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# Investigation of arsenic removal from copper concentrates by alkaline sulphur leaching

# Taha Boyraz <sup>1,2</sup>, Tuğba Türk <sup>1</sup>, İbrahim Alp <sup>1,2</sup>

<sup>1</sup> Karadeniz Technical University, Dept. of Mining Engineering, 61080, Trabzon, Türkiye

<sup>2</sup> Ergold Innovation Eng. Srv. Ltd. Co., Trabzon Teknokent, No: 19/B No: 3301, 61080 Trabzon, Türkiye

Corresponding author: tahaboyraz@ktu.edu.tr (Taha Boyraz)

**Abstract:** This study investigated the removal of arsenic from copper concentrates by alkaline sulphur leaching. The first set of mineralogical investigations was carried out as part of the study. It has been assessed that arsenic is present in the ore in the form of the tennantite and the minerals of the fahlerz group. Chemical analyses, SEM-EDS and XRD analyses support these mineralogical findings. As was observed to be associated with Cu-S. Mineralogical tests indicated that the arsenic in the concentrate could not be removed by flotation. Preliminary flotation tests also showed that arsenic could not be selectively separated from the copper concentrate. Na<sub>2</sub>S and NaOH leaching (alkaline sulphur leaching) were carried out to selectively remove arsenic from the flotation concentrate. It was observed that the arsenic contained in the copper concentrate was recovered with high efficiency (>80%) after the alkaline sulphur leaching process. Furthermore, it was observed that arsenic recovery increased further (>90%) when the flotation concentrate was regrinded and alkaline leached under the same conditions. The leaching mechanism was investigated by chemical, ore microscopy, SEM-EDS and XRD analyses. Finally, elemental sulfur (S) precipitation is an advantage of the process investigated in this study, as it is environmentally friendly and has the potential to be used in different areas.

Keywords: copper concentrates, arsenic removal, alkaline sulphur leaching, Na<sub>2</sub>S, NaOH

## 1. Introduction

Arsenic is found in several different minerals in nature. Arsenopyrite (FeAsS) is perhaps the most significant of the group, with orpiment (As<sub>2</sub>S<sub>3</sub>) and realgar (As<sub>2</sub>S<sub>2</sub>) also being notable examples. In addition to these, numerous others occur in complex sulfide minerals, including enargite (Cu<sub>3</sub>AsS<sub>4</sub>), tennantite (Cu<sub>12</sub>As<sub>4</sub>S<sub>13</sub>), and proustite (Ag<sub>3</sub>AsS<sub>3</sub>) (Mandal and Suzuki, 2002). The present study investigates copper–iron–sulfide concentrates characterised by arsenic in enargite (Cu<sub>3</sub>AsS<sub>4</sub>) and tennantite (Cu<sub>12</sub>As<sub>4</sub>S<sub>13</sub>). These minerals are the most commonly occurring copper-bearing minerals, characterised by their relatively high copper content. Nevertheless, it should be noted that a high arsenic presence also characterises this situation. The existence of this problematic element has a notable effect on the quality and fundamental characteristics of the copper product (Bujňáková et al., 2014).

The release of arsenic into the atmosphere represents a significant environmental concern at most copper smelting facilities worldwide, posing a substantial risk to surrounding ecosystems and human health. In the pyrometallurgical industry, metals containing arsenic require heating to approximately 500 °C. The oxygen flow is meticulously regulated. This process leads to the volatilisation of the resulting arsenic trioxide, followed by its recondensation (Espinosa, 1965). Nevertheless, the volatilised arsenic in this process may consequently constitute a further source of pollution. Furthermore, solidification and stabilisation are no longer considered the optimal approach for toxic waste with arsenic content (Leist et al., 2000). Therefore, it is necessary to develop alternative approaches for processing such materials (Türk, 2016).

In the composition of copper ores, arsenic is frequently found to be present within either tennantite  $(Cu_{12}As_4S_{13})$  or enargite  $(Cu_3AsS_4)$ . Analytical findings have revealed that particular copper-arsenic

minerals contain 51.6% and 48.4% copper, respectively, demonstrating a propensity to float similarly to other copper sulphide minerals, reporting to the concentrate. In light of the recent regulatory developments on the disposal of arsenic-based products, smelters are now obligated to exhibit heightened discernment in their approach to the procurement of concentrates. Furthermore, appropriate financial penalties are imposed for excessive concentrates arsenic levels (Long et al., 2012). Developing a cost-effective technique for removing arsenic-bearing minerals earlier in processing minerals is increasingly important.

