Arsenic contamination levels in drinking water sources in mining areas in Lake Victoria Basin, Tanzania, and its removal using stabilized ferralsols

Gabriel R. KASSENGA * and Rubhera R. MATO

Department of Environmental Science and Management, Ardhi University (ARU), P. O. Box 35176, Dar es Salaam, Tanzania.

*Corresponding author, E-mail: kassengagr@yahoo.com, kassenga@aru.ac.tz, Tel. (255 022) 2771272, 2775004 (Office), Fax: (255 022) 2775448, 2775479

ABSTRACT

Arsenic contamination in drinking water is a public health problem all over the World especially in mining areas. The study herein reported assessed the concentration levels of arsenic in some drinking water sources in the mining areas in the Lake Victoria Basin and investigated the potential for its removal by adsorption using stabilized ferralsols. Randomly collected samples from drinking water sources were analysed for arsenic, phosphorus, iron and pH. Arsenic was detected in 58% of water sources surveyed and 41% of them had arsenic levels equal to or exceeding the Tanzania Drinking Water Quality Standards threshold value of 10 ppb. Arsenic mobilization is suspected to be due to oxidation of arsenopyrite after exposure of the ore to the air partly due to mining activities. Laboratory experiments have shown that adsorption using stabilized ferralsols (1:1 mixture of Portland cement and ferralsols on dry weight basis) is capable of removing up to 97% of arsenic independent of water pH and initial contaminant concentrations used during the study. In view of this, adsorption using stabilized ferralsols may be considered to be a promising technology for removing arsenic from drinking water.

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Keywords: Arsenopyrite, Portland cement, Adsorption isotherms, Groundwater, Efficiency.

INTRODUCTION

Arsenic is a highly toxic metalloid that is brittle, crystalline, odourless and tasteless in its elemental form (Kassenga, 2003). Naturally, arsenic occurs primarily as arsenopyrite (FeAsS), which is the most abundant arsenic mineral, dominantly in mineral veins (Halsey, 2000). Smedley and Kinniburgh (2002) describe different manifestations of arsenic which are basically inorganic and organic forms; inorganic species are mainly arsenite, As(III), and arsenate, As(V). The inorganic species of arsenic (arsenite and arsenate) are the most important species with relevance to toxicity in humans (Halsey, 2000; Kassenga, 2003). Consequently, these two species have been the centre of focus of many arsenic researches (Korte and Fernando, 1991; Vance, 1995; Halsey, 2000) including the present study.

The significance of arsenic speciation is based on toxicity and remedial capability (Vance, 1995). Arsenite is the most toxic of all the species of arsenic. Arsenite is about 60 times more poisonous than arsenate and 70 times more toxic than the methylated species (Korte and Fernando, 1991). Besides, as far as arsenic removal in contaminated water is concerned, arsenite is difficult to be removed by using many conventional and innovative remediation techniques (Korte and Fernando, 1991; Halsey, 2000).

The World Health Organization (WHO) (2001) has set the International Drinking Water Guideline for arsenic at 10 ppb. This is a strict maximum value because of high toxicity caused by this chemical. The
major early manifestations due to acute arsenic poisoning include burning and dryness of the mouth and throat, dysphasia, colicky abnormal pain, projectile vomiting, profuse diarrhoea, and hematuria (Kassenga, 2003). The muscular cramps, facial edema and cardiac abnormalities, shock can develop rapidly as a result of dehydration (Jain and Ali, 2000).

According to Smith et al. (2000), approximately 20 million people are thought to be exposed to arsenic poisoning and another 50 million are considered to be at risk of arsenic poisoning after the installation of over 4 million tube wells in Bangladesh. Arsenic levels higher than 50 ppb are common and levels as high as 2,000 ppb have been reported and generally, many wells were observed to exceed the WHO drinking water standard of 10 ppb (Smith et al., 2000).

