EFFECTS OF VARIED PROCESS PARAMETERS ON FROTH FLOTATION EFFICIENCY: A CASE STUDY OF ITAKPE IRON ORE

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ABSTRACT
The dire need for Itakpe iron ore concentrates of appreciable iron content meets for smelting operation necessitated this study. Core samples of the ore sourced from Itakpe, Kogi State, Nigeria were prepared for petrological analysis followed by chemical and particle size analyses. Froth flotation was done using different collectors at varying particle sizes and pH values. Characterization studies carried out revealed that Itakpe iron ore is a lean ore assaying 36.18% Fe₂O₃ and contains predominantly quartz, sillimanite, and haematite. Its liberation size lies favourably at 75 µm. Processing the ore by froth flotation yielded appreciable enrichment. Optimal recovery (~92%) was achieved using potassium amyl xanthate (PAX) at pH 11 for fine feed sizes (<125 µm) yielding iron concentrate assaying 67.66% Fe₂O₃. Thus, processing at this set-of-conditions is recommended for the industrial production of more enriched Itakpe iron ore concentrates.

Keywords: Process parameters, Froth flotation, Efficiency, Itakpe iron ore

1. INTRODUCTION
Ores are composed of varieties of minerals, among which the mineral of interest lies. Prior to the processing of these ores, it is pertinent to have a grasp of their mineral entities, morphology, the spatial distribution of mineral constituents, particle size distribution, and other attributes [1]. Therefore, a detailed mineralogical characterization does not only reveals economic minerals but also gangue minerals which are deterrent to the exploitation and processing of the ore to salable product [2]. On-premise of the distinctive physical/physicochemical properties of these minerals, the desired mineral(s) can be liberated through successive comminution and subsequently, separated via methods such as magnetic, gravity, and froth flotation [3]. The nature and surface properties of minerals affect to a great extent their susceptibility to froth flotation. These properties are exploited to facilitate the liberation of valuable minerals from gangue minerals [4]. Froth flotation is a wet separation process that segregates mineral particles in a slurry [5]. It is often referred to as a physicochemical process that employs the use of chemical reagents to alter the surface properties of mineral particles towards selective separation [6]. Consequently, the surfaces of selected mineral particles are made hydrophobic (water repellent) and become attached to air bubbles introduced in the pulp via aeration [7]. These are carried to the froth layer and skimmed off while leaving the hydrophilic (wetted) mineral particles depressed in the pulp. Froth flotation is undoubtedly the most versatile mineral separation technique employed in the mineral industry to treat sulphide minerals (such as Galena, Chalcopyrite, etc.), oxide minerals (such as haematite, cassiterite, etc.) and so on; since the mineralogical constituents of these minerals possess distinct surface properties [8]. The enormous growth of industrialization from the eighteenth century until this day has been ascribed

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to the discovery of metals especially iron and the mineral industry as a whole [9]. As the demand for iron concentrate increases globally, there is a need to assess and develop flotation conditions that would best beneficiate viable deposits towards optimal recovery [10]. Of such deposits is Itakpe, Kogi State, Nigeria; which has a total estimated reserve of about 182.5 million metric tonnes and consists mainly of quartzite with magnetite and hematite [11, 12]. The Itakpe deposit has been developed to supply iron ore concentrates to Ajaokuta steel plant and the Delta steel plant, Aladja. The deposit is being processed using magnetic and gravity methods of mineral processing while the flotation plant remains under-developed [13], thus justifying the need for this research.

According to Wills and Napier-Munn [8], in view of assessing the metallurgical performance/efficiency of a process, concentrate grade and recovery are prominent parameters employed. Although, the economic viability of the process depends not only on these parameters but other factors such as smelter's cost are considered [14]. However, froth flotation is a process system whose efficiency depends on the interplay of certain process parameters. It is therefore pertinent to take into consideration these parameters during froth flotation operations. Such parameters include feed rate, pH, collector type, and particle size; which automatically cause changes in other parts of the system such as percent recovery, flotation rate, and pulp density [15]. Therefore, this research aimed to characterize Itakpe iron ore, investigate the effects of varying collectors, particle sizes, and pH values on froth flotation efficiency and also establish the best process parameters that yield optimal recovery.

