INCREASING GOLD LEACHING EFFICIENCY WITH CHANGE OF SOLUTION RHEOLOGICAL PROPERTIES

**Purpose.** Increasing the efficiency of gold heap leaching by changing the solution rheological properties due to its mechanical activation. Semi-industrial tests were carried out to verify the results of laboratory work.

**Methodology.** First, semi-industrial tests were carried out without activation of the solution (basic technology), and then the solution activation was applied. The ore amount in the stockpile was 600 kg, the gold content in the ore is 1.12 g/t. At the same time, the concentration of cyanide was 400 ppm. The activation degree was from 3 to 8 minutes, the leaching time was 2, 4, 6 hours, the solution temperature varied from 13 to 30 degrees.

**Findings.** According to the results of semi-industrial tests, the dependence of gold content, pH value, and residual concentration of the reagent on the leaching time was obtained using both proposed technology and the basic one. Comparison of the semi-industrial test results related to the basic technology and proposed one (with the solution activation) showed that the maximum gold content in the productive solution with the basic technology is 1.83 mg/l, and with the activation of the solution it is 2.10 mg/l. Hence, there is an increase in the gold content by 14.7 % with the solution activation.

**Originality.** Dependence of gold content in productive solution, pH value, and residual concentration of the reagent in solution on the degree of its activation has been obtained.

**Practical value.** The optimum degree of the treatment solution activation during gold heap leaching is determined, which is from 5 minutes to 7 minutes depending on the leaching time, which increases the gold content in the solution by 14.7 %.

**Keywords:** gold, heap leaching, cyanide concentration, mechanical activation, productive solution, leaching solution

**Introduction.** Mineral raw materials, which have more accurate porosity and permeability providing access of cyanide solutions to the surface of noble metals and diffusion of dissolved cyanide metal complexes into productive solution HL, are most suitable for processing by heap leaching (HL) method [1, 2].

In recent years, in the field of gold mining, work has been carried out to involve ores with a low initial content of useful components in processing. It is the heap leaching technology that is of particular interest for processing such raw materials [3].

The application of this process makes it possible to involve poor and out-of-balance ores in the production, open pit stripping soils, and mature fine tailings of processing plants. However, despite all the advantages, a significant disadvantage of such a leaching process was the low recovery rate, the duration of the leaching process, and the dependence on climatic conditions.

The analysis of the works on improving the efficiency of heap leaching makes it possible to draw the following conclusions:

- to intensify the leaching process, various types of effects on ore (physical, chemical, mechanical, biological ones) are used, most of which are aimed at accelerating the transfer of valuable components from ore to productive solution and improving its quality (maintenance);
- physical and mechanical intensification methods are mainly used when reducing the filtration properties of the rock, therefore, the choice of ore size, the method for laying and irrigation of the heap plays an important role;
- to increase the rate of metal dissolution, various catalysts (Fe³⁺ ions, and so on) are introduced into the solution, which allows reducing the duration of heap leaching.

The works by many scientists are devoted to studying the influence of cyanide concentration on the intensity of metal leaching [4, 5]. The use of higher concentration solutions makes it possible to accelerate the leaching process, but at the same time, the consumption is also increased due to its more intense interaction with some minerals [6].

The authors of [7], analyzing the results of further studies, conclude that for each concentration of NaCN the degree of gold recovery and its concentration in solution grow with increasing the number of irrigations to achieve a total of 40–45 % due to the extraction of gold from ore, while the consumption of sodium cyanide in general increases.

Meanwhile, studies are being carried out on the activation of pulp for intensification of cyanide-free leaching of fine-dispersed gold; in particular, preliminary electrohydraulic activation [8] is applied.

It follows from the work [9] that in the temperature range from 20 to 90 °C with an increase in temperature and duration, the degree of gold leaching by “aqua regia” from untreated and activated samples increases. However, activation of samples leads to an increase in the rate of gold extraction: 98 % of gold are extracted in 20 minutes at 90 °C, while the duration of gold extracting for untreated samples at the same temperature is 90 minutes [9].

To intensify the sorption cyanidation process, oxidation is proposed in two stages: physicochemical oxidation (based on directed photo-electro-chemical effects) and bio-oxidation. Sulfide ores, including gold-pyrite, gold-mined ores, occupy a special place in refractory gold-bearing mineral raw materials (according to experts, the share of these ores is 30 % of the world’s gold reserves in the subsoil). In addition to the high dispersion of gold in such sulfides, the reasons for their technological persistence are also the presence of organic carbon. An analysis of mineral and raw material objects of gold ore deposits showed that ores of the black-shale formation of the stringer-porphry mineralization type with fine-dispersed gold in sulfides and carbonaceous matter in the amount of 3–5 % are of greatest interest.