In Ghana, high arsenic concentrations have been noted in soils and river waters close to the gold mining activity (Smedley, 1996). This is because gold containing ore is associated with sulphide mineralization, particularly arsenopyrite (Smedley and Kinniburgh, 2002). Arsenic contamination from mining activities has been identified in numerous areas of the USA, many of which have been summarized in Welch et al. (2000). According to Welch et al., (2000), groundwater from some areas has been reported to have very high arsenic concentrations of up to 48,000 ppb.

The Lake Victoria Basin is the leading gold producing area in Tanzania. There is a number of goldfields in the area (collectively known as Lake Victoria Goldfields) and they are located in the Archean Nyanzian greenstone belt east and south of Lake Victoria in northwest Tanzania (Department of Geology, 1994). A short term assessment study on mercury and other heavy metal pollution in gold mining areas around Lake Victoria conducted in 1994 by Department of Geology of the University of Dar es Salaam, Tanzania showed the presence of arsenic with concentrations, which ranged from 0.5 to 379 ppb in some water sources (Department of Geology, 1994). Suspected source of arsenic pollution in the sediments and water was identified to be oxidation of arsenopyrite in mine tailings dumped into the rivers during gold panning.

Arsenic toxicity has no known effective treatment, but drinking of arsenic free water can help arsenic affected people at early stage of ailment to get rid of the symptoms of arsenic toxicity. Therefore, the most important measure needed is to prevent further exposure of population by providing them with arsenic-free safe drinking water. There exist a number of treatment processes to remove arsenic from drinking water including sulphide precipitation, co-precipitation with iron and metal hydroxides, coagulation and adsorption processes. Methods, which are usually used for removal of arsenic from drinking water in developed countries such as lime softening, ion exchange resins, membrane methods and activated alumina (Johnston and Heijnen, 2001) are often too expensive to be used in developing countries. Socio-economic conditions of Tanzania and other less developed countries demand low-cost as well as small-scale treatment systems that could be implemented in the rural areas at household or community levels. Thus, there is an urgent need to develop simple to use and cost efficient treatment technologies capable of removing arsenic from drinking water.

The general objective of the study for which this paper was prepared was to conduct a rapid assessment on the contamination levels of arsenic in drinking water sources in mining areas of the Lake Victoria Basin in Tanzania and to investigate the potential of stabilized ferralsols in removal of the contaminant by adsorption for future design of household and small community water treatment systems. Specifically, the study quantified the concentration levels of arsenic in drinking water sources and determined the arsenic adsorption isotherms of stabilized ferralsols as well as the removal performance of the adsorbent focusing on the influence of water pH and initial arsenic concentration.

MATERIALS AND METHODS

Study Area

Field studies involved onsite measurements of concentrations of arsenic, iron and phosphorus in drinking water sources. These studies were conducted in the Lake Victoria Basin in Tanzania, which covers some parts of the regions of Shinyanga, Mwanza, and Mara. Water sources sampled were specifically located in districts
of Bariadi, Maswa, Meatu, Kahama and Bukombe in Shinyanga region, Musoma, Buhemba, and Serengeti in Mara region and Mwanza urban, Sengerema, Geita, and Kwimba in Mwanza region as shown in Figure 1. The total population in the sampled districts is around 4.2 million people, which is approximately 12% of the entire population of the country (URT, 2002).

Geochemical Modelling

The influence of pH, ferric iron and phosphorus as well as other geochemical conditions on speciation of arsenic were analysed using PHREEQC computer model (Version 2 for Windows). PHREEQC is a computer program that is designed to perform a wide variety of low-temperature aqueous geochemical calculations. The model is based on an ion-association aqueous model and has capabilities for (i) speciation and saturation-index calculations; (ii) batch-reaction; and, (iii) one-dimensional transport calculations involving reversible reactions (Parkhurst and Appelo, 1999). Model inputs were reactions of arsenic in aquifer media, pH, concentrations of iron, phosphate and arsenic whereas the outputs were concentrations of various complexes of arsenic.

