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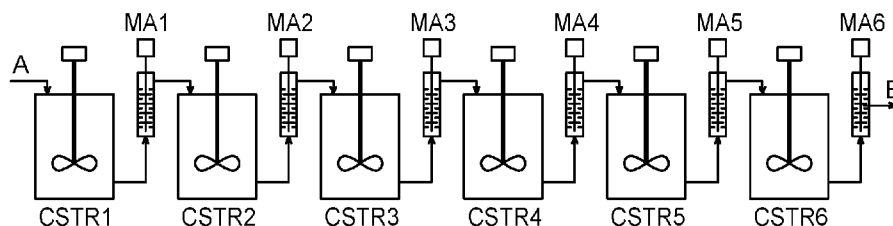
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Fig. 2



(57) Abstract: Alkaline oxidation process for treating refractory sulfide ore or concentrate particles enriched in a metal to be recovered comprising stages in which refractory ore or concentrate particles are surface-oxidized in an alkaline oxidation step in alkaline liquid phase with calcium hydroxide forming an alkaline slurry, which slurry is thereafter mechanically activated to remove passivating coatings from the surface oxidized refractory ore particles.

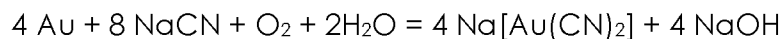


**ALKALINE OXIDATION PROCESS AND DEVICE FOR TREATING REFRACTORY
SULFIDE ORE, IN PARTICULAR REFRACTORY GOLD ORE**

The present invention relates to an alkaline oxidation process for treating refractory sulfide ore particles enriched in a metal to be recovered, i.e., prior to extraction and recovery of such metal, as well as to a device for such alkaline oxidation process.

Gold and other precious metals as well as some base metals are extracted from sulfide ores by treatment with a cyanide solution which solubilizes the gold and said other metals in the presence of oxygen.

In the cyanide leaching process, solid metallic gold (Au^0) is oxidized (to Au^+) by sparged oxygen and is maintained in soluble form as a gold-cyanide complex ($Au(CN)_2^-$) via the overall reaction illustrated below using sodium cyanide as an example, prior to recovery by adsorption onto activated carbon and subsequent processing:

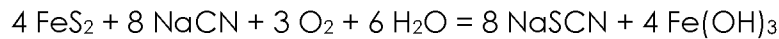


Cyanidation leaching of gold needs to occur at alkaline pH conditions, preferably at pH 11, to maintain cyanide in an anionic solution form (CN^-) for use in the gold dissolution. At pH levels below 11, and particularly below 9, soluble cyanide is converted to hydrogen cyanide (HCN) that is both ineffective as a leaching agent and prone to evaporation as a toxic gas released from the aqueous phase. It is, therefore, imperative to increase the aqueous mineral slurry pH and maintain it at a safe and effective level, which imposes a reagent consumption demand, most often in the form of quicklime (with active reagent CaO) as the most economical alkalizing reagent. Quicklime is converted to its hydrated form, $Ca(OH)_2$, when in contact with water and may be utilized in either $Ca(OH)_2$ or CaO formats.

In some ores, the gold or other metal is termed "refractory" to recovery by cyanidation. There are several features that may cause ores to be refractory: (1) The presence of carbonaceous matter that adsorbs (i.e. robs) gold from the pregnant solution (containing cyanide-dissolved gold) thereby reducing recovery, leading to the term "preg-robbing" ore. The most common

form of refractory ore is caused by the presence of sulfide minerals, such as pyrite or arsenopyrite. Sulfide refractory ore (or concentrate) is the most commonly encountered refractory ore where fine gold particles are co-contained and often occluded within a matrix of sulfide minerals, most commonly pyrite (FeS₂) or arsenopyrite (FeAsS).

Such gold is not sufficiently liberated and therefore not sufficiently accessible to lixiviants, of which cyanide is the most commonly used. Even if sufficiently liberated the presence of sulfide minerals contributes to uneconomically high cyanide reaction consumption because of the overwhelming unwanted side-reaction of sulfide minerals with cyanide, to form thiocyanate (SCN⁻) that is less effective as a gold lixiviant.



Because of the predominance of sulfide refractory ore, the term "refractory" in the rest of this text is meant to refer to sulfide refractory ore (or concentrate) rather than preg-robbing ore.

To solve the aforementioned difficulties, there are several well-known processing options, as pre-treatment before cyanidation, which are available to recover gold from sulfide-refractory ore, most commonly after it has been upgraded to a concentrate via a mineral flotation process [Aylmore et al., 2012. Evaluating process options for treating some refractory ores DOI: 10.13140/2.1.4325.9842 Conference: ALTA 2012 INTERNATIONAL GOLD CONFERENCE, Perth, Australia].

Such available pre-treatment process options are further described in La Brooy et al., Review of gold extraction from ores, Minerals Engineering, [Volume 7, Issue 10, 1994, Pages 1213-1241, [https://doi.org/10.1016/0892-6875\(94\)90114-7](https://doi.org/10.1016/0892-6875(94)90114-7)], and are carried out either via roasting or hydrometallurgical methods that range from high temperature (up to 250°C), pressure oxidation in autoclaves to bacteria catalyzed oxidation at atmospheric pressure and mild temperature conditions (35 – 45°C in the case of mesophilic bacteria and up to 80°C for archaea systems).

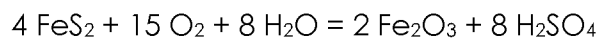
A first option available is a roasting pre-treatment. Roasting is conducted at temperatures of approximately 800°C in the presence of oxygen-containing gas, resulting in the oxidation of sulfides into sulfate solids and gaseous

sulfur dioxide. The generation of sulfur dioxide gas is problematic and needs to be collected, generally by conversion into sulfuric acid. The roasting process and capital requirements are further complicated when arsenic is present in the refractory ore. After calcination, the residual gold-containing material, can be
5 subjected to conventional gold cyanidation processes.

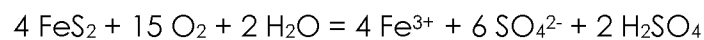
Another option available is pressure oxidation. Pressure oxidation is a hydrometallurgical processing method in which refractory concentrate (typically with a P_{80} 150 μm) is oxidized in an autoclave by the introduction of oxygen gas into aqueous media under elevated pressure and temperature
10 conditions, typically 200°C and 3,100 kPa for up to 60 minutes.

The notation P_x represents a diameter expressed in μm , compared to which the size of $X\%$ by volume of the particles measured are less than or equal to this diameter. For example, P_{80} 150 μm means that 80% by volume of the particles have a diameter smaller than or equal to 150 μm . The particle size
15 distribution of a sample can be determined by different methods, known in the art, for example by laser diffraction according to the standard ISO 13320-2020.

The reaction below illustrates pyrite oxidation (by way of example) as is typically achieved in an autoclave, generating hematite and sulfuric acid.



A portion of the iron contained in pyrite remains in solution as ferric iron sulfate because of the high acid levels preventing all ferric iron precipitating.



A further available option is bacterial bio-oxidation. Bacterial bio-oxidation is a hydrometallurgical processing method in which refractory concentrate (typically $P_{80} < 150 \mu\text{m}$) is oxidized in aqueous media at atmospheric pressure and mild temperature 35 – 45°C, where sulfide mineral oxidation is catalyzed by iron- and sulfur-oxidizing bacteria in a cascade of reactors with a total residence time of approximately 4 days. While bacteria cope well with
25 impurities such as arsenic, they have a low tolerance for chloride (above approximately 1g/dm³), thereby prohibiting the use of seawater (chloride content of approximately 17g/dm³). Similar pyrite oxidation reactions occur
30

during biooxidation as earlier illustrated for pressure oxidation, but the ambient pressure conditions of biooxidation result in iron precipitating as jarosite rather than hematite.

