on observation, the produced water sample from Kolo creek oil field did not contain Cadmium at all. This could be as a result of the location of the field.

Figure 14 is a plot of Pb, Ni, Cd and Cu versus time (mg/L) against time (h) for 300 \( \mu \) particle size. The plots are linear with same negative slopes as the previous one of 150 \( \mu \) size. From the plots, copper (Cu) was completely removed from the sample after 8 h of treatment. Zn, Mn, Ca and Ba were also completely eliminated from the sample after 8 h of treatment according to Table 3.

In summary it was observed that adsorption process is more favorable when the surface area of an adsorbent is large. The larger the surface areas, the smaller the particles and the more the adsorption sites for effective adsorption process. From the research, better results were obtained with 150 \( \mu \) size because of its large surface area compared with 300 \( \mu \) size. Simultaneous mixture of the adsorbents has proved worthwhile in the treatment of produced water from Niger Delta oil field. The results obtained were in accordance with the discharge limits as stated by the regulatory bodies.
Analysis of the results using adsorption isotherms

The Langmuir model

This isotherm model is used to express the relationship that exists between the quantity of extracted material and its equilibrium concentration in bulk of solutions. Langmuir isotherm is only acceptable (valid) for monolayer adsorption on a surface containing finite number of similar sites. Langmuir isotherm model assumes a uniform adsorption on the surface and transmigration in the plane of the surface. Langmuir isotherm can be expressed as:

\[ q_e = \frac{K_L C_e}{1 + b C_e} \]
where $q_e$ is capacity of adsorption at equilibrium in (mg/g), $C_e$ the equilibrium concentration (mg/L) and $K_L$ is the Langmuir constant in (mL/mg) (Table 5).

### The Freundlich model

Freundlich isotherm model is a model that considers adsorption process to occur on heterogeneous surfaces; and the model further states that adsorption capacity is related to the concentration of the adsorbent in question. The Freundlich model can be expression as:

$$q_e = K_F C_e^{1/n}$$

Note $K_F$ is the Freundlich constant and $1/n$ is a constant showing the reaction intensity. These Freundlich parameters $K_F$ and $1/n$ can be estimated graphically from the plot of...
revealed that the sorption of Pb ion on the simultaneous mixture of the adsorbents was found to be a favorable process. The correlation coefficient from Freundlich isotherm plot of $\log q_e$ versus $\log c_e$ as captured in Fig. 16 could not give a better result, showing that Freundlich sorption isotherm model is not suitable for equilibrium study of Pb (Fig. 17).

For the Ni adsorption, the linear plot of $\log q_e$ versus $\log c_e$ indicates that adsorption obeys the Freundlich sorption model as shown in Fig. 18. The values of $K_F$ and $n$ were determined from the slope and intercept of the plot respectively as captured in Table 6. The correlation coefficient ($R^2$) was estimated to be 0.9835. This shows that the experimental values and then applying the Freundlich equation in this form:

$$\ln q_e = \ln K_F + \left(\frac{1}{n}\right) \ln c_e$$

The linear plot of $c_e/q_e$ versus $c_e$ indicates that adsorption process follows the Langmuir model as shown in Fig. 15. The values of $Q$ and $b$ were obtained from the slope and intercept of the plotted data in Table 6. The correlation coefficient ($R^2$) was found to be 0.9676 for Pb and 0.0038 for same Pb as captured in Fig. 16. The results revealed that the Langmuir sorption isotherm model is good for equilibrium study for Pb. This suggests the formation of monolayer coverage of the adsorbate on the adsorbent surface for the metal ion.

The quantity of metal ion extracted per unit mass of the adsorbent rises with the metal concentration as expected and the sorption capacity was also determined. The value of the equilibrium separation factor $R_L$, of 0.812 (Table 6)
Freundlich sorption isotherm is good for equilibrium study of Ni. This suggests the formation of heterogeneous layer coverage of the adsorbate on the adsorbent surface for the metal ion. The values of $K_F$ and $n$ (Table 6) indicate the adsorption process is favorable. Although the Langmuir isotherm model result for Ni adsorption as captured in Fig. 17 indicated an increasing trend for $C_e/q_e$ (g/L) versus $C_e$, it was not a bad result; as fortunately, Freundlich model gave a better result with respect to Ni adsorption on the adsorbent surface.

For Fe (iron) adsorption on the simultaneous mixture of the adsorbents, Freundlich-type isotherm model gives a better result than Langmuir isotherm model. The plot of log $q_e$ versus log $C_e$ indicates that adsorption obeys the Freundlich adsorption model as captured in Fig. 20. The values of $K_F$ and $n$ as indicated on Table 6 indicate that the sorption of Fe on the adsorbent surface is a favorable process (Fig. 20).

**Conclusion**

In this work, an adsorption chamber was successfully designed and used for the treatment of the produced water samples from Niger Delta Oil fields and the adsorption of heavy metals were evaluated based on Langmuir and Freundlich models. Simultaneous mixture of the adsorbents was very effective in the treatment of the produced water samples.

Based on statistical analysis, for the 150 μ size of sample R, the percentage reductions for the metal concentrations (Pb, Ni, Cd, Cu, Fe, Mg, Cr, Zn, Mn, Ca, Ar, B, Sn and Ba) were found to be 100%, 52.7%, 100%, 100%, 85.87%, 19.48%, 100%, 92.8%, 17.74%, 98.86%, 22.32%, 29.56%, 78.06% and 44.74%, respectively, while the reduction in 300 μ size were 1.52%, 97.2%, 71.4%, 17.1%, 43.8%, 45.6%, 7.04%, 89.6%, 35.4%, 99.6%, 0.0001%, 1.19%, 14.19% and 0.002%, respectively. Hence, the finer adsorbents were more effective in reducing the presence of heavy metals in the treated produced water.

The concentration of most of the heavy metals in the water samples were reduced to an acceptable level for discharge; below the standard disposal level (Ca—200 mg/L, Ba—0.01 mg/L, Sn—0.04 mg/L, B—1.0–0.5 mg/L, Ar—5 mg/L, Mn—0.02 mg/L, Zn—1.0 mg/L, Cr—0.02 mg/L, Mg—60 mg/L, Fe—1.0 mg/L, Cu—0.02 mg/L, Cd—0.01 mg/L, Ni—0.03 mg/L and Pb—0.05 mg/L).

Few other metals were found to have raw values (concentration before treatment) less than the discharge limit which is a good development. This method of produced water treatment is strongly recommended for oil industry and other wastewater generating industries. The local materials (adsorbents) are readily available, very economical and are environmentally friendly.

**Recommendations**

Further study should look into modification of the bio-adsorbents to avoid early degradation with time. Degradation of the bio-adsorbents result to loss of active site and this leads to poor adsorption process.

Other waste materials apart from the ones used in this research should be look into to see how effective and efficient they could be in produced water treatment.

The scope of the particle size should be extended and the results compared with the ones used in this research work.
Research attention should be shifted from those conventional ways of produced water treatment. These conventional methods like micro filtration, precipitation, flotation and sedimentation, etc., are very capital intensive and pollute the environment. Researcher should be educated on the need to phase out these methods and imbibe the culture of using the locally and available adsorbent.

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