Laboratory Experiments

Materials

a) Chemicals and reagents

Neat solution of arsenic trioxide \( \text{As}_2\text{O}_3 \), (1 mg/mL) in 0.5 M/L HCl (BDH Chemicals Ltd., Leicestershire, UK) was used to prepare stock solutions of arsenic. Iron and phosphorus standard solutions were obtained from Hach Co. (Loveland, CO). Other chemicals used in this study such as HNO\(_3\), and NaOH were reagent grade and most of them were purchased from BDH Chemicals Ltd. (Leicestershire, UK).

Figure 1: Location maps of Tanzania in Africa (A) and the study area (B).
b) Preparation of adsorbent (stabilized ferralsols)  

Ferralsols used in this experiment was collected and prepared according to the modified procedures specified in Kundu et al. (2004). In brief, ferralsols was soaked and stirred in distilled water and then left in quiescence condition for 1 hour after which, suspended organic matter were removed. The ferralsols was then dried for 24 hours at room temperature of 25 ± 1 ºC.

Three hundreds grams (300 g) of ferralsols and 300 g (ratio of 1:1 by dry weight) of commercially available Portland cement (Tanzania Portland Cement Ltd, Dar es Salaam, Tanzania) were mixed with sufficient amount of distilled water to form a homogeneous slurry mixture. The slurry mixture was then stored for 72 hours in a plastic container at room temperature (25 ± 1 ºC) for hardening.

After the hardened stabilized ferralsols was completely air-dried, it was broken into particles of approximately 300 mm using a sledgehammer. For further stabilization of bonds between cement and ferralsols, the granules were submerged in water for 96 hours.

The granules were further reduced in size using a sledgehammer. The required sizes of stabilized ferralsols were obtained by sieving whereby two sieve sizes of 1.5 mm and 3.0 mm were used to arrive at particle sizes, which ranged between 1.5 and 3.0 mm. The granules were finally oven-dried for 24 hours between 100 and 110 ºC before use.

Adsorption experiments

The main objective of adsorption experiments was to determine the capacity of stabilized ferralsols in adsorbing arsenic under various pH conditions. Adsorption experiments were done using tightly covered 1000 mL glass beakers containing 500 mL of arsenic solution with varying concentrations and 10 g of ferralsols (20 g/L). pH of arsenic solution was adjusted by adding 0.1N HCl or 1M NaOH to obtain pH values of 4, 5, 6, 7, 8, and 9 (Standard error for each measured pH value was ± 0.1). For each pH value, three experiments were performed for a total of 15 isotherm points. Isotherm points were generated by diluting the arsenic solution with an electrolyte solution (0.01 M CaCl₂) using dilutions of 1:1, 1:2, 1:4, 1:8 and 1:16 to obtain concentrations of 20, 10, 5, 2.5 and 1.25 mg/L. Control experiments were also conducted for each pH using the adsorbents and distilled water.

Contents of beakers were thoroughly mixed to facilitate adsorption by stirring at 25 ± 1 ºC for 8 hours at 70 revolutions per minute using the jar tester (PB-700™ Standard JarTester, Phipps & Bird, Inc., Richmond, Virginia), centrifuged and the clear supernatant analyzed for arsenic using Atomic Absorption Hydride System (HC 3000).

Arsenic removal efficiency

Assessment of arsenic removal efficiency by adsorption onto stabilized ferralsols matrix at different pH values (4, 5, 6, 7, 8 and 9) and arsenic concentrations of 2.5, 5.0, 10.0 and 20.0 mg/L was conducted in triplicate by batch experiments. Batch experiments were conducted in a similar manner as adsorption experiments as described above.