Alternatively, there also exist ultra-fine grinding methods (P_{80} 25 μm),
5 such as the Albion process, disclosed in EP 1 171 641. According to this document, ultra-fine grinding methods are employed to improve the leaching rate under atmospheric conditions in the presence of oxygen, but at the expense of increased grinding costs. In this method, all of the material to be treated is subjected to grinding to reduce mineral particle size, prior to leaching in alkaline
10 conditions. Unfortunately, this process is energy intensive. As disclosed in EP 1 171 641, the alkaline leaching of refractory sulfide and/or carbonaceous materials can only be successfully achieved by careful selection of the particle size of the material to be leached. A similar method with previous ultra-fine grinding of the refractory ore is disclosed in WO2006/064350.

15 A more recent alternative is an acidic leaching process by FLSmidt called the Rapid Ore Leaching process (as disclosed in WO 2016/100981) which makes use of alternating grinding (stirred media reactors) and leaching, in acid environments for base metal recovery.

Leaching inside a mill was also proposed in Patent CN1228480
20 which teaches a process in which milling and oxidation occur simultaneously in the same reactor without separation between the oxidation step and the surface attritioning/milling. The problem with such an approach is that the oxidation and milling cannot be independently controlled.

Several of the above hydrometallurgical methods, such as
25 FLSmidt's Rapid Ore Leaching process, will yield acidic conditions in the bulk of the slurry.

Acid generation is a consequence of most aqueous pyrite oxidation processes and has three important negative consequences:

1. The combination of acidic and oxidative process conditions
30 (dissolved oxygen and ferric iron) is particularly corrosive, thus requiring special materials of construction that, in turn, impose significant capital equipment costs for the pressurized reactor vessels. The corrosive conditions are further

exacerbated by the use of chloride-containing water as is contained in seawater. These features increase capital equipment complexity and costs.

2. The generation of acid during hydrometallurgical sulfide mineral oxidation also causes significant dissolution of gangue minerals contained in the ore or concentrate. Dissolved gangue elements such as Fe, Al, Mn, Mg and others, require precipitation for removal from solution prior to process solution re-use or disposal. These gangue element precipitation reactions also typically require a combination of calcium carbonate (effective at pH levels < 4.5 for Fe and Al removal) and lime (effective at pH levels up to 12 for precipitation of elements that precipitate as hydroxides at elevated pH levels). The precipitation reagents required, incur significant costs.

3. Acidic conditions are contrary to the alkaline conditions required in the subsequent gold cyanidation leaching process. The oxidized slurry therefore requires the addition of large quantities of base reagents to render the pH conducive to cyanidation leaching of gold.

As it can be seen from industrial practice, the available hydrometallurgical process options present problems of acid generation. To overcome these challenges, alkaline sulfide mineral oxidation methods have been proposed, where the absence of acid would avoid the problems of corrosion and gangue mineral dissolution [Bahkta et al., 1989. Alkaline oxidative leaching of gold-bearing arsenopyrite ores. Report of investigations (United States of America Bureau of Mines): 9258. Supt. Docs no: I28.23.:9258]. Alkaline reagents that have previously been considered for this purpose have included, amongst others, NaOH, Na₂CO₃, NaOCl and KOH. Unfortunately, the use of these reagents, however, poses a number of problems:

1. The monovalent cations (such as Na⁺) associated with these reagents typically remain soluble and are not readily removed from solution circuits by precipitation reactions. This, in turn, causes secondary environmental, regulatory or license-to-operate issues because of process circuit discharge and seepage into underground- or open water sources.

2. These reagents are relatively expensive.

Further, Bidari et al. studied the alkaline oxidation of pyrite in the presence of calcite and dolomite and highlighted the formation of a surface layer containing Ca (in the case of calcite) and Mg (in the case of dolomite) on the pyrite surface, which ascribed to be the cause of the slowdown in pyrite leaching rate. Pyrite alkaline oxidation rate decreased in the presence of both
5 calcite and dolomite, while a more detrimental effect was observed in the case of calcite (see "*Pyrite oxidation in the presence of calcite and dolomite: Alkaline leaching, chemical modelling and surface characterization*" in Trans. Nonferrous Met. Soc. China 28(2018) 1433–1443).

10 Caldeira et al. studied the alkaline oxidation of pyrite with different alkaline solutions to characterize the nature of a product layer formed at the surface of the oxidized pyrite. A comparison between different alkaline reagents was performed, such as with sodium or calcium hydroxide and sodium carbonate. Caldeira et al., 2003 reported that a "... very low oxidation of pyrite
15 was obtained in the presence of lime" compared to sodium carbonate which does not result in gypsum formation [C.L Caldeira, V.S.T Ciminelli, A Dias, K Osseo-Asare, Pyrite oxidation in alkaline solutions: nature of the product layer, International Journal of Mineral Processing, Volume 72, Issues 1–4, 2003, Pages 373-386].

20 Alkaline oxidation conditions have advantages compared to acidic conditions, but has a number of drawbacks: (1) alkali reagents such as sodium hydroxide, ammonium hydroxide or potassium hydroxide are expensive reagents and pose environmental risks because of the inability to readily remove the associated cation from processing circuits, while (2) calcium based alkali
25 reagents have been demonstrated to cause passivation of pyrite and thereby work counter to the objective of oxidizing refractory ore.

There is, therefore, a need to provide oxidation of sulfide minerals for recovering metal in the sulfide mineral having the minimal impact on environment, either in terms of impact or treatment of disposal, further allowing
30 the use of diverse source of water and where the capital equipment costs remain reasonable.

It is an objective of the present invention to compensate at least partly for these drawbacks by providing an alkaline oxidation process of

refractory sulfide ore particles for recovering metal which is entrapped in or co-occurring such sulfide minerals which involves less expensive alkaline agent, allows the use of diversified sources of water and where the capital equipment costs and energy consumption remain at an acceptable level.

5 To this end, the present invention relates to an alkaline oxidation process for treating refractory sulfide ore particles enriched in a metal to be recovered wherein refractory ore particles are submitted to at least 3 stages in each of which said refractory sulfide ore particles are oxidized in surface by an oxidizing agent in an alkaline oxidation step in alkaline liquid phase and form an
10 alkaline slurry containing surface oxidized refractory sulfide ore particles and in each of which said alkaline slurry is thereafter submitted to a mechanical activation step for removing from the surface oxidized refractory sulfide ore particles at least partly a surface layer containing oxidized matter, said mechanical activation forming a mechanically activated slurry containing
15 oxidized matter, refractory sulfide ore particles from which oxidized matter has been removed, alkaline liquid phase and liberated metal to be recovered by further processing or returned to a next or a previous alkaline oxidation step, said alkaline liquid phase containing calcium hydroxide as alkaline agent.

According to the present invention and against all expectations, it
20 has been possible to perform efficiently a lime based alkaline oxidation step of refractory sulfide ore particles for extracting a metal of interest. While the literature was strongly teaching away from the use of an alkaline calcium reagent compound because the surface coating effect of lime creating a passivation layer that prevents further oxidation of the refractory ore, the yield of
25 the oxidation process according to the present invention was comparable to other acid or alkaline oxidation leaching but more economical and without the drawbacks of the prior art.

Indeed, US Patent 2016/0258038A1 states that "...leaching of iron sulfide materials with lime has been unsuccessful in that leaching is incomplete
30 and subsequent precious metal recovery is low. For example, an earlier study of alkaline oxidation of pyrite for gold recovery using lime achieved only 30 to 40% gold recovery which offered little improvement over direct cyanidation of the

pyrite. This is believed to be due to passivation of the mineral by precipitation of a gypsum/iron oxide layer."

