Assessment of Pollution Potential of Cyanide-Bearing Tailings Dam*

E. K. Asiam and D. Kugblenu


Abstract

This study ascertained the types of chemical species that are formed in tailings material of gold leaching plants and assessed their short and long term pollution potential capabilities. The results revealed that chemical, biochemical and geochemical reactions were occurring in the tailings material; producing complexed cyanides, bicarbonates and nitrates which accumulate with depth in the tailings material while ammonium, carbonate and sulphate ions decreased with depth. Heavy metal accumulation particularly Mn, Pb, Zn, Cu, Fe were also observed to increase with depth. It was concluded that the cyanide-bearing tailings material has short term pollution potential of free cyanides, ammonium ions, sulphates ions and a long term risk of, complexed cyanide, nitrate and increase in water hardness due to bicarbonate ion. Heavy metal pollution and a decrease in pH are also possible.

1 Introduction

Gold normally occurs in association with some amount of waste materials (gangue) and after beneficitation of the ore the gangue which becomes the solid phase of the tailings is deposited in impoundment structures (tailings dam) to maximize space in the beneficiation plant. Therefore, tailings dams are unavoidable structures of any mining operations. During the extraction of gold from quartzite and sulphide ores, cyanide is used as a lixiviant and as a result the tailings will always contain some amount of cyanide.

Cyanide can impact adversely on the environment in three distinct ways namely: producing direct and on-going poisoning through volatilization of hydrogen cyanide gas, interfering with certain metabolic pathways as a result of formation of certain metal thiocyanate complexes and producing toxic levels or deficiencies of certain metal ions in natural water by complexing with them (Clark and Linkson, 1997). Furthermore, if the solid phase of the tailings is not stable (if sulphides are present) then there exists the potential of acid mine drainage, a phenomenon that tends to lower the pH, leach and mobilize heavy metals into the geoenvironment (Marcus, 1997).

Consequently, seepage from such tailings material is generally not of a suitable quality for direct discharge into the environment during and after deposition; as it has the potential to impact adversely on the ecosystem. As such, some attempts are normally made nowadays to detoxify the tailings’ liquid and also line the dam’s surface with imperious materials to contain any liquid and thereby prevent contamination of the geoenvironment. Unfortunately, the liner materials that are used have limited life span and lining is not a suitable option for in-pit tailings impoundment structures because of their irregular configuration (Lane, 2000). In addition, there seems to be lack of adequate data on in-situ transformation of chemical species within the tailings material itself.

As a result of these limitations, liquid contained in tailings is a source of substantial concerns to regulatory agencies since several unplanned releases from tailings dam have occurred around the world such as Los Frailes in Spain in 1998, Aurul in Romania in 2000 and Merriespruit, in South Africa in 1994 among others which have caused extensive destruction to life and property in some cases (Anon, 2002; Trancoso, 1990). Hence, potential chemical impact and contaminant transport from cyanide-bearing tailings material cannot be over emphasised since detoxification and other methods like lining do not ensure total environmental protection.

As gold mining operations nowadays have concentrated on exploiting low grade surface deposits in many countries, large tailings dams are inevitable to store the ensuing tailings.

Hence, there is the need to generate more data on in-situ reactions occurring in tailings material in order to design, operate and decommission tailings facility in a manner that will not only be economical but will also ensure public safety, environmental protection and industry trust.

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This paper therefore attempts to identify the types of chemical species that are formed in tailings dam in a tropical environment and to assess their pollution potential.

2 Location and Climate

The tailings dam on which this study was carried out is about 100 ha and is located in one of the gold mines in Ghana. A tailings material was deposited in this dam for 6 years and thereafter rehabilitated with plant cover. The area is classified as having a humid tropical climate characterized by hot and wet rainy periods and mild temperatures. The mean air temperature is about 26°C and the mean monthly relative humidity increases from 65% in January to a maximum of over 80% in August and September. The mean annual rainfall is about 1800 mm.