Analytical methods

Field analysis of arsenic was done using Arsenic Test Kit (Hach Company, Loveland, CO), which has a detection limit of 5 ppb. Aqueous concentrations of ferrous iron and phosphorus were analysed using a colorimetric method with a portable spectrophotometer (DR 2010; Hach Co., Loveland, CO). Global Positioning System (GPS) was used to record the locations of the sampling points.

Chemical compositions of ferralsols and stabilized ferralsols were measured using Sequential X-ray Spectrometer (SRS 300) from Siemens equipped with a Rh 65 end-window tube (Beryllium, 125 µm, using the Rh-. L-line). Sample preparation involved mixing 5 g of pulverized sample with 8 g boric acid which, was used as a binder. The mixture was then subjected to a pressure treatment of 2 kN/m² for one minute to obtain pellets of the sample with a thickness of 4 mm. Pellets were finally analysed both qualitatively and quantitatively using the Sequential X-ray Spectrometer.

Laboratory analysis of arsenic was done using Atomic Absorption Hydride System (HC 3000) from GBS Scientific Equipment (Arlington Heights, IL). The detection limit of this method is 0.5 ppb.
Computational Framework

Adsorption efficiency

The adsorbed concentration of arsenic was calculated from the measured equilibrium concentrations as:

\[ q_e = \left( c_i - c_e \right) \times \frac{V}{m} \quad (1) \]

where \( q_e \) (mg/kg) is adsorbed concentration; \( c_i \) (mg/L) is initial arsenite concentration; \( c_e \) (mg/L) equilibrium concentration; \( V \) (L) is volume of the solution; and, \( m \) (kg) is weight of the stabilised ferralsols.

Percentage removal of arsenic was calculated from the measured equilibrium concentrations as follows:

\[ \% \text{Removal of arsenic} = \frac{c_i - c_e}{c_i} \times 100\% \quad (2) \]

Adsorption Isotherms

Adsorption of arsenic onto stabilized ferralsols matrix was described using Langmuir and Freundlich models. The linearized version of Langmuir sorption model was used in the present study:

\[ \frac{1}{q_e} = \frac{1}{Q_{\text{max}}} + \frac{1}{bQ_{\text{max}}} \left( \frac{1}{C_e} \right) \quad (3) \]

where \( q_e \) is adsorbed contaminant concentration (mass of contaminant/mass of adsorbent); \( b \) is equilibrium constant for the sorption reaction; \( C_e \) is dissolved contaminant concentration; \( Q_{\text{max}} \) is number of sorption sites (maximum amount of sorbed contaminant).

Nonlinear isotherms were described by the Freundlich equation:

\[ q_e = kC_e^{\frac{1}{n}} \quad (4) \]

Statistical Analysis

ANOVA was used to determine if the effect of pH on removal of arsenic by adsorption onto stabilized ferralsols matrix was indeed significant. The t-test was used to determine the differences in values of coefficient of determination between Freundlich and Langmuir sorption models. The results were considered statistically significant if \( P < 0.05 \).

RESULTS

Levels of arsenic concentration in drinking water sources

A total of 96 drinking water sources were sampled for analysis of arsenic, iron, phosphorus and pH. Table 1 shows measured concentrations of arsenic in the districts surveyed. It should be noted that Table 1 reports arsenic concentrations only in districts where the contaminant was detected.

Composition of ferralsols and stabilized ferralsols

The composition of ferralsols and stabilized ferralsols is depicted in Table 2. Portland cement has the advantage of improving the structure of ferralsols and hence its adsorption capacity by increasing the amount of CaO (Olisio et al., 2002).