Some hydrometallurgical approaches are available for the treatment of enargite and tennantite, including ammonia leaching (Gajam and Raghavan, 1983), bioleaching (Takatsugi et al., 2011), and acid leaching (Dutrizac and Morrison, 1984). Nevertheless, the preferred technique for extracting arsenic from these minerals would appear to be alkaline leaching (Curreli et al., 2009; Awe and Sandström, 2010). The extraction of arsenic from concentrates in sodium sulfide solutions using highly alkaline reducing conditions results from the alkaline leaching process. The processes of conversion or transformation of  $Cu_3AsS_4$  or ( $Cu_{12}As_4S_{13}$ ) to  $Cu_2S$  or CuS are shown in equations (1) and (2), respectively.:

$$2Cu_3AsS_4 + 3Na_2S \rightarrow 3Cu_2S + 2Na_3AsS_4 \tag{1}$$

## $Cu_{12}As_4S_{13} + 2Na_2S \rightarrow 5Cu_2S + 2CuS + 4NaAsS_2$ <sup>(2)</sup>

The leaching medium is composed of two primary constituents: sodium sulfide and sodium hydroxide. The application of Na<sub>2</sub>S has been demonstrated to generate a markedly reducing environment, with a potential of approximately -500 mV being observed. This phenomenon is in marked contrast to the effect of NaOH on the pH values, which remain at a level greater than 12.5. The property of NaOH under investigation here has been shown to inhibit the hydrolysis of sulfide ions. The role of sulfide ions in the substitution of arsenic in enargite and tennantite has been proven (Tongamp et al., 2009). A crucial factor that has been demonstrated to significantly influence the efficiency of arsenic dissolution from enargite and tennantite is the impact of temperature. It has been found that an elevated temperature is associated with an enhancement in the efficiency of this process (Curreli et al., 2009).

In the process of pressure oxidation, the leaching of enargite, which occurs in the forms of  $Ca_3(AsO_4)_2$  or  $Fe_3(AsO_4)_2$  arsenic, is removed by dissolution in an alkaline solution (Na<sub>2</sub>S–NaOH). For precipitation reactions to occur, the pH of the solution must be reduced to a level below 2. (Filippou et al., 2007). In contrast, Tongamp et al. (2010) have proposed an alternative process for removing dissolved arsenic from the Cu<sub>3</sub>AsS<sub>4</sub>–NaHS–NaOH leaching system. In this process, arsenic is precipitated as Na<sub>3</sub>AsS<sub>4</sub> without the need for pH adjustment.

This investigation proposes a solution to processes where arsenic cannot be removed from copper ores by flotation technique. Furthermore, mineralogical evaluations have been conducted to elucidate the mechanism by which physicochemical techniques cannot remove arsenic from such ores. The present paper proposes a methodology for treating arsenic-bearing copper concentrate, entailing the precipitation of the arsenic-containing filtrate obtained after selective alkaline leaching.

#### 2. Materials and methods

## 2.1. Materials

The current study was performed on a run-of-mine ore and copper concentrate. The ore samples examined in this study were obtained from the sulphide copper deposit in Yomra, Trabzon. The ore is primarily composed of pyrite and chalcopyrite, accompanied by galena, fahlerz, bornite, and sphalerite. The main gangue mineral in the ore is quartz. The following chemicals were utilised in this study: calcium hydroxide (Ca(OH)<sub>2</sub>) (ACS, Sigma Aldrich, USA), sodium-diisobutyl dithiophosphinate (Aerophine 3418A, CYTEC), sodium-diisobutyl dithiophosphinate (MIBC, CYTEC), sodium sulfide (Na<sub>2</sub>S.9H<sub>2</sub>O) (98% purity, Sigma Aldrich, USA), sodium hydroxide (NaOH) (>98% purity, Sigma Aldrich, USA), potassium iodide (KI) ( $\geq$ 99.5% purity, Merck, Germany), L-ascorbic acid (99% purity, Sigma Aldrich, USA), sulfate analyser kit (Palintest) and AAS standard solutions (Merck, Germany).

#### 2.2. Methods

#### 2.2.1. Sample preparation

A copper concentrate sample containing arsenic was subjected to fractional analysis, employing the standard sample preparation methods. The separation of the sample into six discrete fractions was achieved through the implementation of a sieve analysis technique, in which each fraction was separated using a specific sieve size. The fractions were -20 microns, 20-38 microns, 38-75 microns, 75-150 microns, and +150 microns. Thereafter, the polished sections of each fraction were prepared for mineralogical investigation.

