Nickel laterite concentration through a non-conventional method with surface sulfidization

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Abstract
Nickel ores are found mainly as sulfides and laterites in oxidized ores, such as iron oxides, which are usually “Ni-bearing”. This investigation determined the physical-chemical conditions necessary to increase the tenor and recovery in lateritic deposits, with the implementation of a new technology that allows the increase of the tenor (a process called “up-grading”). The froth flotation is proposed as a concentrating process to increase the Nickel content in the lateritic deposits. By means of sulfidization and direct flotation, specific hydrophobicity of the mineral is achieved, substantially improving the nickel concentrations in the process, with recoveries close to 86%, which, compared with conventional direct and inverse floats without effecting this activation with \( \text{Na}_2\text{S} \), results in recoveries of 70% and 16%, respectively. The reverse flotation also increases the Nickel content with an approximate recovery of 70%; however, the froth flotation, with activation \( \text{Na}_2\text{S} \) is still better.

Keywords: Ni-bearing; laterites; concentration; sulfidization; froth flotation.

1. Introduction

Research around the world showed 70 percent of worldwide nickel deposits are classified as laterites, while 30 percent are sulfides. According to Geology for Investors, nickel laterite deposits occur in the 23.5 parallels of the north and south of the equator. The deposits are present in these areas for their warm, tropical environments, which in turn allow for chemical and mechanical weathering to take place. Meanwhile, some deposits are also found beyond the tropical belt, such as in Oregon, Russia and Canada and with the characteristic that replacing atoms in the other minerals such as iron oxides, which are called “Ni-bearing”. Almost three-quarters of the world nickel resources belong to laterite type...
deposits, but less than half of the primary production nickel comes from this type of deposits. On the other hand, the total production of nickel has increased more than 10 times since 1950. Whereas sulfide-type ores contributed up to 90% of nickel in the world, in 2009 laterite-type ores marked a production higher than 42% and is expected to reach 72% in the coming years [1].

Globally, deposits with high nickel content become scarce. For this reason, the development of technologies that can process many nickel laterites of lesser degree in the concentration process is being proposed in order to eliminate a substantial amount of gangue that participates in the subsequent processes of metallurgical extraction [1,2]. The conventional practices for the increase of the tenor in lateritic deposits are based on a separation of sizes, which metallurgically is known as “up-grading” [3], presuming that in the range of fine sizes there is an increase in the percentage of Nickel due to the same genesis of the deposit. However, this practice presents serious problems in the recovery, which is unsustainable in the long term [4], and strongly favors the selective mining of the deposits and punishing the reserves. Other techniques such as magnetic separation and gravimetric separation have been tested but have not generated acceptable concentrates mainly for pyrometallurgical processes [5,6] and have been finely tested hydrometallurgical extraction processes as "heap leaching" as an alternative to the extraction in poor ores of nickel [7,8]. Unfortunately, though, the selectivity of the leaching reagents seriously compromises this technology.

The froth flotation as a nickel concentrating process from lateritic deposits has not been as successful as the recovery of Nickel from sulfide deposits. However, several investigations have been conducted trying to find alternatives to increase nickel concentration in laterites and oxidized ores [3,9]. The main regions where nickel deposits are found from lateritic soils in the world and where froth flotation research has been carried out to process these minerals are New Caledonia, Australia, Indonesia, South America (Colombia and Brazil), the Philippines, India and Russia [10] lateritic deposits have become a major attraction of economic interest and the gangue, modifying the chemical and electrochemical environment of the system by means of the appropriate selection of the added chemical reagents: collectors, foamers, activators, depressors or pH modifiers (Fig 1.) [3,22–24]. The pulp is a mixture of solid particles containing mineralogical species, water, and dissolved chemical reagents, however, it must meet certain conditions, that is, that the material is properly ground no greater than 295 μm, nor less than 53 μm [9,22,23] for this specific case. Floation reagents are the most important variable since they are responsible for changing the properties of wettability [29]. Oxide minerals have a greater affinity for oxygen-containing reactive chains, and their surface chemistry is largely determined by ion exchange reactions. Physical adsorption plays an important role together with chemical adsorption; the adsorption of the collector in non-sulfuriferous minerals is much less specific than in the case of sulfide minerals [3,15,17,22,24,25].