Further, in copper sulfide mineral processing, pyrite occurs as an unwanted sulfide mineral that needs to be rejected during processing. This is typically achieved by inducing surface coating effects partly attributed to lime. For example Hu et al. 2000, notes that: "Well known phenomenon of the depression of pyrite by lime is attributed to the surface formation of $\text{Ca}(\text{OH})_2$, CaSO_4 and $\text{Fe}(\text{OH})_3$ as determined by XPS analysis" [Hu et al., J. D. (2000). Surface chemistry of activation of lime-depressed pyrite in flotation. Transactions of Nonferrous Metals Society of China (English Edition), 10(6), 798-803.]

Pyrite oxidation is the main cause of acid mine drainage (also called acid rock drainage). Acid mine drainage prevention therefore focusses on preventing the oxidation of pyrite. An article by Qian 2017 states that: "treatment with lime-saturated water was found to be of paramount importance for maintaining long-term circum-neutral pH, favourable for the formation and preservation of the pyrite surface passivating layer and reduced acid generation rate" [Qian, Gujie & Schumann, Russell & Li, Jun & Short, Michael & Fan, Rong & Li, Yubiao & Kawashima, Nobuyuki & Zhou, Yan & St. C. Smart, Roger & R. Gerson, Andrea. (2017). Strategies for Reduced Acid and Metalliferous Drainage by Pyrite Surface Passivation. Minerals. 7. 1-15. 10.3390/min7030042.]

According to the present invention, it has been identified that it is possible to use lime during alkaline oxidation of refractory sulfide ore particles enriched in a metal to be recovered after at least 3 stages of alkaline oxidation with intermittent mechanical activation. During each alkaline oxidation step, a slurry is formed in which the refractory ore particles are oxidized. Thereafter the slurry is mechanically activated to remove or alter the passivation layer around the refractory ore particles in a mechanical activation step, under the form of an attrition step or milling step. The attrition step or milling step is performed during a relatively short residence time to only remove or alter the passivation layer and to limit the energy requirements for this mechanical activation step. Thereafter, the mechanically activated slurry is again submitted to a new and second stage of alkaline oxidation-mechanical activation. During the alkaline oxidation step of the second stage, the remaining portion of the refractory ore particles is further

surface-oxidized and become coated with a new passivation layer (forming a second slurry). The new passivation layer is further removed or altered during a subsequent attrition or milling step of the second slurry and forms a second mechanically activated slurry. The second mechanically activated slurry is further submitted to a third stage of alkaline oxidation step and mechanical activation step. During the alkaline oxidation step of the third stage, the remaining portion of the refractory ore particles is further oxidized in surface and surrounded with a new passivation layer (forming a third slurry). Thereafter the new passivation layer is removed or alter during a subsequent attrition or milling step of the third slurry and forms a third mechanically activated slurry. The third mechanically activated slurry is either collected for further processing or submitted to a next alkaline oxidation step, possibly by being beforehand returned to (so looping with) the previous alkaline oxidation step through the third mechanical activation step.

The number of stages is at least three and can be further increased depending on the particle size of the refractory ore particles. In addition, the residence time for each alkaline oxidation step can be the same or different from one to each other and will be defined taking into account the constraints of the process.

An unexpected finding of the present invention was that refractory ore particles could be oxidized to a substantial extent with the use of lime as alkaline agent under oxidation conditions that are otherwise known for generating a passivation layer, normally considered as an obstacle to oxidation. This was achieved by facilitating oxidation under alkaline conditions using a lime reagent, and subsequent removal/alteration of the surface passivation layer to further generate a new "active" surface and accordingly, reduce the main diameter of each refractory ore particles step by step by the stages in the process according to the present invention.

In a preferred embodiment of the alkaline oxidation process according to the present invention, said at least 3 stages comprise:

- a series of n alkaline oxidation steps in a series of n agitated reactors, each n^{th} alkaline oxidation forming a n^{th} alkaline slurry, where n is an

integer comprised between 3 and 10, preferably between 4 and 8, more preferably between 5 and 7,

- a series of x mechanical activation steps, each mechanical activation step being a mechanical activation of said n^{th} alkaline slurry, where x is an integer, equal or lower than n , and comprised between 3 and 10, preferably between 4 and 8, more preferably between 5 and 7,

wherein said series of n alkaline oxidation steps comprises at least:

- a first alkaline oxidation step wherein said refractory sulfide ore particles enriched in a metal to be recovered are fed into an agitated reactor and form a first alkaline slurry,
- at least one intermediate alkaline oxidation step fed with one alkaline slurry from a previous alkaline oxidation step,
- a last alkaline oxidation step being said n^{th} alkaline oxidation step fed with one alkaline slurry from a previous alkaline oxidation step,

said series of x mechanical activation steps comprising at least:

- a first mechanical activation step of said first alkaline slurry in a first mechanical activation means to form a first mechanically activated slurry,
- at least one intermediate mechanical activation step of a n^{th} alkaline slurry with said n^{th} alkaline slurry being comprised between a 2^{nd} alkaline slurry and a $(x-1)^{\text{th}}$ alkaline slurry in an intermediate (y^{th}) mechanical activation means to form an intermediate (y^{th}) mechanically activated slurry,
- a last mechanical activation step of said last alkaline slurry being said x^{th} mechanical activation step in a final (x^{th}) mechanical activation means to form a final (x^{th}) mechanically activated slurry.

According to the present invention, said refractory ore is a refractory sulfide ore or concentrate enriched in a metal selected among gold, silver, platinum, palladium, copper, nickel, zinc, cobalt and combinations thereof, particularly pyrite or arsenopyrite containing said metal or combination of metals, such as gold-containing pyrite ore or concentrate, gold containing arsenopyrite ore or concentrate, copper sulfide ore or concentrate, nickel sulfide

ore or concentrate, zinc sulfide ore or concentrate or cobalt sulfide ore or concentrate and combined metal sulfide ore or concentrate such as low grade copper-gold sulfide ores or concentrates.

In addition, according to the present invention, said oxidizing agent
5 is an oxidizing liquid, an oxidizing powder or an oxidizing gas, such as oxygen, ozone, and their mixture.

In case of oxygen, oxygen is preferably in the form of dissolved oxygen in the solution to be available for the reaction. Oxygen may be supplied to the oxidation steps via sparging of gas, under conditions that
10 optimize the mass transfer of oxygen from the gas phase into the liquid phase. Gas sparging may occur via several methods well-known by the skilled person, for example by means of a sparge ring, sparge jet impellers or supersonic gas injectors.

Further, in a particular embodiment, said calcium hydroxide of said
15 alkaline liquid phase is obtained by one or more addition of dry quicklime CaO , hydrated lime $\text{Ca}(\text{OH})_2$ or a milk of lime also known as slaked lime being a slurry of $\text{Ca}(\text{OH})_2$ and water, said one or more addition being chosen amongst an addition upstream one n^{th} reactor, in one n^{th} reactor, upstream a x^{th} mechanical activation means, in a x^{th} mechanical activation means or combination thereof.

20 More specifically, in the process according to the present invention, each intermediate alkaline oxidation step is fed by a previous alkaline oxidation step, optionally via a mechanical activation step.

For example, in one advantageous embodiment of the process according to the present invention, $x = n$ and each n^{th} alkaline oxidation step of
25 said series of n alkaline oxidation step is followed by a x^{th} mechanical activation step of said series of x mechanical activation steps (with the number of the series x^{th} being normally equal to the number of the series n^{th} but not mandatorily), each n^{th} (from second), i.e. each amongst the at least one intermediate alkaline oxidation step and last alkaline oxidation step being fed by a x^{th} mechanically
30 activated slurry where $x = n - 1$.