3 Methodology

The following were carried out:

3.1 Sample Collection

A mechanically powered auger was used to drill four holes: one located at eastern, western, northern and southern sections of the tailing dam. Each hole was 100 m from the embankment. Samples were taken at various depths as drilling progressed until water was encountered when the drilling was stopped.

3.2 Laboratory Investigations

The moisture content of the samples was determined by drying 5 g each at 105°C for 12 hours, cooled in desiccator and then weighed.

A composite sample was prepared from the samples from the four holes and particle size determination was carried out using sieve analysis. Another composite sample for each hole was also prepared for X-ray diffractometry (Siemens D500) analysis, ore microscopy examination (Leitz Sm-Lux-Pol) and for identification of Nitrosomonas and Nitrobacter bacteria using Vinogradsky (1952) method.

About 1 g of the composite sample of the four holes was digested with aqua regia for one hour and then filtered into a-100 ml flask. The filter cake was washed with distilled water and made to the mark. The filtrate was analyzed for some elements.

About 200 g (wet) of each sample from each hole were pulped to 50% solid (taking into account the moisture content of the sample) with distilled water and leached for 24 hours. The pulp was filtered into a-250 ml flask and made to the mark. The following were carried on the sample of the filtrate: Measurement of some physico-chemical properties, elemental analysis using AAS (Philips 480D) and determination of some anions by ion chromatograph (Dx 2000 isp) were carried out on the sample and filtrate.

4 Results and Discussions

During the drilling of the holes water was encountered at depth of 6.5 m, 4.5 m, 3.3 m and 2.5 m for the northern, eastern, southern and western holes respectively. The results obtained for northern hole were similar to the other holes. Therefore, the results for the northern hole were used in the detailed discussions because of its greater depth compared to the rest of the holes.

The results of the particle size analysis showed that 40% of the particles are sand, 55% silt and 5% clay indicating the tailings material can be classified as silty-sand. Thus, it can be inferred that the tailings material will be permeable to liquid and gases and desiccation of the material will be possible. As a result, the moisture content of the samples was found to increase with depth increasing from 12% to 19% within a-6 m depth.

The X-ray diffractometry analysis revealed that the main mineral was quartz while the minor minerals were pyrite, muscovite and kaolinite. The microscopic examinations further showed that many of the pyrite grains have been altered or corroded. Thus, it can be inferred that the pyrites were being weathered or undergoing geochemical reaction.

Consequently, a relatively high amount of sulphate ions was found in the tailings mass. It can be seen in Fig. 1 that the sulphate ion concentration decreased with depth. The decrease could be due to its conversion to gypsum (CaSO₄·2H₂O) by Ca²⁺ ion (see Equation 1) whose concentration was high in the tailings (see Fig. 2). No gypsum was detected in the X-ray diffractometry analysis as it may be in amorphous state. Sulphate can cause water hardness and also impact taste of water.

\[ \text{Ca}^{2+} + \text{SO}_4^{2-} + 2\text{H}_2\text{O} = \text{CaSO}_4·2\text{H}_2\text{O} \] (1)

Fig. 1 Variation of Sulphate Ion Concentration with Depth
The results of the AAS elemental (total) analysis are given in Table 1, while the water soluble elements are plotted in Fig. 2 and 3. It can be seen that the elements in the solid phase are also present in the interstitial liquid phase of the tailings material. Hence, the elements in the water soluble phase might have been derived from the solid phase of the tailings material. Furthermore, the concentration of the water soluble elements increased with depth. Notably, there is almost 50% increase in Manganese concentration at depth of 5 m, iron was 30% and lead 200%. This accumulation could be due to weathering activity in the tailings material. The relatively high level of Zn, Cu and Fe in the liquid and solid phase of tailings material may give rise to more cyanide effectively being retained in the tailings material through complex ion formation thereby stabilising cyanide that would otherwise have been available for hydrolysis, volatilization and eventual venting to the atmosphere.