This investigation studied the flotation of oxide minerals, which contain significant amounts of primary slats, such as clays and iron oxides. In addition, the same valuable minerals are generally very soft forming silts. This can be a problem when carrying out the flotation since it can present high viscosity in the pulp, which is covered with slimes of a mineral “slime coating” on the particles more thicker than another mineral, thereby generating greater collector consumption caused by the indiscriminate adsorption and large areas of mineral surfaces, reducing efficiency by the incorporation of ultrafine particles into the air bubbles and the dilution of the concentrate by means of gangue slats entrained in the foam. In addition, the physical adsorption of sparingly soluble collectors, such as fatty acids [4, 26] is much slower and less efficient in the case of fine particles than in coarser ones.

Likewise, in the froth flotation process for oxides, theories are evaluated to explain the adsorption mechanisms of collectors (cationic and anionic) in non-sulfur mineral surfaces (oxides, silicates, carbonates, phosphates, nitrates, sulfates, and others). In addition to the theory of ionic adsorption, or the formation of hemimicelles, the theory of solubility, and the theory of the formation of ion-molecular complexes [27], the theory of ionic adsorption, or the formation of hemimicelles [26] states that the adsorption of collectors occurs by the electrostatic interaction of the collector ions with the surface of the mineral according to the theory of the electric double layer, and by the association of the hydrocarbon chains of the collector ions by Van der Waals lateral forces, forming hemimicelles.
Table 1.

Works carried out on Nickel flotation from lateritic ores in the world.

<table>
<thead>
<tr>
<th>Country</th>
<th>Conditions and aspects</th>
<th>Ni grade initial (%)</th>
<th>Ni grade final (%)</th>
<th>Recovery (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brazil</td>
<td>Flotation with octanedione dioxin as a collector for three laterite samples [12]</td>
<td>1.2</td>
<td>1.4</td>
<td>63</td>
</tr>
<tr>
<td>New Caledonia</td>
<td>Micro flotation of the fraction 53-74 μm, with cetyl trimethyl ammonium bromide</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>India</td>
<td>Anionic collectors, in addition to sodium silicate [14]</td>
<td>0.6</td>
<td>1.6</td>
<td>60</td>
</tr>
<tr>
<td>India</td>
<td>Cationic collectors and starch to recover Ni from a cyclone overflow of 20 μm [15]</td>
<td>0.8</td>
<td>1.3</td>
<td>86</td>
</tr>
<tr>
<td>India</td>
<td>Pre-concentration of the discharge of a hydrocyclone (coarse fraction) with low amount of Ni. The flotation of this cyclone stream using sulfonate, sodium silicate and pine oil [16]</td>
<td>0.5</td>
<td>1</td>
<td>82</td>
</tr>
<tr>
<td>India</td>
<td>Use of sodium lauryl acetic acid methylamino and sodium salt modified by carboxylic acid [13]</td>
<td>0.5</td>
<td>1</td>
<td>70-80</td>
</tr>
<tr>
<td>India</td>
<td>They used as reagents quinoline hydroxy, dimethylglyoxime, nitroso pyrazolone, 2,2 bipyridal, triethanolamine and sodium oleate [15]</td>
<td>0.5</td>
<td>-</td>
<td>40-60</td>
</tr>
<tr>
<td>Côte d'Ivoire</td>
<td>The segregation of nickel minerals in laterites followed by flotation using sodium silicate as a dispersant [17]</td>
<td>1.2</td>
<td>6-8</td>
<td>35-40</td>
</tr>
<tr>
<td>Australia</td>
<td>Several techniques of preparation of feeding before flotation [18]</td>
<td>1</td>
<td>1.4</td>
<td>43</td>
</tr>
</tbody>
</table>

Source: Adopted from [12-18].

An example is seen in Fig. 1 [15,28,29]. The theory of solubility assumes that the adsorption of collectors in minerals is due to chemical bonds that follow the laws that govern the precipitation of substances of low solubility [30,31]. The theory of the formation of ionomolecular complexes indicates that diverse molecular associations between the two unitary species, ion and neutral molecule, would be formed in solution before adsorption. These associations would be quite simple and would be abundant in different ranges of pH in concentrations lower than the respective critical micellar concentrations (CMC), since at such a point no floats would occur due to collector characteristics. Particularly active at the interface, both solid/liquid and liquid/air, would be the ion-neutral molecule dimers, the so-called ionomolecular complexes [3,23,26].