| S/N | District | Range (ppb) | Mean ± Standard Deviation (ppb) |
|-----|----------|-------------|---------------------------------|
| 1.  | Bariadi  | 5-10        | 8.0 ± 2.7                       |
| 2.  | Maswa    | 5-70        | 18.3 ± 21.14                    |
| 3.  | Shinyanga| 5-50        | 20.7 ± 20.29                    |
| 4.  | Musoma   | 5-40        | 17.2 ± 12.77                    |
| 5.  | Kahama   | 5-30        | 12.5 ± 11.9                     |
| 6.  | Bukombe  | 5-10        | 9.3 ± 1.88                      |
| 7.  | Geita    | 5-30        | 11.3 ± 7.91                     |
In the present study, mixing of Portland cement with ferrasols increased the concentration of CaO by a factor of 47 (Table 2). Furthermore, Table 2 shows that mixing ferrasols with Portland cement reduced the concentration of phosphorus by 33% relative to un-amended ferrasols. Although mixing reduced the concentration of ferric oxide by 15% relative to un-amended ferrasols, the amount present in the mixture is still higher than the minimum threshold value of 1.6%, below which impairing of adsorption by phosphorus may occur.

**Arsenic removal efficiency of the adsorbent**

The effect of pH and arsenic concentration on arsenic removal efficiency is presented in Figure 2.

| Table 2: Concentration and composition of ferrasols and stabilized ferrasols (mixture of Ferrasols and Portland cement at a ratio of 1:1 (w/w)). |
|---------------------------------------------------------------|
| **Constituent** | **Concentration (mol/kg dry weight)** | **Percent Composition (w/w)** |
| | **Mixture** | **Ferrasols** | **Mixture** | **Ferrasols** |
| SiO₂ | 24.79 | 53.31 | 29.8 | 64.08 |
| Al₂O₃ | 6.32 | 11.18 | 12.9 | 22.8 |
| Fe₂O₃ | 1.32 | 1.56 | 4.21 | 4.97 |
| MnO | 0.06 | 0.07 | 0.09 | 0.10 |
| MgO | 0.45 | 0.30 | 0.36 | 0.24 |
| CaO | 30.46 | 0.65 | 34.6 | 0.74 |
| Na₂O | 0.10 | 0.09 | 0.12 | 0.11 |
| K₂O | 0.20 | 0.27 | 0.37 | 0.51 |
| P₂O₅ | 0.03 | 0.04 | 0.08 | 0.12 |
| SO₃ | 1.84 | 0.08 | 2.95 | 0.13 |
| TiO₂ | 0.19 | 0.26 | 0.56 | 0.79 |
| LiO | 30.08 | 11.29 | 13.8 | 5.18 |
| Minor constituents | - | - | 0.09 | 0.03 |
| **Total** | 100.00 | 100.00 |

**Figure 2**: Influence of pH and arsenic concentration on arsenic removal efficiency
Adsorption isotherms

Adsorption isotherms of arsenic are shown in Figure 3 and 4 whereas Table 3 presents parameter values of the Langmuir and Freundlich adsorption models.

DISCUSSION

Arsenic contamination levels

Arsenic was detected in 58% of samples analyzed. In water sources, which tested positive, the observed arsenic concentration values ranged from 5 to 70 ppb (Table 1). Forty-one percent (41%) of samples analyzed had arsenic concentrations exceeding the Tanzania Drinking Water Quality Standards threshold value, which is 10 ppb. Arsenic contamination in the study area was observed to be less serious compared to the situation in other parts of the world reported elsewhere (Welch et al., 2000; Smith et al., 2000). For example, arsenic concentration of 50 ppb is common in Bangladesh and concentrations of up to 2,000 ppb have been reported in the country (Smith et al., 2000). In USA, concentrations as high as 48,000 ppb were observed in Fairbanks gold mining district in Alaska (Welch et al., 2000). Both America and Bangladesh have the maximum permissible limit of arsenic in drinking water of 50 ppb, which is technically and politically acceptable bearing in mind that the levels of the contaminant are high and its spatial distribution is wide.