Consequently, the total cyanide and the weak acid dissociable (WAD) cyanide levels were found to increase with depth, while free cyanide concentration was almost constant at all depth (see Fig. 4). Thus, it can be inferred that most of the cyanides were either strongly or weakly complexed. The complexed cyanide species are not toxic but may break down in the long term, producing free cyanide which is toxic (Smith and Mudder, 1991).

<table>
<thead>
<tr>
<th>Element</th>
<th>Fe</th>
<th>Mn</th>
<th>Zn</th>
<th>Cu</th>
<th>Pb</th>
<th>Ca</th>
</tr>
</thead>
<tbody>
<tr>
<td>Concentration (ppm)</td>
<td>17806.6</td>
<td>54.3</td>
<td>10.4</td>
<td>16.3</td>
<td>6.8</td>
<td>28.7</td>
</tr>
</tbody>
</table>
The analysis of NH$_4^+$ and NO$_3^-$ is presented in Fig. 5. It can be observed that ammonium concentration decreased while that of nitrate ion increased with depth. The results of microbial identification confirmed the presence of *Nitrosomona* and *Nitrobacter* bacteria in the tailings material. This information suggests that bio-chemical reactions are occurring in the tailings namely: NH$_4^+$ is converted to NO$_2^-$ by *Nitrosomona* bacteria, while the NO$_2^-$ was further converted to NO$_3^-$ by *Nitrobacter*. Nitrate can produce methemoglobinemia in babies (Chanlett, 1977). Ammonium ion can impact odour to water.

The level of CO$_3^{2-}$ and HCO$_3^-$ in the tailings is presented in Fig. 6. It can be seen that while CO$_3^{2-}$ level decreased with depth in contrast, the HCO$_3^-$ level is increased with depth. The pH of the tailings slurry during the time of deposition was greater than 10 and lime was used to control the pH. The carbon dioxide of the atmosphere can react with the excess lime as follows:

$$Ca(OH)_2 + CO_2 = CaCO_3 + H_2O \quad (2)$$

The carbonate can also be converted to bicarbonate through a reaction like;

$$CaCO_3 + CO_2 + H_2O = Ca^{2+} + 2HCO_3^- \quad (3)$$

It is also likely that part of the bicarbonate may come as by-product of cyanide attenuation at pH ≥ 10.

$$CN^- + 2H_2O = NH_3 + HCO_3^- \quad (4)$$

Furthermore, at pH between 6.0 and 6.5 carbonate can be converted to bicarbonate as follows:

$$CO_3^{2-} + H^+ = HCO_3^- \quad (5)$$

The pH of the tailings material is within this range (Table 2). These reactions may explain the formation of more bicarbonate with depth.

The foregoing discussions established that geo-chemical, biochemical and chemical reactions were occurring in the tailings material producing new chemical species. As a result, the physico-chemical analysis (Table 2) showed that the conductivity and TDS in the tailings pore water increased with depth while the pH of the tailings material decreased with depth.

### 4 Conclusions

The study confirmed that the tailings material is a silty-sand and therefore liquid can be mobilized through it. It was also established that chemical, biochemical and geochemical processes were occurring in the tailings material producing chemical species that tend to accumulate with depth. These phenomena have brought about increase in conductivity and total dissolved solid values.

The relatively high level of water soluble Zn, Cu and Fe in liquid and solid (as it weathers) phases of the tailings material may retain cyanide through complex ion formation and thereby delay its attenuation through hydrolysis, volatilization and microbial activity.

The short term pollution potential was identified as free cyanide, ammonium ion, sulphates and carbonates while the long term risks will be complexed cyanides, nitrates, increase in bicarbonates (water hardness) and accumulation of heavy metals like lead, manganese, zinc and lowering of pH. The decommissioning of such tailings material has to deal with these issues.

### References


Authors

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