Due to the oxide compounds present in the study mineral and the little reaction with traditional methods require special treatment, sulfidization activation is carried out with sodium sulfide (Na₂S), followed by flotation using xanthate and others tio-collectors. Sulfidizing agents such as sodium sulfide (Na₂S), sodium sulfhydrate or ammonium sulfide ((NH₄)₂S) are usually added in stages for greater efficiency and control. A common example in this technique is for copper oxide ores, since they do not respond well to traditional concentration methods [17].

The process of sulfidization activation is quite attractive, but in practice, care should be taken not engender two important disadvantages. The first is the dosage of the sulphiding agent: excess produces a depression in both the sulfide minerals and in the oxides, and an insufficient quantity produces a poor recovery. The second disadvantage pertains to the oxide minerals, since they respond differently to sulfidization [17]. Fig. 2(a) shows an outline of the activation process using in aqueous solution, as shown in the eq. (1):

\[ \text{Na}_2\text{S} \rightarrow 2\text{Na}^+ + \text{S}^{2-} \]  

The sulfur ion is adsorbed on metal oxides that include the Ni-bearing and do not compromise the surface alteration of the silicates. The result is a surface sulfide of Fe and Ni mainly, which can be hydrophobicized selectively by a thio-collector. In order to have efficiency in sulfidization activation, the pulp must be conditioned at a neutral to alkaline pH no higher than 10.5 [32], and preferably with caustic soda instead of lime. The variation of the collectors

Figure 1. Schematic representation of the electric double layer in the presence of an anionic collector.
Source:[26].

Figure 2. (a) Activation process by means of Na₂S in aqueous solution. (b) Tio-collector Adsorption.
Source: The Authors.
can range from ethyl and amyl xanthates to the promoters AERO 8474, 8475 and 8649 among others, which are dithiocarbamates, especially the thio-collector reagents that favor chemisorption on metallic bonds, as shown in Fig. 2(b) for the flotation of metal sulphides frequently used in the flotation of metal sulfides with a concentration of 250 g/ton and using Aerofroth 65 as a foaming agent, which is a polyglycol that exhibits strength in the flotation circuits [29].

2. Methodology

2.1. Samples

The mineral with which the froth flotation tests were carried out belongs to a lateritic soil extracted from a deposit located in the northwest of Colombia, in the Córdoba department (Fig. 3). The soil was dried and then ground, taking it to an optimum granulometry for froth flotation tests. All tables and figures occupy the entire width of the column. Use figures and tables of two columns of width only when necessary. The SG of the mineral was 2.475. With the atomic absorption spectrometry (AAS) ICETM 3000, the concentration of each of the analytical elements in the sample was determined. Table 2 shows the results of the chemical analysis by atomic absorption performed on the mineral in percentage by weight. This was in order to observe what amount of material of interest was retained in each of the meshes, which were made with a series of Taylor (USA STANDARD TEST SIEVE) screens in the Rotap RX-29® sieve. Approximately 20 tests were performed in SEM (scanning electron microscopy) type JEOL JSM-5910LV, with a quantitative and qualitative analyzer for the three-dimensional observation of high-resolution images, surface analysis at high magnifications, grain shape, and chemical composition. Fig. 4 shows the microanalysis for the mineral that accompanies the nickel in the lateritic deposit.

![Figure 3. A mineral of interest from the lateritic deposit of Ni. Source: The Authors.](image)

The mineral has an approximate content of 1.5% in the mineral, with a range from 1.2% to 1.7%, with 39% insoluble and $SiO_2/MgO$ ratio of 1.8. For XRD tests, X’pert-pro analytical equipment was carried out with X’pert-pro data collector software. For the measurement of the ore, it was crushed and then 10 grams of mesh 200 were taken to finally be analyzed, resulting in what is shown in Fig. 5 and Table 3.

![Figure 4. Scanning electron microscope (SEM) for the float mineral. Source: The Authors.](image)

Table 2. Chemical analysis by atomic absorption.