When processing refractory raw materials with gold nano-droplets, the main causes of the valuable component loss include: 1) methods of mechanical disintegrating of the solid raw material matrix that does not allow grinding the material
with a particle size of less than 0.001 cm and does not provide opening the gold micron inclusions; 2) hydrometallurgical and pyrometallurgical processes occurring at low redox potential, whose level does not provide oxidation of the flouried part of the raw material matrix; 3) absorption of dissolved gold from the liquid phase of the pulp at standard cyanidation with clay minerals and carbonaceous substance activated during ore preparation without additional methods suppressing sorption properties of complexes.

The main problems of extracting ultrafine gold from ores concentrates, and fabricated raw materials using chemical processes are as follows: providing access to the valuable component of the leaching solution due to the formation of a sufficiently developed system of supply micro-cracks and pores in the carrier mineral crystals; the choice of an effective system of oxides and complexing agents that disturb the primary chemical bonds between gold atoms and mineral-forming atoms as well as form more stable bonds with it, compared to those elements to which it is associated initially but in the mineral matrix; providing the sequential and long-term output of gold into liquid phase due to various forms of its location, and accordingly, different conditions of its dissolution and precipitation of part of metal dissolved in the first minutes to minerals-sorbents.

In this case, it is possible to improve the efficiency of ultrafine gold extraction by increasing the activity of oxygen, chlorine and hydrogen-containing complexes in solution (liquid phase). This is primarily related to the fact that the diffusion of the solution's active components deep into the mineral structure plays a decisive role in minerals with increased micro-fracture (capillarity) and porosity (layered micro-structure). Second, disruption of the bonds between the gold and the corresponding elements in the superfine layers of the sorbent minerals depends on the oxidizing capacity of the oxygen-containing free radicals. Thirdly, for example, montmorillonite activated during standard cyanidation, i.e. without additional complexes suppressing its sorption properties, can absorb already dissolved gold from the liquid phase of pulp [9].

Laboratory studies on the effect of activation before metal leaching prove the effectiveness of the proposed unconventional methods for opening the mineral matrix. The increase in gold extraction after the combined sulfide and arsenopyrite oxidation technology was 18%, and the duration of bio-oxidation during the preliminary photo-electrochemical oxidation before leaching was reduced approximately by 2 times [10, 11].

The efficiency of processing refractory gold-bearing raw materials by combined methods of cuvette and heap leaching based on gold extraction. The authors of works [15, 16] propose combination technology to increase the intensity of heap leaching. The essence of the given technology lies in the fact that after ore preparation processing of mineral mass by the solution of leaching reagent and extraction of gold are carried out in two stages: preliminary mineral mass is mixed in cuvettes with hydro-electro-activated starting solution with subsequent injection of cyanide solution into the pulp, and its electro-cavitation treatment. On average, despite the increase in gold recovered for ion exchange media in the amount of 72.3 kg (17%), obtained in experimental production lines, the development of the activation effect is heavy [12, 13].

The well-known cyanide technology by heap leaching (HL) also has several disadvantages: 1) low metal recovery (at the level of 50–80%); 2) relatively long duration of the process (from 30 days to 1–2 years); 3) the need for significant space; 4) negative impact of atmospheric phenomena and complexity of operation under intense climatic conditions; 5) not known types of gold-bearing ores are suitable for cyanidation under HL conditions, and so on. That led to the idea of combining HL and cuvette leaching to take advantage of each other and reduce their disadvantages. However, the technology itself, even in combination, does not solve the problems of gold refractoriness [14].

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As the results of previous studies show, in order to increase the gold extraction during leaching, an increase in the cyanide concentration is necessary, which leads to an increase in the cost of leaching. One way to increase the rate of metal dissolution is to intensify the leaching properties of the solution, that is, intensify its activation. To increase metal extraction without additional costs for leaching, a technique for changing the rheological properties of a working solution based on mechanical activation was proposed [17].

Activation of the leaching solution means the increase in the solution’s chemical activity after its treatment in a cavitation plant. Due to the leaching solution activation, the transfer of both the useful component and the colmatage elements into the liquid phase increases, which reduces the likelihood of their precipitation in the form of colmatants. Cavitation in the liquid is accompanied by various chemical reactions. While some chemical reactions are accelerated, others are initiated. These reactions occur in an extremely short time [18].

Cavitation is the formation of fluid discontinuities as a result of local pressure reduction. The essence of cavitation tech-
nology lies in the fact that leaching solution is cavitated before supply to ore mass. As a result of the mechanical effect (cavitation) on continuum, its structure and temperature change, which is accompanied by the breakdown of bonds between atoms and the destruction of the crystal lattice.

**Technique.** To determine the effect of the mechanical activation process on the efficiency of heap leaching, laboratory studies were carried out. Relationships between useful component content in the productive solution, the temperature of the product solution, the reaction time, the concentration of reactants in the leaching solution were obtained.