In another exemplary embodiment according to the present invention, $x = n$ and each n^{th} alkaline oxidation step of said series of n alkaline oxidation step is followed by a x^{th} mechanical activation step of said series of x mechanical activation steps (with the number of the series x^{th} being normally
5 equal to the number of the series n^{th} but not mandatorily), each n^{th} (from second), i.e. each amongst the at least one intermediate alkaline oxidation step and last alkaline oxidation step being fed by a $(n - 1)^{\text{th}}$ slurry from a $(n - 1)^{\text{th}}$ agitated reactor to which a $(n - 1)^{\text{th}}$ mechanically activated slurry is returned.

In a further exemplary embodiment according to the process of the
10 present invention, x is different from n , hence lower than n . In this exemplary embodiment, the process comprises several intermediate alkaline oxidation steps and some intermediate alkaline oxidation steps of said series of n alkaline oxidation step are followed by a mechanical activation step. In one sub-variant of this exemplary embodiment, each n^{th} intermediate alkaline oxidation step
15 forming a n^{th} slurry is fed by a $(n-1)^{\text{th}}$ slurry from the $(n- 1)^{\text{th}}$ agitated reactor, optionally subsequently mechanically activated. In another sub-variant of this exemplary embodiment, each n^{th} alkaline oxidation step forming a n^{th} slurry is fed from a $(n - 1)^{\text{th}}$ agitated reactor optionally to which a x^{th} mechanically activated slurry with $x < (n - 1)$ is returned.

20 According to the present invention, it is foreseen that mechanical activation is achieved by at least one comminution device, i.e., can be made by a single comminution device or two, three or more sequentially ordered comminution devices in parallel and/or in series, between two sequential alkaline oxidation steps.

25 In an advantageous embodiment of the process according to the present invention, at least one alkaline oxidation step, preferably more than one alkaline oxidation step and more preferably each alkaline oxidation step of said series of n alkaline oxidation steps is performed at atmospheric pressure.

30 In another advantageous embodiment of the process according to the present invention, at least one alkaline oxidation step, preferably more than one alkaline oxidation step and more preferably each alkaline oxidation step of said series of n alkaline oxidation steps is performed at a temperature

comprised between 70 and 100°C, preferably between 80 and 95°C, more preferably between 80°C and 90°C.

In a further particular embodiment of the process according to the present invention, at least one alkaline oxidation step, preferably more than one
5 alkaline oxidation step and more preferably each alkaline oxidation step of said series of n alkaline oxidation steps is performed with a dissolved oxygen content comprised between 1 and 30 mg/dm³, preferably between 5 and 25 mg/dm³, more preferably higher than 10 mg/dm³ and in particular between 10 and 20 mg/dm³ of liquid phase.

10 In the process according to the invention it is particularly possible to apply oxygen-limiting conditions to facilitate pyrite oxidation to result in ferric iron precipitate, that is substantially reduced in sulfate content (for example hematite rather than jarosite), and in sulfur, that is preferably soluble (i.e. thiosulfate or other soluble polysulfide rather than gypsum). Such conditions allow to obtain a
15 reduction of sulfate-containing iron precipitates and gypsum precipitate. In these conditions the alkaline oxidation steps are preferably performed with a dissolved oxygen content comprised between 1 and 5 mg/dm³, advantageously of 4 mg/dm³ of liquid phase.

Preferably, in the process according to the present invention, at
20 least one alkaline oxidation step, preferably more than one alkaline oxidation step and more preferably each alkaline oxidation step of said series of n alkaline oxidation steps is performed at a pH comprised between 10 and 12.5, preferably comprised between 10 and 12 and in particular between 10.5 and 11.5, said pH being controlled with a controlled addition of said alkaline agent which as
25 above disclosed is a lime reagent.

More preferably, in the process according to the present invention, at least one alkaline oxidation step, preferably more than one alkaline oxidation step and more preferably each alkaline oxidation step of said series of n alkaline oxidation steps is performed with solid content in the stirred reactor comprised
30 between 10 and 70 wt%, preferably, between 20 and 70 wt%, more preferably between 35 and 70 wt%, more particularly between 40 and 65 wt% with respect to the total weight contained in said stirred reactor.

In particular, in the process according to the present invention, at least one of said series of x mechanical activation steps is performed in a vertical mill, a vertical stirred mill, a horizontal mill, an attritor, a stirred ball mill or a horizontal stirred mill.

5 Advantageously, in the process according to the present invention, said refractory sulfide ore particles fed to said first alkaline oxidation step are obtained by a previous crushing and grinding, advantageously down to having 80% of the particles with a diameter in the range between 25 and 200 μm , preferably between 25 and 150 μm . most preferably between 30 and 120 μm .
10 Such P_{80} is measured by laser diffraction according to the standard ISO 13320-2020.

 Preferably, in the process according to the present invention, said crushed and ground refractory ore particles are subjected to mineral flotation to produce a concentrate of refractory sulfide ore particles fed to said first alkaline
15 oxidation step.

 Other embodiments of the process according to the present invention are mentioned in the appended claims.

 The present invention also relates to a device for alkaline oxidation of refractory sulfide ore particles enriched in a metal to be recovered comprising:

- 20 • a series of n agitated reactors for alkaline oxidation where n is an integer comprised between 3 and 10, preferably between 4 and 8, more preferably between 5 and 7,
- a series of x mechanical activation means where x is an integer equal or lower than n and comprised between 3 and 10, preferably between
25 4 and 8, more preferably between 5 and 7, and
- a collection means for feeding a process for recovering said metal,

said device further comprising a lime reagent feeding means connected to at least one agitated reactor or to at least one mechanical activation means or both.

30 By the wording "said device further comprising a lime reagent feeding means connected to at least one agitated reactor or to at least one mechanical activation means or both", it is meant that lime can be fed to all or

any number of the alkaline oxidation reactors and/or mechanical activation means.

Advantageously, in the device according to the present invention, at least one agitated reactor, preferably at least 3 agitated reactors, more preferably each agitated reactor, comprises sparging means for feeding an oxidizing gas.

In a particular embodiment of the device according to the present invention, said lime feeding means comprises a further stirred vessel to suspend calcium hydroxide powder in a liquid phase connected to a lime reagent pump.

In a particular embodiment of the device according to the present invention, at least one intermediate n^{th} agitated reactor of said series of n agitated reactors is connected on one side to a previous $(n-1)^{\text{th}}$ agitated reactor, optionally via a mechanical activation means, and on the other side to a following x^{th} mechanical activation means of said series of x mechanical activation means.

In another particular embodiment of the device according to the present invention, at least one intermediate n^{th} agitated reactor of said series of n agitated reactor is connected on one side to a previous $(n-1)^{\text{th}}$ agitated reactor, optionally via a mechanical activation means, and on the other side to a following $(n+1)^{\text{th}}$ agitated reactor of said series of n agitated reactors.

For example, in one exemplary embodiment of the device according to the present invention, $x = n$ and each n^{th} (from second), i.e. each amongst the at least one intermediate agitated reactor and last agitated reactor of said series of n agitated reactor is connected to a x^{th} mechanical activation means of said series of x mechanical activation means for feeding the x^{th} mechanical activation means with a n^{th} slurry of surface oxidized refractory ore particles, said x^{th} mechanical activation means being connected to a $(n + 1)^{\text{th}}$ agitated reactor for feeding said $(n + 1)^{\text{th}}$ agitated reactor with a x^{th} mechanically activated slurry.

In another exemplary embodiment of the device according to the present invention, $x = n$ and each n^{th} (from second), i.e. each amongst the at

least one intermediate agitated reactor and last agitated reactor of said series of n agitated reactor is connected to a x^{th} mechanical activation means of said series of x mechanical activation means for feeding the x^{th} mechanical activation means with a n^{th} slurry of surface oxidized refractory ore particles, said
5 x^{th} mechanical activation means being connected (inwards and outwards) to said n^{th} agitated reactor for returning (looping) said x^{th} mechanically activated slurry to said n^{th} agitated reactor, the $(n + 1)^{\text{th}}$ agitated reactor being connected to said n^{th} agitated reactor to be fed by a flow from said n^{th} agitated reactor.