<table>
<thead>
<tr>
<th>Mesh</th>
<th>% Fe$_2$O$_3$</th>
<th>% Ni</th>
<th>% MnO</th>
<th>% CaO</th>
<th>% Al$_2$O$_3$</th>
<th>% MgO</th>
<th>Insoluble</th>
</tr>
</thead>
<tbody>
<tr>
<td>-70/+100</td>
<td>12.099</td>
<td>1.697</td>
<td>0.201</td>
<td>0.180</td>
<td>0.689</td>
<td>13.491</td>
<td>43.050</td>
</tr>
<tr>
<td>-100/+170</td>
<td>11.970</td>
<td>1.749</td>
<td>0.186</td>
<td>0.185</td>
<td>0.667</td>
<td>12.304</td>
<td>43.420</td>
</tr>
<tr>
<td>-200/+230</td>
<td>11.928</td>
<td>1.711</td>
<td>0.179</td>
<td>0.183</td>
<td>0.667</td>
<td>13.152</td>
<td>43.650</td>
</tr>
<tr>
<td>-230/+270</td>
<td>12.684</td>
<td>1.779</td>
<td>0.174</td>
<td>0.144</td>
<td>0.647</td>
<td>13.308</td>
<td>41.160</td>
</tr>
<tr>
<td>-270/+325</td>
<td>12.787</td>
<td>1.802</td>
<td>0.175</td>
<td>0.143</td>
<td>0.634</td>
<td>12.393</td>
<td>42.630</td>
</tr>
<tr>
<td>-325</td>
<td>12.588</td>
<td>0.169</td>
<td>0.140</td>
<td>0.140</td>
<td>0.737</td>
<td>13.151</td>
<td>42.220</td>
</tr>
</tbody>
</table>

Source: The Authors.

![Figure 5. DRX for the mineral. Source: The Authors.](image)

Table 3. XRD (X-ray Diffraction) results.

<table>
<thead>
<tr>
<th>Name of the compound</th>
<th>Quantity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quartz</td>
<td>1.2</td>
</tr>
<tr>
<td>Magnesioferrite</td>
<td>55.7</td>
</tr>
<tr>
<td>Lizardite</td>
<td>11.5</td>
</tr>
<tr>
<td>Hematite</td>
<td>17.1</td>
</tr>
<tr>
<td>Liebenbergite</td>
<td>14.4</td>
</tr>
</tbody>
</table>

Source: The Authors.
2.2. Flotation reagents

Three different mechanisms of froth flotation were performed in which each of these was used as a type of reagent [22,23,32–34]. Oxide-type mineral collectors must emphasize better efficiency for the recovery of fine particles and aid in the improvement of the differential flotation of complex oxide ores, having greater selectivity in anionic flotation in relation to silica. The reagents that appear in Table 4 were used [35] in the investigation.

Froth flotation tests were performed under forward and reverse mechanisms. For direct flotation, oleic acid was used as collector at a concentration of 20% solution at 250 g/t, similarly, they floatations with activation were performed by sulfidization with isopropyl xanthate sodium at a concentration of 20% in solution 250 g/t and for the reverse flotation the Aero 825 promoter was used with a concentration of 20% in solution at 250 g/t. Likewise, for the three tests, Aerofroth 65 at 200 g/t was used. Finally analyzing each of the results and given that recoveries for the two cells are similar it was decided to perform laboratory tests with Denver cells.

2.3. Hydrophobicity mineral mechanisms

The first mechanism has been formed by the collector adsorption on the surface of the particle to be floated, as loads are increased. The second mechanism raises generate a laterite ore interface modifying the particle surface, achieving hydrophobizing mineral by thio-collectors. It is also necessary that measurements of contact angle and ζ are carried potential, which helps determine the mechanisms of the mineral hydrophobicity.

Testing contact angle measurement was performed with the goniometer type OCA 15 PLUS computer with software SCA 20 and measurements of ζ potential on the computer model Zeta Sazer nano z90 zenh3690 Malvern. The mineral was used in aqueous solution by varying the pH with sodium hydroxide to the basic area.

2.4. Laboratory process

A methodology is proposed as shown in Fig. 6 based on the processing and analysis of the samples, modifying variables during the different stages of the process. Initially, a quartering of the feed sample is performed along with the chemical and mineralogical characterization for the identification of the useful and gangue species, their associations, and release size. Once the characterization stage is completed, the granulometry of the mineral species useful for their concentration is determined. Subsequently, the appropriate cell was chosen for the flotation process and its volume was determined, the ore is subjected to grinding to take advantage of the optimum nickel mineral release size, without forgetting also the variables involved in the flotation process such as of flotation reagents, reagent dosing, pulp density, aeration and conditioning, pH, water type, temperature, and flotation time, in addition, contact angle measurements were made with different reagents to determine the hydrophobicity of the particles.