The ore samples used in the flotation tests were dried in an oven. Due to the coarse size of the runof-mine ore, a laboratory jaw crusher was used to reduce the particle size to -2 mm. Half of the crushed sample was retained as a reference. Wet grinding was performed using a laboratory-type rod mill with an internal diameter of 19.5 cm and an internal length of 30 cm. The ore sample was ground in a rod mill for different durations (5, 10, and 15 minutes), and the average particle size was calculated. In order to prevent oxidation of the ore, a grinding operation was carried out prior to each flotation test. Appropriate sampling and reduction techniques were employed to ensure that the flotation concentrate samples utilised in leach tests are reduced to the requisite quantity.

#### 2.2.2. Flotation tests

The present study examined the performance of the traditional Denver D-12 type device by implementing flotation tests. In the experiment conducted, the pH value of the pulp was measured using a pH meter. Before the flotation experiments, the solid ratio of the ore subjected to wet grinding in the rod mill was adjusted to a range of 30-35%. Each reagent used in the flotation tests was transferred to a volumetric flask on a precision balance. The volume of the flasks to which 2.5 g of reagent has been added is 250 ml with distilled water. The reagents were prepared in this manner to achieve a concentration of 1% in 250 ml volumetric flasks. The amount of reagents used in the experiments was calculated in g/t. The conditioning step initiated pH adjustment (using Ca(OH)<sub>2</sub>) as the primary procedure. Subsequently, the flotation process was initiated by adding the collector (sodium-diisobutyl dithiophosphate) and frother (Methyl Isobutyl Carbinol). After completing the flotation tests, filtration was conducted using a vacuum pump. The solid residue was then dried in an oven at 105 °C. The dried materials were weighed on a high-precision scale. Finally, the metal contents of the solid samples were determined, and the flotation efficiencies of these samples were calculated.

#### 2.2.3. Alkaline leaching tests

This study aimed to examine the applicability of alkaline leaching techniques for removing arsenic from the flotation concentrate. Leaching studies were conducted within a jacketed reactor, each with a nominal capacity of 250 millilitres and equipped with vortex preventers affixed to the inner walls. The temperature was controlled through the utilisation of a heating-cooling circulator, whereby water was circulated within the jacket of the reactor. During the leaching experiments, the reactors were sealed by applying suitable covers to prevent evaporation losses from the ambient temperature. The mixing process was conducted using a mechanical mixer configured with torque control and a digital speed monitor. In this study, a Teflon (PTFE)-coated impeller was utilised to ensure the impeller's resilience to leaching conditions, particularly elevated temperatures and alkalinity. The schematic representation of the leaching process is presented in Fig. 1. In the experimental process, the quantity of the initial solution was designated as 125 millilitres, and the solid ratio was established at 20% by weight/volume (w/v). The flotation concentrate sample was introduced into the reactor upon attaining the requisite temperature, triggering the experimental procedure. The mixing rate was constant and set at a speed of 500 rpm. The experiment was conducted at a temperature of 90 °C to ensure the optimal conditions for leaching. The leaching time was held constant at six hours. In addition, the impact of particle size on leaching process efficiency was investigated through the performance of distinct experiments utilising concentrate particle sizes (d<sub>80</sub>) of 118 and 57.8 µm, respectively. At predetermined intervals (15, 30, 60, 120, 240, and 360 minutes) throughout the leaching process, samples were collected (2-3 mL) and subjected to centrifugation at 4100 rpm for three minutes. This procedure facilitated the separation of

solid and liquid phases. The extracted solution was then subjected to a process of dilution prior to undergoing a metal analysis. Following the conclusion of the leaching tests, the leach pulp was processed through a centrifuge at 4100 rpm for eight minutes. The objective of this step was to facilitate the separation of the loaded leach solution. After centrifugation, the transparent solution was transferred to sample storage bottles for further analysis. The solid phase was subjected to a hot purified water wash and oven-dried at 65 °C for 12 hours. The leaching performances were determined by analysis of the metal quantities present in the pregnant solution at the culmination of the leaching process, the metal quantities remaining in the waste material, and the metal quantities in the sample before leaching.



Fig. 1. Schematic diagram of the leaching system

## 2.2.4. Arsenic precipitation experiment

The precipitation of the arsenic from the pregnant leach solution can be achieved by means of a short and simple process. The arsenic concentration was increased to greater than 35 g/L to precipitate the arsenic using elemental sulphur (S) from the loaded alkaline leach solution. Double alkaline extraction experiments were performed to achieve this result under identical conditions, using the same leach solution. The findings of the experimental trials indicated that a leach solution with an arsenic concentration of 40.3 g/L was successfully produced. In this stage of the process, the solid (flotation concentrate) and liquid (arsenical alkaline leaching solution) phases were efficiently separated by solidliquid separation following alkaline leaching. The pregnant leach solution was then conditioned at 90 °C for 20 minutes. The initial 10 minutes of the 20 minutes were used for the mixing stage. Concurrently, with the commencement of the second 10-minute period, elemental sulfur was incorporated into the solution at a concentration equivalent to 1.5 times that of the arsenic. After adding sulfur, the mixture was left undisturbed for 10 minutes. The conditioning phase was followed by initiating the precipitation process, which was carried out by slowly reducing the temperature in a laboratory setting.