Ashanti gold mining region in Ghana has higher arsenic concentrations (from 1 to about 140 ppb) than those observed in the Lake Victoria Basin and no significant health effects have been reported in the region (Smedley, 1996; Mukherjee et al., 2006).
Table 3: Values of adsorption parameters.

| pH  | Temperature (ºC) | Langmuir Constants | Freundlich Constants |
|-----|------------------|---------------------|----------------------|
|     |                  | Q<sub>max</sub> (µg/g) | b (µg/g) | R<sup>2</sup> | k | l/n | n | R<sup>2</sup> |
| 4±0.1 | 25±1 | 769 | 0.0087 | 0.890 | 16.66 | 0.58 | 1.7 | 0.980 |
| 5±0.1 | 25±1 | 625 | 0.0550 | 0.853 | 13.53 | 0.6 | 1.7 | 0.920 |
| 6±0.1 | 25±1 | 833 | 0.0088 | 0.990 | 15.92 | 0.67 | 1.5 | 0.998 |
| 7±0.1 | 25±1 | 769 | 0.0049 | 0.971 | 10.29 | 0.67 | 1.5 | 0.980 |
| 8±0.1 | 25±1 | 769 | 0.0170 | 0.920 | 17.98 | 0.75 | 1.3 | 0.940 |
| 9±0.1 | 25±1 | 833 | 0.0190 | 0.970 | 16.15 | 0.89 | 1.1 | 0.970 |
| Mean |                  | 766 | 0.0190 | 0.930 | 15.10 | 0.692 | 1.5 | 0.965 |
| Standard Deviation (±) | 76 | 0.0180 | 0.055 | 2.76 | 0.013 | 0.2 | 0.029 |
| Coefficient of Variation | 0.099 | 0.980 | 0.059 | 0.183 | 0.163 | 0.1 | 0.030 |

The fact that the observed arsenic concentrations in the study area are inconsequential compared to those reported in other gold mining areas should not be the reason for complacency due to the following reasons:

(i) WHO (2001) asserts that arsenic is carcinogenic even at very low concentrations and the effect is cumulative (even at the WHO guideline of 10 ppb, the health risk is still 0.2 %);

(ii) the population in the Lake Victoria basin of around 4.2 million (about 12.4 % of the population of Tanzania) (URT, 2002), which is potentially exposed to arsenic is high and as such even a small risk could be significant. However, the proportion of the population exposed to arsenic in Tanzania is significantly lower compared to that in Bangladesh where the percentage of population exposed to drinking water with arsenic concentration of more than 10 ppb has been reported by Lokuge et al. (2004) to be about 58.8 %;

(iii) majority of Tanzanians (61 %) and especially in rural areas do not have access to safe water and as such they rely on sources with compromised quality including those containing arsenic and other contaminants. It is worth noting that in the Lake Basin more than 67 % of the households live more than a kilometer from a safe drinking water source (URT, 2002); and,

(iv) the lake Basin is generally dry with high evaporation rates (Sutcliffe and Petersen, 2007) thus there is a possibility of concentrating the contaminant in water sources.

Therefore, if left unattended the arsenic contamination problem could sooner than later reach a crisis level.

Arsenate and arsenite, which are arsenic species of significant public health concern, were detected in the sampled water sources in the study area. Observed mean pH value of samples analysed was 7.58 ± 1.188. This pH value favors the formation of arsenate (specifically HAsO<sub>4</sub>²⁻) over arsenite. In all samples analysed arsenate was a more predominant species than arsenite. Indeed the output values of the model for speciation of arsenic showed that average concentration of arsenate (0.19 nMol/L) was six orders of magnitude higher than that of arsenite (7.2 × 10<sup>-8</sup> nMol/L). The dominance of arsenate over arsenite is basically because pH and redox potential of most aquifers favors the formation of the former over the latter (Bhumble and Keefer, 1994; Smedley and Kinniburgh, 2002). Conversely, arsenate, the thermodynamically stable form, generally predominates in oxic surface waters (high redox potential) and moderate pH range (Bhumble and Keefer, 1994; Smedley and Kinniburgh, 2002). Therefore, the findings of the present study are in agreement with observations made by Smedley and Kinniburgh (2002) and Korte and Fernando (1991) who also observed the predominance of arsenate over arsenite. In addition, low positive Pearson’s correlation coefficient (r = 0.1) also suggests that formation of the two arsenic species is uncorrelated.