To prepare working reagents, 5 g of potassium iodide is dissolved in 100 ml of distilled water. The obtained indicator is poured into a drip with a strained plug. 1.733 g of silver nitrate is dissolved in distilled water, poured into a measuring flask for 1000 ml, and brought to the mark with water. With a 10 ml pipette, 10 ml of productive or working solution is taken, poured into a 100 ml conical flask, or into a 50 ml beaker. Then 3–4 drops of potassium iodide solution are added. We titrate with a silver bath until the appearance of persistent yellowish turbidity.

A generator unit was prepared for the mechanical activation of the solution (Fig. 1). The generator is mounted on a platform, where a generator unit with inlet and outlet branch pipes and an electric motor are coaxially placed. Start and stop are performed by the activator control unit, to which the electric network is connected.

To establish the operability of the laboratory plant created for mechanical activation, test activations of liquids were carried out as follows: a working liquid was poured into the reservoir and an electric pump was turned on, which circulated the liquid through the generator along the MNC circuit. A change in the rheological properties of the liquids (solutions) used was recorded by measuring the mechanic-chemical parameters at certain intervals. The solution passing under pressure through the activator and returning to the container over time denatured rheological properties. There was no flow of solution through the docking units of the plant, which confirmed the tightness of the closed circuit.

When testing the laboratory installation, we detected that, when the solution drained after the experiments, part of it remained in the vertical pipe and was not removed due to the presence of a reverse valve. This is caused by design features of the used pump. Based on this experimental result, an electric pump with a drain plug was used. This allowed draining the next volume of cyanide solution into a separate tank complete after its treatment and washing the plant with water to prevent corrosion of the pump parts.

To perform treatment of the following volumes, a minimum of 1.5 liters of solution with the required concentration was poured into the container. It was determined that the preliminary stirring time was within 7–10 minutes to reach the temperature.

Laboratory work was carried out with basic technology as well, i.e. without activation of the solution. During the testing the concentration of cyanide was 400 ppm (a decrease in the cyanide concentration in the leaching solution with an increase in the activation degree is explained by an increase in the solution temperature). The activation rate altered from 3 to 8 minutes, the leaching time was 2, 4, 6 hours, the temperature of the cyanide varied from 13 to 30 degrees.

**Research results.** The laboratory results are summarized in Table 1 to select the optimal degree of the working solution activation.

As Table 1 shows, with an increase in activation extent from 3 to 7 minutes and at leaching time 2 hours the gold content in productive solution increases from 0.49 up to 0.64 mg/l. Further increase in concentration degree, on the contrary, leads to a decrease in the content of gold in productive solution up to 0.6 mg/l. The same change in the gold content of the production solution was observed at a reaction time of 4 and 6 hours. From the laboratory research it can be concluded that the optimal degree of the working solution activation ranges from 5 to 7 minutes depending on the leaching time.

Data processing of laboratory works gave dependences of gold content in productive solution on extent of leaching solution activation and leaching time (Fig. 2).

| Cavitation time, minutes | Gold content, mg/l | pH | t, °C | CN, ppm |
|--------------------------|--------------------|----|------|---------|
|                          |                    |    |      |         |
| with a leaching time of 2 hours |
| Without activation       | 0.42               | 10.10 | 17.0 | 400     |
| 3                        | 0.49               | 10.09 | 20.2 | 350     |
| 5                        | 0.57               | 10.06 | 26.3 | 310     |
| 7                        | 0.64               | 10.02 | 27.5 | 285     |
| 8                        | 0.60               | 10.01 | 30.8 | 260     |
| with a leaching time of 4 hours |
| Without activation       | 0.56               | 10.12 | 17.0 | 400     |
| 3                        | 0.70               | 10.08 | 20.2 | 365     |
| 5                        | 0.79               | 10.05 | 26.3 | 325     |
| 7                        | 0.79               | 10.02 | 27.5 | 305     |
| 8                        | 0.80               | 10.00 | 30.8 | 280     |
| with a leaching time of 6 hours |
| Without activation       | 0.70               | 10.09 | 13.0 | 400     |
| 3                        | 0.72               | 10.08 | 15.0 | 340     |
| 5                        | 0.80               | 10.06 | 21.4 | 300     |
| 7                        | 0.76               | 10.05 | 23.5 | 280     |
| 8                        | 0.75               | 10.04 | 26.9 | 250     |

**Fig. 1. Solution activation generator**

**Fig. 2. Dependence of gold content in the productive solution on the degree of the alkaline solution activation and leaching time**
To verify the results of laboratory work, semi-industrial tests were carried out. Sampling was performed from freed ore in a quarry by an exhaustion method. There was about 1200 kg of ore in total.