## 2.2.5. Characterization

The samples were analysed using laser particle size analysis (Malvern Mastersizer 2000) to determine their size distribution. Microscopic examinations were conducted in the KTU Mining Engineering Department Laboratory using a Nikon Eclipse polarising microscope. In addition, the samples were characterised through scanning electron microscopy (Thermo Scientific Apreo 2S) and energy-dispersive X-ray spectroscopy (EDS; Thermo Scientific Ultradry EDS). X-ray diffraction measurements were conducted using the X'Pert3 Powder Diffractometer (PANanalytical), which is equipped with a

 $\theta/\theta$  goniometer, CuK $\alpha$  radiation (45 kV, 40 mA) and a Pixcel 1D detector. The scanning rate was set at 10 degrees per minute, ranging between 5 and 80 degrees. FE-SEM and XRD analyses were carried out at KTU Central Research Laboratory.

#### 2.2.6. Chemical analysis

Before chemical analysis, the samples were dried in an oven. In order to identify the chemical composition of the samples, acid dissolution was carried out using a microwave digestion instrument (Milestone Ethos Easy). The metal contents of the samples were determined by atomic absorption spectrophotometry (PerkinElmer AAnalyst 400 AA). Arsenic analyses were conducted by integrating an atomic absorption spectrometer (AAS) and a hydride production device. (Perkin Elmer FIAS 100 model). The calibration solutions were prepared using standard solutions with a concentration of 1000 ( $\pm$ 5) mg/L to determine metals with AAS. The standard solutions were prepared in a medium of 2M HCl. The analyses were carried out by calibrating the instrument with a blank solution containing 2M HCl. The standards were prepared in this way. In the calibration process, analyses were performed by providing a correlation coefficient of  $\geq$ 0.999 for each metal. Each sample was analysed using a dualreading method, with two readings obtained from each replicate sample.

Sulfur (S) analysis was carried out using a LECO CS744 carbon and sulfur analyser. In this device, carbon and sulfur are determined in inorganic substances by post-combustion infrared identification technique. Additionally, the analysis of soluble sulfates ( $SO_4^{2-}$ ) in solution was conducted using a UV Spectrophotometer (Palintest 5000). In this method, a BaCl<sub>2</sub> tablet was added to the solutions within the appropriate pH range and mixed for two minutes. The tubes were subsequently subjected to a second mixing, after which a photometer measured the light transmittance (T) value at a wavelength of 520 nm. The sulphate ion concentration was then determined as milligrams per litre (mg/L) from the transmittance (T) value using the conversion table. Sulfate analyses were performed in duplicate, and the mean values of each analysis are presented herein.

#### 3. Results and discussion

#### 3.1. Ore characterisation

The current flotation tests were performed on a run-of-mine ore, with the chemical compositions of the samples presented in Table 1. The X-ray diffraction (XRD) pattern of the ore sample is shown in Fig. 2. As a result of the Rietveld analysis, it was determined that the dominant mineral in the Sample was quartz. However, the ore also contained pyrite, chalcopyrite, barite, sphalerite and tennantite (Fig. 2).

			,					
Element	Unit	Cu	Zn	Pb	Fe	As	Ag (g/ton)	S
Ore	%	2.1	1.6	0.3	21.6	0.21	113	25.3

Table 1. The elemental analysis values of a run-of-mine ore

The SEM image and EDS analysis results of the ore sample are shown in Fig. 3. The major peaks in the EDS analysis have been identified as Si, S, Fe, K, Ca, Ba, Cu, Zn and As. The results of the SEM-EDS data and the chemical and XRD analysis confirm each other. The EDS analysis revealed that the prominent spectral peaks were indicative of Si with quartz, S with pyrite, chalcopyrite, sphalerite, tennantite, Cu with chalcopyrite, tennantite, As with tennantite, Zn with sphalerite and Ba with barite.