In yet another exemplary embodiment according to the present
10 invention, x is different from n , hence lower than n . In this exemplary embodiment, the process comprises several intermediate agitated reactors and some intermediate agitated reactors of said series of n agitated reactors are followed by a mechanical activation step. In one sub-variant of this exemplary embodiment, each n^{th} intermediate agitated reactor forming a n^{th} slurry is
15 connected to a $(n - 1)^{\text{th}}$ agitated reactor and fed by a flow from said $(n - 1)^{\text{th}}$ agitated reactor, optionally subsequently mechanically activated. In another sub-variant of this exemplary embodiment, each n^{th} intermediate agitated reactor forming a n^{th} slurry is fed from a $(n - 1)^{\text{th}}$ agitated reactor optionally to which a x^{th} mechanically activated slurry with $x < (n - 1)$ is returned.

20 According to the present invention, it is foreseen that a mechanical activation means is made by at least one comminution device, i.e., can be made by a single comminution device or two, three or more sequentially ordered comminution devices in parallel and/or in series, between two sequential alkaline oxidation steps.

25 Preferably, in the device according to the present invention, at least one agitated reactor, preferably each agitated reactor of said series of n agitated reactor comprises heating means.

30 More particularly, in the device according to the present invention, at least one mechanical activation means of said series of x mechanical activation means is a vertical mill, a vertical stirred mill, a horizontal mill, an attritor, a stirred ball mill or a horizontal stirred mill.

Other embodiments of the device according to the present invention are mentioned in the appended claims.

Other characteristics and advantages of the present invention will be derived from the non-limitative following description, and by making
5 reference to the drawings.

Figure 1 is a schematic view of the status of a refractory ore material particle during one stage of alkaline oxidation and mechanical activation.

Figure 2 is a flow chart of one embodiment of the process according to the present invention, showing the device to carry out such
10 process.

Figure 3 is a flow chart of another embodiment of the process according to the present invention, showing the device to carry out such process.

Figure 4 is a flow chart of yet another embodiment of the process according to the present invention, showing the device to carry out such
15 process.

Figure 5 is a flow chart of a further embodiment of the process according to the present invention, showing the device to carry out such process.

Figure 6 is a flow chart of a variant embodiment of the process according to the present invention, showing the device to carry out such
20 process.

Figure 7 is a graph showing the extent of gold extraction by cyanidation leaching (or other target method) as a function of extent of pyrite
25 oxidation.

In the drawings, the same reference numbers have been allocated to the same or analogue element.

The present invention relates to an alkaline oxidation process for treating refractory sulfide ore particles enriched in a metal to be recovered.

The refractory ore particles are submitted to at least 3 stages, preferably 4, 5, 6, 7 or even 8 or up to 10 stages, in which refractory ore particles are oxidized in surface in an alkaline oxidation step in alkaline liquid phase
30 containing calcium hydroxide for forming an alkaline slurry containing surface

oxidized refractory ore particles. Thereafter the slurry is mechanically activated to remove at least partly a surface layer from the surface oxidized refractory ore particles. The surface layer contains oxidized matter. The mechanical activation forms a slurry which is called a mechanically activated slurry and which contains
5 oxidized matter, refractory sulfide ore or concentrate particles from which said surface layer has been removed, alkaline liquid phase and liberated metal to be recovered for further processing.

The process according to the present invention is contrary to the conventional industrial practice of using lime-based reagents, where it is
10 generally used to retard sulfide mineral oxidation. The present invention is based on the use of a lime-based reagent to induce alkaline conditions for sulfide mineral oxidation by explicitly overcoming the gypsum and iron oxide coating effect, thereby facilitating high rates of sulfide mineral oxidation.

Refractory ore is preferably crushed and milled in order to typically
15 obtain at least 80% of particles having a diameter between 25 and 150 μm . This ore can either be fed directly into the first alkaline oxidation reactor, or can first be subjected to a mineral flotation process and thereby upgraded to a concentrate. The upgraded concentrate A, containing the metal and sulfide mineral of interest, is then be fed into the first alkaline agitated oxidation reactor.

The present invention is herein illustrated with gold refractory ore, without being
20 limited thereto. Other precious metal refractory ore can also be treated in the process according to the present invention, for example refractory ores containing silver, palladium or platinum, but also base metal sulfide ore, such as zinc sulfide ore, copper sulfide ore, cobalt sulfide ore, nickel sulfide ore or
25 combined metal sulfide ore such as low grade copper-gold sulfide ores.

In refractory sulfide gold ore, the main sulfide minerals in the concentrate are most typically pyrite or arsenopyrite. Lime as dry quicklime, dry hydrated lime (also known as slaked lime or dry slaked lime), or a milk of lime (a slurry of $\text{Ca}(\text{OH})_2$ particles and water) or a paste of lime may be added to at
30 least one agitated reactor and/or to at least one mechanical activation means, preferably to at least 3 agitated reactors and/or to at least 3 mechanical activation means, more preferably to each agitated reactors and/or to at least each mechanical activation means. Lime (in any form) is added to as many of

the process units (agitated reactors and/or mechanical activation means) as is required to maintain the pH throughout the processing circuit at the target pH setpoint. Practically, if the lime is added to each oxidation reactor, to maintain the pH at the setpoint, the lime consumption can be used as an indicator of the extent of mineral oxidation. Therefore it could be a very important parameter to monitor the performance of the process circuit.

It is also possible that the lime is added to each stage of the process by adding it to the attritioner feed. In this way the mechanical activation means acts to mill and disperse added lime.

The sulfide mineral concentrate (1) is subjected to a succession of alkaline oxidation ((CSTR)_n) and surface attritioning ((MA)_n) in order to reduce the impact of surface coating and passivation effects, illustrated in figure 1. The surface layer (2) or passivation layer (2) contains ferric iron hydroxide (Fe(OH)₃) and gypsum (CaSO₄·2 H₂O). The surface attritioning forms a slurry which is called a mechanically activated slurry and which contains oxidized matter (4), refractory sulfide ore or concentrate particles from which oxidized matter has been removed (3), alkaline liquid phase and liberated metal to be recovered (5).

The sulfide mineral concentrate is subjected to an elevated temperature alkaline oxidation process into which oxygen gas (preferably with an oxygen content greater than 95% v/v) is sparged and lime is added.

A combination of surface attritioning (mechanical activation) and alkaline oxidation, prior to gold leaching, is thereby induced. The rationale for this approach is to limit milling/grinding to achieve mineral surface attritioning (also called herein mechanical activation and encompassing both mechanical activation or attritioning) in a stirred media reactor sufficient to disrupt/alter or remove gypsum and iron oxide surface coating and passivation. In effect, the alkaline oxidation process is applied to mineral surfaces and milling (mechanical activation) is only used to expose fresh mineral surfaces ("active" layer) to sustain elevated mineral oxidation rates, as illustrated in figure 1. The process of mechanically activation also results in a high degree of strain being introduced into the sulphide mineral lattice, increasing the number of grain boundary fractures and lattice defects in the mineral. The introduction of strain lowers the

activation energy for the oxidation of the sulfides and enables oxidation under atmospheric conditions. The rate of oxidation is also enhanced, due to the increased mineral surface area.

The process according to the present invention can occur in a number of different circuit configurations, with the basic feature of alternating alkaline oxidation (shown in CSTR reactors below) with inter-stage surface attritioning (also called mechanical activation, MA). The number of stages can range from 3 – 10, preferably from 3 to 8. More preferably from 4 to 7.