Flotation tests are carried out in conventional semi-batch (Denver) and column cells for direct and inverse. First, direct flotation tests were carried out in which the material of interest comes out through the foam and the gangue is depressed and exits through the reactor, followed by inverse flotation tests in which the material that comes out through the foam is the gangue and by the tailings the material of interest, in this case, this process was done to float the greater amount of material that is not of interest. In each of the aforementioned tests, there was a previous analysis of reagents, dosage, pH, conditioning time, pulp density and collection time. The two currents of each of the tests both the concentrate and the tails were duly dried, weighed, and finally analyzed by atomic adsorption in order to determine the recovery that was obtained and, in this way, to determine the representative variable for the process (Fig. 6).

In addition, direct flotation tests were carried out on semi-batch flotation cells (Denver) and column with sulfidization activation using as sulfidizing agent the Na2S making the material behaves as a sulfide material. This was achieved with modifications in dosing variables, pH, and time of

Table 4. Flotation reagents.

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aero 825 Promoter</td>
<td>Traditional oil sulfonates, which must be dispersed in water at 82°C to reduce viscosity and improve handling characteristics [32].</td>
</tr>
<tr>
<td>Aeromine 3030c</td>
<td>It is a cationic promoter, it can be used in acidic or alkaline media. It is associated with micas [32].</td>
</tr>
<tr>
<td>Aerofroth 65</td>
<td>Soluble polyglycol that exhibits strength and persistence in flotation circuits. It is used for floating with difficult foaming obtaining a froth with low reagent consumption [32].</td>
</tr>
<tr>
<td>Aerofroth 70</td>
<td>Low molecular weight alcohol foam when the selectivity is important for the feed that contains a percentage of fines higher than normal [32].</td>
</tr>
<tr>
<td>Starch or Dextrins</td>
<td>They possess strong flocculating properties, but when they are negatively charged by substitution of an anion, we obtain negatively charged macromolecules which are used as selective depressants of talc and siliceous minerals [36].</td>
</tr>
</tbody>
</table>

Source: The Authors.

Figure 6. Flow chart of the methodology implemented for the development of the research work.
Source: The Authors.
collection. These were also collected properly by each of the streams, dried, and analyzed, thus making determining their recovery possible.

3. Results and discussion

3.1. Contact angle and z potential values

Fig. 7(a) shows the measurement of the contact angle of the surface of a quartz particle of the mineral of interest, which gave a very low angle, guarantees that the surface is hydrophilic and can be taken as a value of the reference. The collectors used for the modification of the quartz surface that provided the greatest hydrophobicity were the Aero 825 and the Aeromine 3030C both reactive with a concentration of 20% in solution and pH of 7.

Figs. 7(b) and Fig. 7(c) show the contact angle measurements obtained on the modified quartz surface with the Aero 825 and Aeromine 3030c manifolds, respectively. In Fig. 8, the summary of contact angle measurements on surface modified quartz occurs by immersion for 5, 10, 15, 20, 25, 30 minutes Aero 285, Aeromine 3030c collectors, and others like Aero 845.

According to Fig. 8, it is observed that the efficient conditioning time for the collector to act on the surface of the ore is 20 minutes, reaching a higher value of 82° the Aero 825 promoter. Likewise, the Aeromine 3030c promoter the 30 minutes reaches an angle with a value of 63° and the Genamin TAP 100 at 20 minutes reaches a maximum value of 42 °, which indicates that greater hydrophobicity of the surface is obtained with the Aero 825 collector, which help as a reference for doing the reverse flotation.

With the results obtained from the contact angle measurement, the wettability of the surface was considered with the type of conditioner and the conditioning time. An important reason to determine the type of collector used for this type of flotation.

The result of the mineral z potential is shown in Fig. 9, it is observed that the surface charge of the mineral is negative at any pH used for this measurement, so it would be convenient to use a cationic collector to induce hydrophobicity to the mineral. However, it should be noted that the test established to determine the z potential was not carried out with the pure phase of the mineral; therefore, the oxide in which this metal is found is not individually discriminated.

3.2. Froth flotation tests

Conventional test cell (Denver) and cell column are performed; however, the results were similar in the two cell types as shown in Table 5. The flotation tests for each of the cells were performed under the same conditions.