Microscopic examination of the fraction above 150 microns revealed that pyrite and chalcopyrite were the dominant minerals. It was found that the liberation was inadequate. The association of chalcopyrite, fahlerz and pyrite was determined in grain with dimensions of 290×150 microns. Different compounds were also observed, including pyrite, chalcopyrite, fahlerz and sphalerite (Fig. 4a-4b). Pyrite with a tectonic cataclastic texture of 55×150 microns was monitored. Fahlerz, chalcopyrite, and pyrite complex particles with a size of 75×55 microns are present in these grains; sphalerite and fahlerz formed separately from chalcopyrite. The fahlerz-chalcopyrite substitution was determined (Fig. 4c). The microscopic examination of the particles between 38-75 microns showed that the liberation was better than the upper sieve fractions. The relative grains of chalcopyrite and pyrite were observed to be lower,

and most of the pyrites were free. However, the substituted fahlerz chalcopyrite textures were not liberated (Fig. 4d). As a result of the examination of the relevant sections, it was found that the most common gangue mineral was quartz (Fig. 4).



Fig. 2. X-ray diffraction pattern of the copper ore



Fig. 3. SEM image (a), EDS spectra, elemental mapping and analysis of copper ore

# 3.2. The flotation behaviour of arsenic and other sulfide minerals

Economic considerations indicate that achieving a separation of minerals containing arsenic at an initial step in processing is favourable. In the flotation process, when the ore particle size ( $d_{80}$ ) values were 148.6 µm, 81.2 µm, 64.8 µm and 53.2 µm, the Cu recoveries were 90.7%, 97.2%, 95.3% and 96.5% respectively (Fig. 5). As illustrated by the results obtained, the ore mineral particle size has a significant impact on the flotation process (Wang et., 2023). Incomplete liberation diminishes the flotation efficiency for particles measuring above 80 µm. It has been observed that minerals, particularly arsenic, zinc (Zn), lead (Pb), iron (Fe), and silver (Ag), present in the ore, act in concert and are retained in the concentrate. Using flotation, a physicochemical technique, it was determined those minerals including As, Zn, Pb, Fe, and Ag remained in the concentrate due to their tendency to associate with Cu (Liu et., 2025). Following a comprehensive mineralogical examination (Section 3.1) and a series of preliminary flotation tests, it was determined that the production of arsenic-free copper concentrate by flotation was



Fig. 4. Microscopic images of copper ore (a: +150 µm, b: +150 µm, c: +75 µm, d: +38 µm)

not a viable option (Fig. 5). Moreover, in the recent past, research into the selective flotation of Asminerals has concentrated on enargite. The primary mechanisms by which to achieve enargite selectivity comprise the implementation of selective flotation reagents (Tapley and Yan, 2003), selective oxidation processes (Guo and Yan, 2005; Li et., 2019; Gan et., 2022), and the modulation of pulp potential (Bruckard et., 2007; Panayotova, 2023). Nevertheless, no commercially viable and universally applicable separation method at the flotation step has been developed. (Plackowski et al., 2012).

As a result of mineralogical studies, the presence of copper minerals such as chalcopyrite and bornite, as well as secondary copper minerals such as covelline and chalcocite, were determined in the ore structure (Section 3.3). Secondary copper minerals dissolve in water to form  $CuSO_4$ , which has been shown to activate the sphalerite found in the ore (Wan et al., 2023). In order to prevent this situation, the objective was to prevent the activation of sphalerite by adding Na<sub>2</sub>S to the medium. For this purpose, flotation tests were carried out by Na<sub>2</sub>S to the rod mill at a different dosage using optimum grain size ( $d_{80}$ : 64.8 µm) and reagent (collector & frother) amounts. It has been established that utilising Na<sub>2</sub>S



Fig. 5. The flotation efficiencies obtained at different ore particle sizes

results in a certain degree of inhibition of the revitalisation process of sphalerite. In these tests conducted for the separation of Cu and Zn minerals, no change in Cu recovery was observed, while a significant decrease in Zn recovery was noted (Fig. 6). Selective oxidation processes have been identified as a viable application for these types of ores (Guo and Yan, 2005; Gan et., 2022). The chalcopyrite and other minerals must be separated for this process to be applied selectively. In this section, it was observed that sphalerite and pyrite could not be wholly separated from chalcopyrite (Fig. 6). Also, the use of Na<sub>2</sub>S did not affect the arsenic in the ore. In light of the findings regarding the flotation technique, it is recommended that alternative processes be investigated for the separation of arsenic minerals from copper concentrate.