As it can be seen from the figures 2 to 6, alkaline sulfide oxidation process for treating refractory ore particles enriched in said metal to be collected comprises said at least 3 stages comprising:

- a series of n alkaline oxidation steps in a series of n agitated reactors (CSTRs) forming n alkaline slurries, where n is an integer comprised between 3 and 10, preferably between 4 and 8, more preferably between 5 and 7,
- a series of x mechanical activation steps in a series of x mechanical activation means (MA), each mechanical activation step being a mechanical activation of a n^{th} alkaline slurry, where x is an integer, equal to or lower than n , and comprised between 3 and 10, preferably between 4 and 8, more preferably between 5 and 7.

The series of n alkaline oxidation steps comprises at least:

- a first alkaline oxidation step where said refractory ore particles enriched in a metal to be recovered are fed into an agitated reactor and form a first alkaline slurry,
- at least one intermediate alkaline oxidation step fed with one alkaline slurry from a previous alkaline oxidation step,
- a last alkaline oxidation step being said n^{th} alkaline oxidation step fed with one alkaline slurry from a previous alkaline oxidation step.

Said series of x mechanical activation steps comprises at least:

- a first mechanical activation step of said first alkaline slurry in a first mechanical activation means to form a first mechanically activated slurry,

- at least one intermediate mechanical activation step of a n^{th} alkaline slurry with said n^{th} alkaline slurry being comprised between a 2^{nd} alkaline slurry and a $(x-1)^{\text{th}}$ alkaline slurry in an intermediate (y^{th}) mechanical activation means to form an intermediate (y^{th}) mechanically activated slurry,
- a last mechanical activation step of said last alkaline slurry being said x^{th} mechanical activation step in a x^{th} mechanical activation means to form a last mechanically activated slurry.

A first embodiment is illustrated in figure 2. As it can be seen, the series of n reactor comprises 6 agitated reactors. Each agitated reactor (CSTR 1 to CSTR 6) is followed by one mechanical activation means (MA 1 to MA 6). In this embodiment of figure 2, $x = n = 6$. The first agitated reactor is fed with refractory ore particles, having preferably a P_{80} lower than $150 \mu\text{m}$. Each alkaline oxidation step of said series of n alkaline oxidation step is followed by a mechanical activation step of said series of x mechanical activation steps. Each n^{th} alkaline oxidation step forms a n^{th} slurry of surface oxidized refractory ore particles. Each n^{th} alkaline oxidation step from $n = 2$ is fed by a x^{th} mechanically activated slurry where $x = n - 1$. Each x^{th} mechanical activation step forms a x^{th} mechanically activated slurry.

In other words, the first alkaline oxidation step in agitated reactor CSTR 1 forms a first slurry of surface oxidized refractory ore particles and is followed by a first mechanical activation step in a mechanical activation means MA 1. The first mechanical activation forms a first mechanically activated slurry. The first alkaline oxidation step is fed by refractory ore particles.

The second alkaline oxidation step in agitated reactor CSTR 2 forms a second slurry of surface oxidized refractory ore particles and is followed by a second mechanical activation step in a mechanical activation means MA 2. The second mechanical activation forms a second mechanically activated slurry. The second alkaline oxidation step is fed by the first mechanically activated slurry.

The third alkaline oxidation step in agitated reactor CSTR 3 forms a third slurry of surface oxidized refractory ore particles and is followed by a third mechanical activation step in a mechanical activation means MA 3. The third

mechanical activation forms a third mechanically activated slurry. The third alkaline oxidation step is fed by the second mechanically activated slurry.

The fourth alkaline oxidation step in agitated reactor CSTR 4 forms a fourth slurry of surface oxidized refractory ore particles and is followed by a
5 fourth mechanical activation step in a mechanical activation means MA 4. The fourth mechanical activation forms a fourth mechanically activated slurry. The fourth alkaline oxidation step is fed by the third mechanically activated slurry.

The fifth alkaline oxidation step in agitated reactor CSTR 5 forms a fifth slurry of surface oxidized refractory ore particles and is followed by a fifth
10 mechanical activation step in a mechanical activation means MA 5. The fifth mechanical activation forms a fifth mechanically activated slurry. The fifth alkaline oxidation step is fed by the fourth mechanically activated slurry.

The sixth alkaline oxidation step in agitated reactor CSTR 6 forms a sixth slurry of surface oxidized refractory ore particles and is followed by a sixth
15 mechanical activation step in a mechanical activation means MA 6. The sixth mechanical activation forms a sixth mechanically activated slurry. The sixth alkaline oxidation step is fed by the fifth mechanically activated slurry.

In another alternative arrangement as illustrated in figure 3, the attritioners or mechanical activation means (MA 1 to MA 6) will not be between
20 the alkaline oxidation stages (CSTR 1 to CSTR 6), but rather in parallel with the oxidation stages, forming an internal loop between the alkaline oxidation reactor and the mechanical activation reactor.

The equipment arrangement would have preferably 4 to 6 reactors for alkaline oxidation, with 4 to 6 attritioners "next to" the reactors.

25 In this case, the slurry is drawn from a n^{th} agitated reactor, sent through the attritioner, which then returns the slurry to the same n^{th} reactor.

This arrangement offers the added benefit that the circuit may continue to operate even if an attritioner needs to be taken offline for maintenance or replacement. It also provides more flexibility to alter the relative
30 extent of attritioning vs oxidation, by adjusting the return (to the same oxidation reactor) versus the forwarding (to the next oxidation reactor in the series) flow rate. Also, not all alkaline oxidation stages may require a mechanical activation step, therefore some mechanical activation steps can be omitted (see figure 5).

Indeed, as illustrated in figure 3, in this arrangement, the number of agitated reactors in said series is preferably 6 and the number of mechanical activation means in said series is also preferably 6.

Accordingly, $x = n = 6$. Each alkaline oxidation step of said series of n alkaline oxidation step is followed by a mechanical activation step of said series of x mechanical activation steps. Each n^{th} alkaline oxidation step forms a n^{th} slurry of surface oxidized refractory ore particles. The n^{th} alkaline oxidation step is fed by a $(n - 1)^{\text{th}}$ slurry from a $(n - 1)^{\text{th}}$ agitated reactor to which a $(n - 1)^{\text{th}}$ mechanically activated slurry is returned.

In other words, the first alkaline oxidation step in agitated reactor CSTR 1 forms a first slurry of surface oxidized refractory ore particles and is followed by a first mechanical activation step in a mechanical activation means MA1. The first mechanical activation forms a first mechanically activated slurry. The first alkaline oxidation step is fed by refractory ore particles and by said first mechanically activated slurry.

The second alkaline oxidation step in agitated reactor CSTR 2 forms a second slurry of surface oxidized refractory ore particles and is followed by a second mechanical activation step in a mechanical activation means MA 2. The second mechanical activation forms a second mechanically activated slurry. The second alkaline oxidation step is fed by the first slurry from the first agitated reactor and by the second mechanically activated slurry.

The third alkaline oxidation step in agitated reactor CSTR 3 forms a third slurry of surface oxidized refractory ore particles and is followed by a third mechanical activation step in a mechanical activation means MA 3. The third mechanical activation forms a third mechanically activated slurry. The third alkaline oxidation step is fed by the second slurry from the second agitated reactor and by the third mechanically activated slurry.

The fourth alkaline oxidation step in agitated reactor CSTR 4 forms a fourth slurry of surface oxidized refractory ore particles and is followed by a fourth mechanical activation step in a mechanical activation means MA 4. The fourth mechanical activation forms a fourth mechanically activated slurry. The fourth alkaline oxidation step is fed by the third slurry from the third agitated reactor and by the fourth mechanically activated slurry.

The fifth alkaline oxidation step in agitated reactor CSTR 5 forms a fifth slurry of surface oxidized refractory ore particles and is followed by a fifth mechanical activation step in a mechanical activation means MA 5. The fifth mechanical activation forms a fifth mechanically activated slurry. The fifth alkaline oxidation step is fed by the fourth slurry from the fourth agitated reactor and by the fifth mechanically activated slurry.