The presence and concentration levels of iron in sampled water sources may provide some clues on mobilization of arsenic in the study area. Major source of arsenic in the
study area is suspected to be oxidation of arsenopyrite (FeAsS), which is exposed to the air as a result of gold mining activities (Halsey, 2000). Arsenopyrite oxidation releases arsenite (subsequent oxidation to arsenate may occur), sulphate, and ferrous iron, though some oxidation to ferric iron is evident (Halsey, 2000; Smedley and Kinniburgh, 2002). Analytical results of drinking water samples from the study area showed strong positive Pearson’s correlation between concentrations of arsenic and iron (r = 0.9) thus supporting this assertion.

**Potential effects of phosphorus on adsorption of arsenic**

The importance of phosphate in analysis of arsenic adsorption stems from the fact that it competes with arsenate for adsorption sites. This competition basically originates from their striking resemblance and it is governed by the Law of Mass Action (Smith et al., 2002). Under normal conditions, phosphate will be preferentially adsorbed onto adsorbent sites by virtue of its higher activity coefficient than that of arsenic (Kassenga, 2003).

According to Smith et al. (2002) the competition between arsenate and phosphates is also affected by the content of oxides of iron in the adsorbent. For an adsorbent with low iron content (less than 100 mmol /kg dry weight) the minimum threshold concentration of phosphate in water, which may impair arsenic adsorption is about 0.16 mmol/L (Smith et al., 2002). The average concentration of phosphate in sampled water sources was observed to be 0.15 mmol/L and the mode was 0.1 mmol/L. Since stabilized ferralsols had iron oxide concentration of 1,560 mmol/kg (Table 2), which is significantly higher than 100 mmol/kg, the adsorbent should therefore be capable of adsorbing arsenic with minimum impairment of its removal efficiency by phosphate.

**Effect of pH on adsorption of arsenic**

The supernatant from the control experiment showed that insignificant concentration of arsenic (< 0.01 ppb) was present in either stabilized red soil and/or distilled water; therefore, spiked arsenic trioxide was the only source of arsenic in the reaction vessels.

The influence of pH on arsenic adsorption was found to be insignificant as Figure 2 shows. Significance of the effect of pH on arsenic adsorption was evaluated using Analysis of Variance (ANOVA). Indeed the influence of pH on arsenic adsorption on stabilized ferralsols for all concentrations used during the experiment was found to be insignificant (F_{crit} > F: 2.7728 > 1.6145 and P = 0.2068) as illustrated in Table 4.

The possible reason for this observation could be that the buffer created by the carbonate ion present in the red soil and Portland cement, negates the dependence of initial pH on arsenic adsorption as Pogany and Davies (2001) also observed. The mean removal was 98.07 ± 1.633 % for all pH values and arsenic concentrations used during the experiment. These results suggest that the adsorbent is capable of removing arsenic effectively over a wide range of pH.

### Table 4: ANOVA results.

| Groups | Count | Sum          | Average | Variance |
|--------|-------|--------------|---------|----------|
| pH 4   | 4     | 392.2825     | 98.07062| 5.227077 |
| pH 5   | 4     | 387.6303     | 96.90758| 4.184644 |
| pH 6   | 4     | 392.1557     | 98.03893| 1.56918  |
| pH 7   | 4     | 388.3189     | 97.07972| 1.245141 |
| pH 8   | 4     | 397.1436     | 99.2859 | 0.034876 |
| pH 9   | 4     | 396.1527     | 99.03817| 1.856862 |

| Source of Variation | SS      | df  | MS           | F       | P-value | F crit |
|---------------------|---------|-----|--------------|---------|---------|--------|
| Between Groups      | 18.99449| 5   | 3.798897     | 1.614516| 0.2068  | 2.772853|
| Within Groups       | 42.35334| 18  | 2.352963     |         |         |        |
| Total               | 61.34783| 23  |              |         |         |        |
Adsorption Modeling