Fig. 6. The impact of utilising Na<sub>2</sub>S as a modifier on the flotation performance

## 3.3. Microscopic structure of copper concentrate

A sieve analysis was implemented on the concentrate, calculating an average particle size (d<sub>80</sub>) of 118 microns. The analysis of the collected data indicates that 9.9% of the Sample displays a size above 150 microns, whereas 29% of the Sample exhibits a size below 20 microns. Upon examination of particles measuring 150 microns and above, it was observed that chalcopyrite was the prevalent mineral. Following this, the density of pyrite, bornite, sphalerite, chalcopyrite and chalcopyrite were analysed. The presence of such mineral phases as bornite, sphalerite, pyrite and chalcopyrite was identified, with the latter reaching a size of 300×140 microns. In addition, compositions comprising fahlerz and chalcopyrite are available, with a size of 300×75 microns. The pyrite mineral is typically seen as a granular material within larger crystals. Within this specific range of particle sizes, the texture of the substitution is characterised by a high degree of complexity or multifaceted nature (Fig. 7a). Although the replacement texture is abundant in this grain size range, it was also observed that the number of interlocked grains is significant. Tennantite-tetrahydrate minerals were present in this particle size range. It was also observed that enargite and luzunite were present in the fahlerz structure. (Fig. 7b).

Replacement minerals have been identified in fractions of grains measuring between 75 and 150 microns. It was determined through observation that an extensive quantity of chalcopyrite was present within this specific particle size range. Conversely, only a portion of the sphalerite was observed to be liberated. It has been observed that the sphalerite, fahlerz, bornite and covelin minerals exhibit an interlocking relationship (Fig. 8a). In addition, it has been noted that gangue minerals, mainly quartz, are not commonly found. A thorough analysis determined that these sections primarily contain ore minerals. A microscopic examination of particles measuring between 38 and 75 microns revealed the presence of chalcopyrite, fahlerz and sphalerite. Following a thorough examination of the relevant mineral particle, it was determined that grinding below 20 microns was required for liberation (Fig. 8b). Furthermore, it was observed that compositions of fahlerz and chalcopyrite, interpreted as replacement textures are prevalent occurrences. It is widely acknowledged that releasing these replacement tissues

poses significant challenges. A microscopic evaluation of particles measuring between 20 and 38 microns was conducted. It was observed that free chalcopyrite and pyrite granules were prevalent in this particular section.



Fig. 7. Unliberated pyrite-chalcopyrite-sphalerite-fahlerite-bornite grain (Cp: Chalcopyrite, P: Pyrite, Sp: Sphalerite, B: Bornite, Fh: Fahlerz) (a) Fahlerz-enargite-luzunite-chalcopyrite granules (b)



Fig. 8. Interlocking (bonded) sphalerite, fahler, bornite and covelline minerals (a) Interlocking sphalerite, fahlerite and bornite minerals (b)

Furthermore, the observation of the interlocking of chalcopyrite, fahlerite, sphalerite and pyrite, which had been previously noted in larger grain sizes, was confirmed at this specific particle size. Following comprehensive analysis, it was determined that certain complex-bound grains (bornite, fahlerz, sphalerite, and pyrite) contained disseminated grains of pyrite measuring less than 5 microns in size (Fig. 9a). Microscopic examination of the fraction below 20 microns revealed that the majority of the pyrite and chalcopyrite had been liberated. It was also noted that various free-bornite and fahlerz minerals were present. Despite the prevalence of bound particles, a limited number of chalcopyrite, sphalerite, and chalcopyrite amalgamated particles was detected (Fig. 9b).

A detailed mineralogical examination of the copper concentrate was conducted using a polarised microscope. The mineral composition of the copper concentrate was determined through comprehensive analytical procedures conducted on the fractions obtained (Fig 7-9). The minerals that could be associated with the arsenic in the chemical composition of the concentrate were evaluated using a polarising microscope. As a result of these evaluations, it was understood that the concentrate contained fahlerz group minerals such as tennantite containing arsenic. In the SEM-EDS examinations conducted to verify this situation, the presence of sections containing elevated concentrations of Cu (36.6%-%32.8), As (%13.9-%12.3) and S (%41.4-%43.5) was determined by the point analysis technique (Fig 10). The results obtained in this section are compatible with the XRD analysis results (Fig 2.), determining the mineralogical composition of the run-of-mine ore.