2. METHODOLOGY

A representative chart of methodologies employed to achieve the aim of this research is presented in Figure 1.

2.1 Material Sourcing and Preparation

Fifty (50) kg sample of the crude ore was sourced from the deposit located at Itakpe, Kogi State, Nigeria having geological co-ordinates of latitude 7°36'52"N and longitude 6°19'7"E. The sample was crushed using a sledgehammer to 10 mm size and charged into the Denver laboratory jaw crusher (Model: Denver D12) for further reduction to 2 mm. Then, thorough mixing was carried out to obtain a homogenized sample. The homogenized sample was then ground to three particle sizes; 63 µm, 75 µm, and 125 µm. These sizes were selected because they fall within 250 – 45 µm, as documented by Wills and Napier-Munn, which depicts the favourable size range for the flotation of iron ores [8].

Figure 1: Typical process flowchart of the froth flotation of Itakpe iron ore
2.2 Compositional Analysis with Energy Dispersive X-ray Fluorescence (ED-XRF)

Chemical constituents of the ore were appraised using Energy Dispersive X-ray Fluorescence Spectrometer (PANanalytical Minipal 7). Twenty grams (20 g) of the ore sample was finely ground to 100% passing 200 mesh and thereafter thoroughly mixed with cellulose flakes binder in the ratio of 5.0 g sample(s) to 1.0 g binder and pelletized at a pressure of 10-15 tons/square inch in a pelletizing machine. The pelletized samples were stored in a desiccator for analysis. ED-XRF machine was switched on and allowed to warm up for 2 hours. Finally, appropriate programs for the various elements of interest were employed to analyze the sample material(s) for their presence or absence. The result of the analysis was reported in percentage (%) for the concentration of minor and major elements.

2.3 Petrological Analysis

Standard size rock samples were cut from the deposits after which their surfaces were ground using emery paper of grit size 500 µm and 1000 µm successively. The samples were mounted on a slide and viewed using a Leica Petrographic Microscope (Model: EGB 100 DMX) to reveal the ore’s microstructure and inherent mineral constituents.

2.4 Particle Size Analysis

Fractional Sieve Analysis technique was adopted to ascertain the particle size distribution of the ore to determine its liberation size. Set of sieves were properly cleaned to avoid contamination of the mineral sample and arranged in conformity with $\sqrt{2}$ series ranging from 500 – 63 µm [16]. 500 g crude sample was charged onto the upper sieve and the set of sieves were agitated for 30 minutes using an Automated Sieve Shaker (Model: Endecott AS400 control). After agitation, the sieves were separated and the retained on each sieve was weighed and recorded. The sieve fractions were also analyzed using ED-XRF.

Plate 1: Denver Flotation Cell
2.5 Froth Flotation

Laboratory flotation experiments were performed using a Denver Flotation Cell, Model: D-12 (Plate 1), having a capacity of up to 800 ml. 100 g sample of 63 µm fraction was charged into a 1000 ml Pyrex cylinder containing up to 500 ml of water under constant stirring speed (600 rpm) to prepare a slurry of 5% pulp density. The initial pH of the slurry was measured as 8.2 using a pH meter and then adjusted to 9 by adding a pH modifier (Sodium hydroxide). The slurry was agitated for 2 minutes after which 2 drops of corn starch (depressant) were added followed by further agitation for 2 minutes. Then, 2 drops (2 mm/kg) of potassium amyl xanthate (PAX-collector) were added and further agitated for 2 minutes. The slurry was then transferred into the flotation cell. The later processes were repeated for the other collector types; sodium ethyl xanthate (SEX) and oleic acid. The impeller speed was set to 1000 rpm and the slurry was further agitated for 2 minutes before adding 2 drops of methyl isobutyl carbinol (MIBC) into the slurry which serves as the frother; it gets adsorbed on the air-water interface and reduces surface tension to facilitate the formation stable air-bubbles. The total agitation time was 8 minutes per flotation process. The air valve of the flotation cell was then opened to introduce air into the slurry for about 30 seconds. This causes the formation of froth at the top layer of the slurry. The froth was then skimmed into trays after every 1 minute until no froth was formed. The above procedure was repeated at pH values of 10 and 11. After which, other sized samples (75 µm and 125 µm fractions) were also processed accordingly. The products (Froth and depressed) were filtered, dried, and randomly sampled for chemical analysis.