Adsorption of arsenic on stabilized ferralsols was described well with both, Freundlich and Langmuir sorption models as illustrated by plots in Figure 2 and 3. However, the Freundlich model consistently gave significantly ($t_{Crit.} < t : 2.156 < 2.571$ and $P = 0.041$) higher coefficient of determination ($R^2 = 0.965 \pm 0.0291$) than Langmuir sorption model ($R^2 = 0.932 \pm 0.0552$) at the concentration range of 1.25 – 20 mg/L as shown in Table 2.

This phenomenon could have been caused by the fact that at a high concentration range Langmuir and Freundlich sorption models behave differently. However, at low solute concentration the sorbed concentration increases linearly with increasing solute concentration for both models. The reason for this behaviour is that Langmuir sorption model assumes that there are a limited number of sorption sites whereas the Freundlich sorption model assumes that the number of sorption sites is large and infinite relative to the number of contaminant molecules (Wiedemeier, 1996).

It can be observed from Table 2 that the Langmuir sorption model gave lower values of $b$ ($0.0190 \pm 0.0180 \ \mu g/g$), equilibrium constant for the sorption reaction, which signify that the value of $Q_{max}$ ($766 \pm 76$), maximum number of sorption site, is very high compared to $q_e$ (sorbed contaminant concentration). This observation is consistent with the fact that the adsorbent had a higher affinity for arsenic as attested by the observed high percent removal values (Figure 2).

Likewise, the Freundlich sorption model yielded higher values of the distribution coefficient, $k$ ($15.1 \pm 2.76$), and the chemical-specific quantity, $n$ ($1.5 \pm 0.2$), which again indicate that the adsorbent (stabilized ferralsols) has a high affinity for arsenic. This implies that within the concentration range used ($0 - 20 \ \text{mg/L}$) arsenic adsorption on stabilized ferralsols was not constrained by the number of sorption sites. In view of this, stabilized ferralsols should be able to effectively remove the contaminant from water sources in the study area since the highest arsenic concentration observed in the sampled water sources was just 70 ppb.

Conclusions

In the present study, arsenic was detected in 58% of water sources sampled and 41% of them had arsenic levels equal to or exceeding the Tanzania Drinking Water Quality Standards threshold value of 10 ppb. Modeling of data has shown that arsenate is more dominant than arsenite, which is the most toxic species. Arsenic mobilization is suspected to be due to oxidation of arsenopyrite after exposure of the ore to the air partly due to mining activities. The arsenic contamination problem in the study area may be aggravated in a near future by high population exposed to the contaminant, lack of reliable and safe water sources and dry weather conditions, which tend to concentrate the contaminant.

Stabilized ferralsols was capable of removing via adsorption up to 99.7% of arsenic regardless of pH and initial concentration. In view of this, adsorption using stabilized ferralsols may be considered to be a promising technology for removing arsenic from drinking water.

Generally, arsenic occurrence in groundwater has a large spatial and temporal variability especially in terms of species and concentration thus making it difficult to draw meaningful conclusion from a handful set of data based on only one sampling effort. It is, therefore, recommended that more comprehensive surveillance studies should be performed for proper quantification of the magnitude of the arsenic contamination problem.

Studies reported in the current paper were conducted under controlled laboratory conditions, which did not mimic actual field conditions reasonably well. In view of this, further studies involving column experiments are recommended to be conducted under actual field operating conditions which could affect adsorption efficiency such as contact time, particle size, competing species, and temperature. It is also worth investigating the possibilities of regeneration of the adsorbent.

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