Fig. 9. Free pyrite-chalcopyrite granules and bound fahlerz-chalcopyrite-sphalerite-bornite granules (a) Predominantly free minerals and limited bound chalcopyrite and fahlerz (<20 µm) (b)



Fig. 10. Elemental mapping (a-b) and EDS-point analysis of copper concentrate (75-150 µm) (c-d)

## 3.4. Arsenic removal from copper concentrate by hydrometallurgical process

The investigation revealed that the copper concentrate utilised in this study exhibited high concentrations of arsenic (Table 2). It is acknowledged that copper concentrates with a commercial value must contain a maximum of 0.5% arsenic. As demonstrated in Section 3.1.1, microscopic analysis indicated that the arsenic present in the concentrate could not be removed by flotation. Consequently, Na<sub>2</sub>S and NaOH leaching (alkaline leaching) were utilised to select arsenic from the flotation concentrate, and the leaching test results are addressed in this section. The leaching mechanism of tennantite is shown in equation (2). In the context of the alkali leaching process, the presence of Na<sub>2</sub>S generates a reducing environment with a potential of <-500 mV. On the other hand, NaOH hinders the hydrolysis of arsenic-sulfur ions within the enargite mineral structure, thereby maintaining a pH level above 12.5 (Tongamp et al., 2009; Blanco-Vino et al., 2024). Bujňáková et al. (2014) demonstrated that increased temperature during the alkaline leaching process enhanced arsenic dissolution efficiency.

Furthermore, they ascertained that the maximum arsenic dissolution occurred at 90 °C in atmospheric conditions. Consequently, the application of hot (90 °C) alkaline leaching tests, employing a flotation concentrate containing 17969 ppm (1.8%) arsenic at 200 g/L Na<sub>2</sub>S and 50 g/L NaOH concentrations, resulted in an As content of 4700 ppm (0.47%) in the concentrate, achieving an 82.1% As removal efficiency. While the arsenic in the concentrate was eliminated to a substantial degree, the time-leach efficiency curve in Fig. 11. demonstrates that the other metals remained entirely unleached. The concentrated sample contained 17969 ppm (1.8%) of arsenic. It is hypothesised that the Sample, which is mechanically activated, possesses a reduced particle size, an enhanced specific surface area, and an improved level of reactivity (Welham, 2001).



Fig. 11. Alkaline Leach Efficiency (unground concentrate) at 200 g/L Na<sub>2</sub>S and 50 g/L NaOH concentrations and 90  $^{\circ}$ C

The enhancement of leaching processes in extractive metallurgy when high-energy milling is employed is primarily attributable to increases in specific surface area, decreases in mineral crystalline phase content and reductions in particle size (Baláž, 2000). Subsequently, the material was subjected to milling ( $d_{80}$ : 57.8 µm), and the leaching test was replicated within the original conditions. Following six hours, the arsenic content of the concentrate was found to be 1802 ppm (0.18%), while the arsenic removal efficiency increased to 90% (Fig. 12). It was observed that the chemical analysis results of the copper concentrate before and after leaching and the chemical analysis of the loaded leach solution were in confirmation (Table 2). In addition, it is evaluated that the S passing into the solution after leaching is caused by NaAsS<sub>2</sub> passing into the solution phase due to the reaction (Equation 2) of tennantite and Na<sub>2</sub>S. This mechanism explains the decrease in S percentage in the solid sample after leaching.

It was determined that the minerals forming the concentrate composition did not undergo morphological change after leaching in an intense alkaline environment (Fig. 13). In addition, arsenic is transferred to the solution phase with Na<sub>2</sub>S from most arsenic minerals, particularly tennantite (Cu<sub>12</sub>As<sub>4</sub>S<sub>13</sub>) and enargite (Cu<sub>3</sub>AsS<sub>4</sub>), which are found in the copper concentrate. This situation has also

Table 2. The distribution percentage of concentrate elements when subjected to alkaline leaching is determined through chemical analysis using the atomic absorption spectrophotometer (AAS)

Element	Unit	Cu	Zn	Pb	Fe	As	Ag (ppm)	S
Before Alkaline Leaching	%	13.02	8.41	1.98	22.04	1.80	561	48.1
After Alkaline Leaching	%	12.84	8.15	1.85	22.01	0.18	526	45.6
Leaching Solution	mg/L	0.3	0.6	1.3	0.55	3200	2.1	4850



Fig. 12. Alkaline Leach Efficiency (ground concentrate) at 200 g/L Na<sub>2</sub>S and 50 g/L NaOH concentrations and 90  $^{\circ}$ C



Fig. 13. SEM image of copper concentrate before (a) and after (b) alkaline leach treatment

been verified using SEM-EDS analysis and chemical analysis. When the EDS spectrum was examined, it was found that the arsenic peak in the concentrate prior to alkaline leaching had almost disappeared in the concentrate after leaching (Fig. 14). In addition, EDS elemental analysis results showed that the arsenic content decreased from 1.6% (Fig. 14a) to 0.3% (Fig. 14b) with alkaline treatment. These data confirm the results of atomic absorption spectroscopy (AAS).