3. RESULTS AND DISCUSSION

3.1 Results

The results obtained are presented in Tables 1-3 and Figures 2-6.

<table>
<thead>
<tr>
<th>Compounds</th>
<th>Al₂O₃</th>
<th>SiO₂</th>
<th>K₂O</th>
<th>CaO</th>
<th>V₂O₅</th>
<th>MnO</th>
<th>Fe₂O₃</th>
<th>CuO</th>
<th>ZnO</th>
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<tr>
<td>% composition</td>
<td>4.20</td>
<td>53.05</td>
<td>0.24</td>
<td>0.559</td>
<td>0.008</td>
<td>0.068</td>
<td>36.18</td>
<td>0.034</td>
<td>0.006</td>
</tr>
</tbody>
</table>

Table 2: Mineralogical Modal Analysis of Petrographic Slide of Itakpe Iron Ore

<table>
<thead>
<tr>
<th>Mineral</th>
<th>No. of Counts</th>
<th>Modal Count</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1st View</td>
<td>2nd View</td>
</tr>
<tr>
<td>Quartz (Q)</td>
<td>43</td>
<td>41</td>
</tr>
<tr>
<td>Sillimanite</td>
<td>10</td>
<td>1</td>
</tr>
<tr>
<td>Haematite and other Opaque minerals (H, Op)</td>
<td>67</td>
<td>65</td>
</tr>
<tr>
<td>Total</td>
<td>120</td>
<td>107</td>
</tr>
</tbody>
</table>

Figure 2: Photomicrographs of crude Itakpe iron ore sample with plane-polarized light

Figure 3: Graph showing the plot of % cumulative weight retained and passing against sieve size for the sieve analysis of Itakpe iron ore
Table 3: Fractional Sieve Size Analysis of Itakpe Iron Ore

<table>
<thead>
<tr>
<th>Sieve Size Range (μm)</th>
<th>Nominal Aperture (μm)</th>
<th>Weight Retained (g)</th>
<th>% Weight Retained</th>
<th>% Cumulative Weight Retained</th>
<th>% Cumulative Weight Passing</th>
<th>% Fe₂O₃</th>
</tr>
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<tr>
<td>+500</td>
<td>500</td>
<td>38.50</td>
<td>7.70</td>
<td>7.70</td>
<td>92.30</td>
<td>35.10</td>
</tr>
<tr>
<td>-500+355</td>
<td>355</td>
<td>106.40</td>
<td>21.28</td>
<td>28.98</td>
<td>71.02</td>
<td>26.34</td>
</tr>
<tr>
<td>-355+250</td>
<td>250</td>
<td>38.35</td>
<td>7.67</td>
<td>36.65</td>
<td>63.35</td>
<td>20.20</td>
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<tr>
<td>-250+180</td>
<td>180</td>
<td>65.50</td>
<td>13.10</td>
<td>49.75</td>
<td>50.25</td>
<td>15.81</td>
</tr>
<tr>
<td>-180+125</td>
<td>125</td>
<td>66.45</td>
<td>13.29</td>
<td>63.04</td>
<td>36.96</td>
<td>12.83</td>
</tr>
<tr>
<td>-125+75</td>
<td>75</td>
<td>59.40</td>
<td>11.88</td>
<td>74.92</td>
<td>25.08</td>
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<tr>
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<td>63</td>
<td>50.90</td>
<td>10.18</td>
<td>85.09</td>
<td>14.91</td>
<td>11.06</td>
</tr>
<tr>
<td>-63</td>
<td>-</td>
<td>74.55</td>
<td>14.91</td>
<td>100</td>
<td>0</td>
<td>20.56</td>
</tr>
</tbody>
</table>

Figure 4: Graphs showing (a) recovery and (b) assay of iron concentrate at varied pH values and particle sizes for the flotation of Itakpe iron using PAX.