Two technological types represented the ores: primary gold-sulfide-quartz and oxidized. The results of group sample analyses showed that the concentration of harmful impurities is low in gold-bearing ores. The gold content of the ore is equal to 1.12 g/t.

For semi-industrial testing, a heap leach unit made of iron sheets was manufactured, which consists of a tray for the productive solution, the base of the plant, a container for the source solution, the main pipeline, and irrigation pipelines. The unit was installed in the laboratory.

To imitate the muck pile, a polyethylene film was glazed onto the plant. Semi-industrial tests were first carried out without solution activity (basic technology), then it was done with solution activity. The amount of ore in the pile was 600 kg, the average gold content in the ore was 1.12 g/t, the cyanide concentration was 600 ppm, and activation degree was 5 minutes.

The leaching time was 20 hours; the first sample was taken after 6 hours, further samples were taken every 2 hours.

Samples of the production solution after the tests were sent to the ore mine laboratory to determine the gold content, pH value, and residual reagent concentration.

Results of semi-industrial tests without activation of solution are given in Table 2.

As can be seen from Table 2, the gold content in the production solution increases from 0.3 to 1.83 mg/l with an increase in the leaching time from 6 to 16 hours. But the gold content in the production solution decreases to 1.43 mg/l with a further increase in the leaching time up to 20 hours. At the same time, the pH value is slightly reduced, and the residual concentration of the reagent decreases from 600 to 80 ppm.

Processing of semi-industrial testing data resulted in the dependence of gold content, pH value, and residual reagent concentration on leaching time at basic technology (Fig. 3).

The results of semi-industrial tests with solution activation are given in Table 3.

As can be seen from Tables 1, 3, with an increase in leaching time from 6 hours to 16 hours, there is an increase in gold content in the production plant from 0.35 to 2.10 mg/l. A further increase in the leaching time resulted in a reduction in the gold content of the production solution up to 1.65 mg/L, with the residual reagent concentration decreasing from 600 ppm in the initial solution to 120 ppm at the end of the leaching.


data (Fig. 4).

Conclusions.

1. Mechanical activation increases the activity of the working solution and reduces the time of crevice.

2. Dependence of gold content in productive solution on the degree of leaching solution activation and leaching time has been obtained. In particular, with an increase in the activation degree from 3 to 7 minutes and with a leaching time of 2 hours, the gold content in the production solution alters from 0.49 to 0.64 mg/l. Further increase in the concentration degree vice versa leads to the gold content reduction in the production solution up to 0.6 mg/l.

3. The maximum gold content of the product solution in the basic technology is 1.83, and it is 2.10 mg/l when the solution is activated, i.e., there is an increase in the gold content when the solution is activated by 14.7 %.

4. The residual concentration of the reagent at the end of the semi-industrial tests in the production plant with the basic technology is 80, and it is 120 ppm when the solution is activated, i.e., when the solution is activated, the gold content in the solution is higher at lower specific consumption of the reagent.

Results of semi-industrial tests with solution activation

| Leaching time, hour | Gold content, mg/l | Residual reagent concentration, ppm | pH value |
|---------------------|--------------------|------------------------------------|----------|
| 6                   | 0.3                | 410                                | 10.1     |
| 8                   | 0.59               | 310                                | 10.1     |
| 10                  | 0.82               | 260                                | 10.2     |
| 12                  | 1.34               | 215                                | 10.2     |
| 14                  | 1.71               | 170                                | 10.2     |
| 16                  | 1.83               | 135                                | 9.7      |
| 18                  | 1.57               | 100                                | 9.30     |
| 20                  | 1.43               | 80                                 | 9.1      |

Fig. 3. Dependencies of gold content, pH value, and residual reagent concentration on leaching time at basic technology

Fig. 4. Dependencies of gold content, pH value, and residual reagent concentration on leaching time when the solution is activated
Підвищення ефективності вилуговування золота зі зміною реологічних властивостей розчину

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**Мета.** Підвищення ефективності купного вилуговування золота шляхом зміни реологічних властивостей розчину за рахунок його механічної активації. Для перегляді результатів лабораторних робіт були проведени на півпромислові випробування.

**Методика.** Напівпромислові випробування спочатку були проведени без активації розчину (базова технологія), потім з активацією розчину. Кількість руди в навалі становила 600 кг, вміст золота в руді становить 1,12 г/т. При цьому концентрація ціаніду становила 400 ppm, ступінь активації від 3 до 8 хвилин, час вилуговування становив 2, 4, 6 годин.

**Результати.** За результатами напівпромислових випробувань отримані залежності вмісту золота, величині рН і залишковій концентрації реагенту від часу вилуговування за базовою та пропонований технології (з активацією розчину).

**Протягом результатів лабораторних робіт були проведени на півпромислові випробування.**

**Ключові слова:** золото, купне вилуговування, концентрація ціаніду, механічна активація, продуктивний розчин, вилуговувач розчин

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