The sixth alkaline oxidation step in agitated reactor CSTR 6 forms a sixth slurry of surface oxidized refractory ore particles and is followed by a sixth mechanical activation step in a mechanical activation means MA 6. The sixth mechanical activation forms a sixth mechanically activated slurry. The sixth alkaline oxidation step is fed by the fifth slurry from the fifth agitated reactor and by the sixth mechanically activated slurry.

Figure 4 illustrates a variant embodiment compared to figure 2, where the number of agitated reactors in said series of n agitated reactor is not the same as the number of mechanical activation means of said series of x mechanical activation means. This arrangement can be carried out as such or at any location in the series and can be carried out as illustrated by construction or because one mechanical activation means needs to be maintained. In this case, a by-pass is realized between two consecutive agitated reactors.

As it can be seen, in this arrangement, x is different from n , hence lower than n . $x = 5$ while $n = 6$. Some alkaline oxidation step of said series of n alkaline oxidation step are followed by a mechanical activation step of said series of x mechanical activation steps, but not each alkaline oxidation steps. Each n^{th} alkaline oxidation step forms a n^{th} slurry of surface oxidized refractory ore particles and is fed by a $(n-1)^{\text{th}}$ slurry from the $(n-1)^{\text{th}}$ agitated reactor, optionally subsequently mechanically activated.

In other words, the first alkaline oxidation step in agitated reactor CSTR 1 forms a first slurry of surface oxidized refractory ore particles and is followed by a first mechanical activation step in a mechanical activation means MA1. The first mechanical activation forms a first mechanically activated slurry. The first alkaline oxidation is fed by refractory ore particles.

The second alkaline oxidation step in agitated reactor CSTR 2 forms a second slurry of surface oxidized refractory ore particles. The second alkaline oxidation step is fed by the first mechanically activated slurry.

5 The third alkaline oxidation step in agitated reactor CSTR 3 forms a third slurry of surface oxidized refractory ore particles and is followed by a second mechanical activation step in a mechanical activation means MA 2. The second mechanical activation forms a second mechanically activated slurry. The third alkaline oxidation step is fed by the second slurry from the second alkaline oxidation step.

10 The fourth alkaline oxidation step in agitated reactor CSTR 4 forms a fourth slurry of surface oxidized refractory ore particles and is followed by a third mechanical activation step in a mechanical activation means MA 3. The third mechanical activation forms a third mechanically activated slurry. The fourth alkaline oxidation step is fed by the second mechanically activated slurry.

15 The fifth alkaline oxidation step in agitated reactor CSTR 5 forms a fifth slurry of surface oxidized refractory ore particles and is followed by a fourth mechanical activation step in a mechanical activation means MA 4. The fourth mechanical activation forms a fourth mechanically activated slurry. The fifth alkaline oxidation step is fed by the third mechanically activated slurry.

20 The sixth alkaline oxidation step in agitated reactor CSTR 6 forms a sixth slurry of surface oxidized refractory ore particles and is followed by a fifth mechanical activation step in a mechanical activation means MA 5. The fifth mechanical activation forms a fifth mechanically activated slurry. The sixth alkaline oxidation step is fed by the fourth mechanically activated slurry.

25 Figure 5 illustrates a variant embodiment compared to figure 3, where the number of agitated reactor in said series of n agitated reactor is not the same as the number of mechanical activation means of said series of x mechanical activation means. This arrangement can be carried out as such or at any location in the series and can be carried out as illustrated by construction
30 or because one mechanical activation means needs to be maintained, as afore explained.

As it can be seen, in this arrangement, x is different from n , and lower than n . $x = 5$ while $n = 6$. Some alkaline oxidation step of said series of n

alkaline oxidation steps (CSTR 1 to CSTR 6) being followed by a mechanical activation step of said series of x mechanical activation steps (MA 1 to MA 5). Each n th alkaline oxidation step forming a n th slurry of surface oxidized refractory ore particles and being fed from a $(n - 1)$ th agitated reactor optionally to which a x th mechanically activated slurry with $x < (n - 1)$ is returned.

In other words, the first alkaline oxidation step in agitated reactor CSTR 1 forms a first slurry of surface oxidized refractory ore particles and is followed by a first mechanical activation step in a mechanical activation means MA 1. The first mechanical activation forms a first mechanically activated slurry. The first alkaline oxidation is fed by refractory ore particles and by said first mechanically activated slurry.

The second alkaline oxidation step in agitated reactor CSTR 2 forms a second slurry of surface oxidized refractory ore particles and is followed by a second mechanical activation step in a mechanical activation means MA 2. The second mechanical activation forms a second mechanically activated slurry. The second alkaline oxidation step is fed by the first slurry from the first agitated reactor and by the second mechanically activated slurry.

The third alkaline oxidation step in agitated reactor CSTR 3 forms a third slurry of surface oxidized refractory ore particles. The third alkaline oxidation step is fed by the second slurry from the second agitated reactor.

The fourth alkaline oxidation step in agitated reactor CSTR 4 forms a fourth slurry of surface oxidized refractory ore particles and is followed by a third mechanical activation step in a mechanical activation means MA 3. The third mechanical activation forms a third mechanically activated slurry. The fourth alkaline oxidation step is fed by the third slurry from the third agitated reactor and by the third mechanically activated slurry.

The fifth alkaline oxidation step in agitated reactor CSTR 5 forms a fifth slurry of surface oxidized refractory ore particles and is followed by a fourth mechanical activation step in a mechanical activation means MA 4. The fourth mechanical activation forms a fourth mechanically activated slurry. The fifth alkaline oxidation step is fed by the fourth slurry from the fourth agitated reactor and by the fourth mechanically activated slurry.

The sixth alkaline oxidation step in agitated reactor CSTR 6 forms a sixth slurry of surface oxidized refractory ore particles and is followed by a fifth mechanical activation step in a mechanical activation means MA 5. The fifth mechanical activation forms a fifth mechanically activated slurry. The sixth alkaline oxidation step is fed by the fifth slurry from the fifth agitated reactor and by the fifth mechanically activated slurry.

Figure 6 shows another arrangement in which a combination of several variants is illustrated.

In this embodiment, the first alkaline oxidation step in agitated reactor CSTR 1 forms a first slurry of surface oxidized refractory ore particles and is followed by a first mechanical activation step in a mechanical activation means MA1. The first mechanical activation forms a first mechanically activated slurry. The first alkaline oxidation is fed by refractory ore particles and the first mechanically activated slurry.

The second alkaline oxidation step in agitated reactor CSTR 2 forms a second slurry of surface oxidized refractory ore particles and is followed by a second mechanical activation step in a mechanical activation means MA 2. The second mechanical activation forms a second mechanically activated slurry. The second alkaline oxidation step is fed by the first slurry from the first agitated reactor and by the second mechanically activated slurry.

The third alkaline oxidation step in agitated reactor CSTR 3 forms a third slurry of surface oxidized refractory ore particles. The third alkaline oxidation step is fed by the second slurry from the second agitated reactor.

The fourth alkaline oxidation step in agitated reactor CSTR 4 forms a fourth slurry of surface oxidized refractory ore particles and is followed by a third mechanical activation step in a mechanical activation means MA 3. The third mechanical activation forms a third mechanically activated slurry. The fourth alkaline oxidation step is fed by the third slurry from the third agitated reactor.

The fifth alkaline oxidation step in agitated reactor CSTR 5 forms a fifth slurry of surface oxidized refractory ore particles. The fifth alkaline oxidation step is fed by the third mechanically activated slurry from the third mechanical activation means.