Figure 5: Graphs showing (a) recovery and (b) assay of iron concentrate at varying pH values and particle sizes for the flotation of Itakpe iron using SEX.

Figure 6: Graphs showing (a) recovery and (b) assay of iron concentrate at varying pH values and particle sizes for the flotation of Itakpe iron using Oleic acid.
3.2 Discussion

The compositional analysis of crude Itakpe iron ore is presented in Table 1. From the result, it can be deduced that the ore contains predominantly 36.18% Fe₂O₃, 53.05% SiO₂, and 4.20% Al₂O₃ alongside other trace compounds with negligible phosphorus content. It can be said that the mineral of interest (Fe₂O₃) is embedded within a quartz dominated matrix, thus emphasizing the need for comminution to facilitate the complete liberation of mineral entities before processing. Therefore, it can be inferred that Itakpe iron ore is a lean non-acidic haematite rich ore assaying 36.18% Fe₂O₃.

Petrological analysis results of Itakpe iron ore are presented in Figure 2 and Table 2; showing the photomicrographs and mineralogical modal analysis obtained respectively. It was deduced that the ore matrix contains coarsely packed grains of quartz (SiO₂), sillimanite (Al₃SiO₅), haematite (Fe₂O₃)/opaque minerals having a relative abundance of 38%, 3%, and 59% respectively. The ore’s photomicrographs further revealed the presence of specks of haematite (reddish-brown) within the rock which indicates a low degree of weathering that may not be visible with a simple eye observation. These findings tailor accordingly to the result of chemical analysis carried out, thus affirming that the ore is indeed predominated with haematite mineral and quartz as the major associated gangue.

Table 3 and Figure 3 reveal the size distribution of crude sample of Itakpe iron ore and a plot of % cumulative weight retained and passing against sieve size respectively. From Table 3, it can be deduced that the ore particles are well distributed within the size range of 355 – 63 µm. This indicates that mineral particles embodied in the ore are nearly fine-sized which enhances the ore’s amenability to froth flotation. The polynomial curves evident in Figure 3 are mirror images of each other having R squared values (R²) of 0.983 and 0.9776 respectively. These values depict that the data closely fit the regression lines/models with an accuracy of ~ 98%; this value satisfies the standard R squared value of >75% which rates the significance of data for analysis [17]. Chemical analysis of each sieve fractions as shown in Table 3 revealed that the actual liberation size of the ore lies at 75 µm. This obtained particle size lies favourably within the size ranges suitable for effective separation by froth flotation documented in literatures [8, 14, 18].

Figure 4 reveals the result obtained from the froth flotation of Itakpe iron ore using potassium amyl xanthate (PAX) in terms of percentage recovery and assay (%Fe₂O₃) at varying particle sizes and pH values. It can be deduced from Figure 4a that an increment in % recovery occurred as the pH value increases from 9 to 11 for all particle sizes. At pH 11, recoveries were 94.80%, 89.79%, and 91.56% for 63 µm, 75 µm, and 125 µm respectively. This implies that floating the ore using PAX in a more alkaline environment enhances % recovery at fine feed sizes. Figure 4b reveals a comparative analysis of the assay of the crude and concentrates obtained for all set-of-conditions. It can be deduced that the crude has been successfully enriched from 36.18% Fe₂O₃ to a concentrate assaying averagely 65% Fe₂O₃ via froth flotation using PAX. Also, a progressive increment of the concentrate assay was observed for all particle sizes as pH increases from 9 to 11. At pH 11, the assays obtained at feed sizes of 63, 75, and 125 µm were 66.13%, 66.26%, and 67.66% Fe₂O₃ respectively. The assays obtained at this pH value were also considerably high relative to other pH values. Therefore, it can be inferred from these findings that processing of Itakpe iron ore using PAX at fine feed sizes (<125 µm) and pH 11 yields enriched concentrate assaying 67.66 % Fe₂O₃ at a recovery of about 92%.