Table 6 shows the results of the reagents used for the different flotation tests and in Fig. 10 the recovery percentage of each of these reagents.
Table 5.
Flotation tests.

<table>
<thead>
<tr>
<th>pH=3 Recovery (%)</th>
<th>pH=7 Recovery (%)</th>
<th>pH=10 Recovery (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Direct Flotation</strong></td>
<td><strong>Reverse Flotation</strong></td>
<td><strong>Sulfidization Activation</strong></td>
</tr>
<tr>
<td>Denver Cell</td>
<td>15</td>
<td>24</td>
</tr>
<tr>
<td>72</td>
<td>70</td>
<td>59</td>
</tr>
<tr>
<td>69</td>
<td>86</td>
<td>84</td>
</tr>
<tr>
<td><strong>Column Cell</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Direct Flotation</td>
<td>17</td>
<td>30</td>
</tr>
<tr>
<td>Reverse Flotation</td>
<td>71</td>
<td>70</td>
</tr>
<tr>
<td>Sulfidization Activation</td>
<td>68</td>
<td>86</td>
</tr>
</tbody>
</table>

Source: The Authors.

Table 6.
Reagents and type of flotation.

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Direct flotation</th>
<th>Reverse flotation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Oleic acid</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>2 Genamin</td>
<td></td>
<td>x</td>
</tr>
<tr>
<td>3 TAP 100</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4 Aero 845 Promoter</td>
<td></td>
<td>x</td>
</tr>
<tr>
<td>5 Sulfate 869</td>
<td></td>
<td>x</td>
</tr>
<tr>
<td>6 Dodecamin</td>
<td></td>
<td>x</td>
</tr>
<tr>
<td>7 Aero 825 Promoter</td>
<td></td>
<td>x</td>
</tr>
<tr>
<td>8 Xantato</td>
<td></td>
<td>x</td>
</tr>
<tr>
<td>9 sulfidization</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10 Aeromine 3030c</td>
<td></td>
<td>x</td>
</tr>
</tbody>
</table>

Source: The Authors.

3.2.1. Direct flotation

The direct flotation process consists of a separation of hydrophobic minerals (material of interest) from hydrophilic minerals (Gangue or tailings) by air injection, in order to obtain a concentration of mineral of interest in the foam, in Fig. 11 observe the results of a direct flotation process. Tests direct flotation were used as collector oleic acid at 250 g/t, foaming the Aerofroth 65 to 200 g/ton changing acidic pH with sulfuric acid and basic pH with sodium hydroxide, with a variety of pH between 3 and 9. In this test the results reported in the literature [13] it is confirmed that the direct flotation is not efficient for oxides in laterite deposits.

3.2.2. Inverse flotation

The inverse flotation process selectively hydrophobes the gangue, which is collected in the foam, therefore, the mineral of interest is depressed, in this case, it is desired to float the largest amount of sterile ore with the minimum percentage of nickel [37]. Fig. 12 show the results obtained with the Aero 825 promoter, with variation in pH and collection time. Reverse flotation tests 825 Aero promoter was used at a concentration of 20% solution at 250 g/t, the foaming Aerofroth 65 to 200 g/t and the mineral depress with corn starch.

3.2.3. Sulfidization activation

The results of the sulfide flotation are shown in Fig. 13 with variation in two variables such as pH and collection time. Sulfide flotation was accomplished by direct mechanism, in which direct flotation test was performed with pH neutral to alkaline, using isopropyl xanthate collector sodium with an amount of 250 g/ton and foaming the Aerofroth 65.

Fig. 14 shows the recovery results adjusted to the García-Zuñiga model for the flotation process by sulfidization.
activation. For this case in the investigation García-Zuñiga model was performed for a batch operation in a conventional Denver cell, represented by the eq. (2):

\[ R = R_\infty [1 - e^{-kt}] \]  

(2)

where \( K \) represents the kinetic constant of flotation and \( R_\infty \) is the maximum recovery achievable in a real process. Since the flotation constant is 0.12 min\(^{-1}\) and is relatively small, it implies that for short flotation times the recovery in the cell is low.

Noting the results and those obtained by previous authors, can admit that this work progresses to a proposal oriented towards recovery and enrichment (Ni-bearing) in laterites, because today the plants of lateritic nickel have no formal processes concentration of minerals, which is considered as an exception to mining-metallurgical plants, it is necessary then to introduce this concept, because several of the concentration processes have failed and it only remains to work techniques related to surface chemistry including froth flotation and selective agglomerations (floculation with polymers or coagulation with salts).