Fig. 15 presents the results of the X-ray diffraction analysis of the initial copper concentrate with an arsenic content of 1.80 wt% alongside the leach residues obtained after leaching. The analysis of the obtained leach residues indicates a near-total absence of the distinctive diffraction peaks of tennantite  $(Cu_{12}As_4S_{13})$ . As can be seen from the Rietveld plot of the samples before and after the alkali treatment, the tennantite mineral almost disappeared with the alkali treatment and covellite (CuS) peaks formed instead (Fig. 15). It is evident that these data correspond with a high degree of similarity to the arsenic leaching profile of the sample as illustrated in Fig. 12. In addition, the presence of typical peaks of covellite (CuS) was identified in the leach residue obtained following a six-hour extraction process. These results support the conversion of tennantite ( $Cu_{12}As_4S_{13}$ ) to  $Cu_2S$  in Na<sub>2</sub>S media, as shown by the reaction (Equation 1) above. Tongamp et al. (2009) removed arsenic from the mineral enargite ( $Cu_3AsS_4$ ) using NaHS. Their study also found that the mineral structure changed with alkaline leaching, forming Cu-S mineral. The researchers confirmed this change by employing XRD analysis.



Fig. 14. EDS spectra, elemental mapping and analysis of copper concentrate samples before (a) and after (b) alkaline leaching



Fig. 15. X-ray diffraction pattern of the copper concentrate before (a) and after (b) the alkaline leach treatment

# 3.5. Arsenic precipitation

The study demonstrated that approximately 80% of the arsenic was precipitated from the loaded alkaline leach solution following the arsenic precipitation process, as outlined in the test procedure. The

alkaline solution from the precipitation process, which contains negligible arsenic, can be reused in the leaching stage. The precipitation of arsenic from solutions containing dissolved arsenic from the mineral enargite (Cu<sub>3</sub>AsS<sub>4</sub>) by alkaline NaHS leaching was investigated by Tongamp et al. (2010). This process facilitates the recycling of the alkaline solution and simplifies the management of the arsenic present. The pH and Eh of the media during all performed experiments were kept consistently above 13 and below -700 mV, respectively. Under such conditions of significant alkalinity, the dissolution of arsenic trisulphide (As<sub>2</sub>S<sub>3</sub>) was observed to occur. The Na<sub>3</sub>AsS<sub>4</sub> salt, which is characterised by low solubility (Equation 5), is precipitated as a solid salt as a result of the reduction of added elemental sulfur (Equation 3) and the associated oxidation of thioarsenite (AsS<sub>3</sub><sup>3-</sup>) to thioarsenate (AsS<sub>4</sub><sup>3-</sup>) (Equation 4) (Tongamp et., 2010b). As illustrated in Fig. 16., the process flow chart for the precipitation of arsenic from the solution is as follows.

Sulfur 
$$(S^0) \rightarrow S^2$$
 (sulfur reduction) (3)

$$AsS_{3^{3-}} + S^{2-} \rightarrow AsS_{4^{3-}} (As^{III} \text{ to } As^{V})$$

$$\tag{4}$$

$$3Na + AsS_4^{3-} \rightarrow Na_3AsS_4(\downarrow)$$
(5)



Fig. 16. The process of removing arsenic from the leach solution

## 4. Conclusions

It was assessed that arsenic in the copper concentrate investigated in this study was in the form of enargite mineral and fahlerz group minerals. No more than 0.5% arsenic is permitted in saleable copper concentrate. Due to its mineralogical structure, the arsenic contained in the copper concentrate is not expected to be removed by methods such as flotation. As a result, Na<sub>2</sub>S and NaOH leaching (alkaline leaching) were used to remove arsenic from the copper concentrate selectively, and at the end of the process, the arsenic concentrate. Further analysis suggests the mechanism of selective leaching of As in combination with its mineralisation state. Alkaline sulphur leaching is a promising process for removing arsenic from copper concentrates, offering numerous benefits such as improved environmental performance, increased smelter efficiency, economic benefits, lower operating costs and greater sustainability. It supports the production of higher-quality copper while minimising the risks and challenges associated with arsenic contamination.

The arsenic extracted from the flotation concentrate through alkaline leaching assumes a solution form. Due to the difficulty of storing the leach solution in systems such as tailings dams, recovery of arsenic in solid form from the leach solution is required. Moreover, there has recently been a demand for arsenic as a raw material. Consequently, it is imperative to recover high arsenic concentrations in solution and solid form. A solid precipitate was produced due to the arsenic precipitation test carried out in the present study.

Further research is required to conduct detailed studies on removing arsenic from the pregnant leach solution. Different arsenic precipitation techniques will be investigated, particularly the characterisation of the precipitates formed. In addition, future research is needed to evaluate the arsenic sediments from an environmental and economic perspective.

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