Figure 5 presents the result of recovery and assay of the mineral of interest (Fe₂O₃) gotten from the froth flotation of Itakpe iron ore using sodium ethyl xanthate as collector at varying particle sizes and pH values. From Figure 5a, a somewhat erratic pattern was observed for recovery as pH increases from 9 to 11. At particle size of 63 and 125 µm, recovery reduces from 89.79 – 70.13% and 77.87 – 74.81% respectively, however, at 75 µm, recovery increases from 49.68 – 85.15 % as the pH value increases from 9 to 11. Figure 5b shows the assay of concentrates obtained for all set-of-conditions. Likewise, significant enrichment of the crude ore from 36.18% Fe₂O₃ to concentrate assaying about 66% Fe₂O₃ was realized when processed with SEX. Also, no significant variation in concentrate assay was observed for all set-of-conditions. Therefore, it can be inferred that processing Itakpe iron ore using SEX is best carried out at 63 µm and pH 9 to yield concentrate assaying about 67.81% Fe₂O₃ at a recovery of 89.79%. From these findings, it is noteworthy that these values fall below those obtained when PAX was utilized, and also
SEX nullifies the suitability of the ore to processing at 125 µm but only at 63 µm. This implies that more energy will be expended to completely comminute the ore to 63 µm compared to 125 µm, thus more cost is incurred which renders the process uneconomical.

Having processed the ore using oleic acid as collector, the obtained result is presented in Figure 6. Figure 6a shows the trend of recovery for different particle sizes as the pH value increases from 9 to 11 while Figure 6b also reveals the assay of concentrates obtained for all set-of-conditions relative to that of the crude. A progressive reduction in % recovery was observed as the pH value increases from 9 to 11 for all particle sizes. This implies that the processing of Itakpe iron ore using oleic acid yields less recovery as the pulp becomes more alkaline. This can be ascribed to the neutralization reaction occurring between the collector and the alkaline pulp which mitigates the ability of the collector to make the mineral of interest more hydrophobic. This phenomenon is very common with acidic collectors which limit their usage in alkaline pulp, rather they are employed in an acidic environment where neutralization effect is negligible [19]. From Figure 6b, it is quite obvious that significant enrichment of the crude ore from 36.18% Fe₂O₃ to concentrate assaying averagely 55% Fe₂O₃ was achieved. Also, a progressive increase in assay was obtained for all particle sizes as the pH value increases from 10 to 11. From these findings, the inverse relationship between percent recovery and concentrate assay can be established. Moreso, processing at pH 9 for all particle sizes yielded optimal recoveries. Therefore, it can be inferred that processing of Itakpe iron ore using oleic acid at pH 9 and particle sizes below 125 µm yields enriched concentrate assaying averagely 57.8% Fe₂O₃ at a recovery of 85%. However, it is noteworthy that these values fall below those obtained for PAX and SEX.

4. CONCLUSION
The suitability of Itakpe iron ore to froth flotation at varying process parameters has been investigated and conclusions drawn include:

i. Significant enrichment was actualized when the ore was processed using PAX, SEX, and oleic acid at varying particle sizes and pH values.

ii. Also, froth flotation efficiency was observed to vary with respect to the collector used such that floating the ore using PAX gave the best result.

iii. Processing the ore using PAX at pH 11 and fine feed size (< 125 µm) was established as the best process condition yielding optimal recovery of about 92% at a concentrate grade of 67.66% Fe₂O₃.

5. REFERENCES


[7]. Glembotskii, V. A., Klassen, V. I., and Plaksin, I. N. Flotation, Primary Sources, New York, 1972