According to the objectives, an innovative methodology is presented, which has not been used previously in the flotation of laterites, which consists of generating a laterite mineral interface formed by an iron sulfide (FeS), which can be hydrophobicized by means of the use of thiocollectors in intervals of neutral and alkaline pH, which is not reported in the literature, which usually operates at acid pH intervals mainly. Together with the above, the literature [3,18] does not reach to present appreciable enrichment and recovery, and the actions that lead to pre-treatments that motivate the selectivity of the hydrophobicization of the Nickel ores are not clear; The direct flotation by them effected to selectively separate from the complex of lateritic oxides is not entirely clear.

This research had to deal with a change in the conception of mineralogy commonly held in lateritic nickel ores in Colombia. A very determining role for the composition of the minerals of Nickel to the oxides is usually assigned; however, the characterization led to conclude that nickel ores are presented as are silicates, which induces a difficulty in selectivity hydrophobing related to two families of minerals containing nickel (oxides and silicates).

In addition to the above, no minerals are presently nickel, if not iron oxide-type minerals mainly, so it can be deduced that Nickel is mainly in Iron replacements, and possibly Magnesium, which confirms the hypothesis of Ni-bearing. Therefore, not all the nickel of the ore is in the form of simple oxides accompanying the iron oxides as it appears in the literature [11]. Table 3 shows the results of the characterization of nickel associated with silicates, which will prevent a high recovery of nickel under a hydrophobicity strategy of simple oxides associated with iron.

On the other hand, in the reverse flotation where it is intended to hydrophobized silicates (mainly quartz), the hydrophobicization also has the opportunity to act on nickel silicates (Liebenbergite). Sulfidization of simple oxides allows adsorption of thiocolecule-type surfactants, such as those used in the research (Table 6), and in which, in the natural pH flotation scenario, they do not interact with silicates. In this sense, sulfiziding generates a sulfide-metallic surface that masks nickel's oxidized ores, which accounts for the losses of this metal in the tails.

The measured Z potential cannot be used as a tool for the selection of physical-chemical flotation conditions as shown in Fig. 9 since this value does not discriminate individually the oxide in which this metal is found. The use of the Z potential would involve the isolation in the laboratory of nickel ore only, and this process is one of the main difficulties in laterite deposits.

On the other hand, for pyrometallurgical nickel extraction processes, the ratio \( SiO/MgO \) should not be higher than 2, since they are conditions for the melting and efficiency of the process [52], this investigation reports a ratio \( SiO/MgO \) of 1.8 so it is located in the desired range 1.5–2.0. The direct and inverse flotation without activation with a recovery of 16% coincide with the results of the literature previously had Table 1 [3,13]. It is for this reason that pretreatments are
suggested to the ores before the concentration processes. From the above, this research, as well as the literature conclude that the absence of activation decreases recovery and enrichment reasons.

4. Conclusions

Through the sulfidization and direct mechanisms of the Ni-bearing ores, specific hydrophobicity of this type of minerals is achieved by substantially improving the Nickel concentrations in the process, with recoveries close to 86% as shown in Fig. 13(b), which compared with conventional direct and reverse flotation without performing this activation with Fig. 11 and Fig. 12(a) give recoveries of 70% and 16% respectively. The reverse flotation also reaches an increase in the nickel content (Fig. 12(a)) with an approximate recovery of 70%; however, the froth flotation with activation is still better.

The interfacial adsorption mechanisms were evaluated with different types of collectors without pretreatment and no selectivity is presented. Therefore, the flotation theory of oxides for this type of ores is not validated by direct flotation, while by sulfidization activation and the use of thiocollector the mechanism approached the theory of metal sulfide flotation where the role of oxygen has been shown to be very important. Nickel concentrates are obtained using froth flotation, with innovative technologies with pretreatment of nickel ores with sulfide, which does not report results in the literature at present.

The froth flotation for concentration Nickel laterite soils showed better selectivity in processes reverse flotation for processes usually used directly froth flotation reaching approximately 72% recovery (Fig. 12(a)) using as Aeromine 825 promoter, a natural pH of 7.0, using starch as a depressant and Aerofroth 65 as a foaming agent.

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