The sixth alkaline oxidation step in agitated reactor CSTR 6 forms a sixth slurry of surface oxidized refractory ore particles and is followed by a fourth mechanical activation step in a mechanical activation means MA 4. The fourth mechanical activation forms a fourth mechanically activated slurry. The sixth alkaline oxidation step is fed by the fifth slurry from the fifth agitated reactor.

The seventh alkaline oxidation step in agitated reactor CSTR 7 forms a seventh slurry of surface oxidized refractory ore particles and is followed by a fifth mechanical activation step in a mechanical activation means MA 5. The fifth mechanical activation forms a fifth mechanically activated slurry. The seventh alkaline oxidation step is fed by the fourth mechanically activated slurry from the fourth mechanical activation means.

According to the present invention, further possible separation steps can be foreseen between two agitated oxidation reactors or between one mechanical activation means and one agitated reactor, such as for example to remove a portion of said liquid phase and keeping a higher concentration in solid matter in a further alkaline oxidation step.

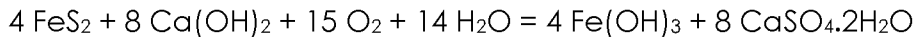
In the process according to the present invention, the conditions for alkaline oxidation process are preferably as follows:

- Pressure: atmospheric
- Solids concentration: 40 – 65% wt
- Temperature: 80 – 90°C
- Dissolved oxygen: 2 - 30mg/dm³ (preferably > 10 mg/dm³)
- pH control level: 11 with controlled lime addition

Lime reagent addition into the oxidation reactors (CSTR) and stirred media (attritioning or mechanical activation) reactors can be achieved either by addition of dry quicklime (CaO – in powder form), lime hydrate (Ca(OH)₂ – in powder form), or milk of lime also known as slaked lime slurry (a suspension of Ca(OH)₂). The lime addition rate would be dependent upon reaction demand and maintaining the pH at 11 in all alkaline oxidation reactors. Said lime addition in the oxidation reactors and mechanical activation means is performed in an usual known manner which is not illustrated on the Figures.

The heat for maintaining the temperature at or above 80°C is derived from the stirred media reactor as well as the exothermic chemical reactions and is sustained by managing the various heat balance factors of the processing circuit. These heat balance factors are specific to each application scenario and should be evaluated for each individual scenario as key inputs into the process design and selection of lime reagent type. The mass feed and subsequent rate of sulfide oxidation, as well the type of lime reagent used are particularly relevant to the heat balance.

The alkaline oxidation reaction, summarized by the overall reaction below, is exothermic ($\Delta H -1732\text{kJ/mol @ } 30^\circ\text{C}$) thereby contributing to heat generation in the agitated reactor.



The hydration reaction of quicklime is also exothermic ($\Delta H -65\text{kJ/mol @ } 30^\circ\text{C}$).



The choice of lime reagent in the form of CaO, compared to already-hydrated Ca(OH)₂ may impact the heat balance, and may be used to manage the amount of heat in the process.

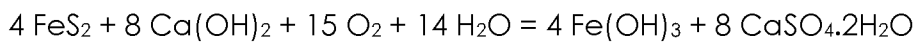
Oxygen is supplied preferably in each oxidation reactor in a known manner which is not illustrated on the Figures. Oxygen utilization efficiency may also be improved by making use of a cascade of multiple oxidation reactors where the oxygen gas from one reactor is captured and introduced to a next reactor vessel in the cascade.

A key design element of the processing circuit is the extent of pyrite (or other sulfide mineral) oxidation to be targeted, as this is the key driver for determining the extent surface attrition, the extent of oxidation and therefore also the extent of lime and oxygen reagent requirement. In turn, these factors influence the process residence time and thereby the reactor size. These factors thus also influence the operational and capital costs of the processing circuit.

The optimal solids content is determined by optimal viscosity of the solids slurry. A too viscous slurry will reduce the efficiency of oxygen mass transfer and would negatively impact the efficiency of attritioning. The design of the process circuit should take into account that the that the solids content, and thus
5 viscosity, of the slurry will increase through the circuit as a result of increased gypsum generation. The process design should take into consideration this increased viscosity and the optimal viscosity for oxygen transfer and attritioning or mechanical activation. Dilution process water may also be introduced along
10 the process circuit to reduce the solids concentration and viscosity if required to maintain optimal conditions

The extent of gold extraction by cyanidation leaching (or other target method) should be determined as a function of extent of pyrite oxidation, as is known in the art and shown by way of example in the graph in figure 7.

The process circuit design, and in particular the extent of surface
15 attritioning and oxidation is based on obtaining key process design parameters experimentally. Lime consumption (to maintain the pH at a setpoint of 11), can be used as an indicator or proxy of the extent of pyrite oxidation because of the correlation via the reaction mechanism:



20 Once initial oxidation is commenced, using the concentrate feed A at particle size as obtained from the mineral flotation process, lime and/or oxygen consumption is monitored (according to the above reaction) until a plateau is reached indicating the onset of the particle coating effect caused by gypsum and iron oxides. The mineral slurry is then transferred to a surface
25 attritioning step to induce surface cleaning for a monitored period of time, that can be adjusted as required. Oxidation is again resumed in the same way until a lime/oxygen consumption plateau is reached. The stage of said series of at least 3 stages is repeated until no further oxidation occurs.

The lime consumption is correlated with the oxygen consumption
30 via the reaction mechanism, which allows for the oxygen demand for the system

to be calculated and the oxygen supply system to be designed for the oxidation reactors.

It should be understood that the present invention is not limited to the described embodiments and that variations can be applied without going
5 outside of the scope of the claims.

CLAIMS

1. Alkaline oxidation process for treating refractory sulfide ore particles enriched in a metal to be recovered wherein refractory sulfide ore particles are submitted to at least 3 stages in each of which said refractory sulfide ore particles (1) are oxidized in surface by an oxidizing agent in an alkaline oxidation step (CSTR) in alkaline liquid phase and form an alkaline slurry containing surface oxidized refractory sulfide ore particles and in each of which said alkaline slurry is thereafter submitted to a mechanical activation step (MA) for removing from the surface oxidized refractory sulfide ore particles at least partly a surface layer (2) containing oxidized matter, said mechanical activation forming a mechanically activated slurry containing oxidized matter (4), refractory sulfide ore particles from which oxidized matter has been removed (3), alkaline liquid phase and liberated metal (5) to be recovered by further processing or returned to a next or a previous alkaline oxidation step, said alkaline liquid phase containing calcium hydroxide as alkaline agent.

2. Alkaline oxidation process for treating refractory sulfide ore particles enriched in said metal to be recovered according to claim 1, wherein said at least 3 stages comprise:

- a series of n alkaline oxidation steps in a series of n agitated reactors, each n^{th} alkaline oxidation forming a n^{th} alkaline slurry, where n is an integer comprised between 3 and 10, preferably between 4 and 8, more preferably between 5 and 7,
- a series of x mechanical activation steps, each mechanical activation step being a mechanical activation of said n^{th} alkaline slurry, where x is an integer, equal or lower than n , and comprised between 3 and 10, preferably between 4 and 8, more preferably between 5 and 7,

wherein said series of n alkaline oxidation steps comprises at least:

- a first alkaline oxidation step wherein said refractory sulfide ore particles enriched in a metal to be recovered are fed into an agitated reactor and form a first alkaline slurry,

- at least one intermediate alkaline oxidation step fed with one alkaline slurry from a previous alkaline oxidation step,
- a last alkaline oxidation step being said n^{th} alkaline oxidation step fed with one alkaline slurry from a previous alkaline oxidation step,

5 said series of x mechanical activation steps comprising at least:

- a first mechanical activation step of said first alkaline slurry in a first mechanical activation means to form a first mechanically activated slurry,
- at least one intermediate mechanical activation step of a n^{th} alkaline slurry with said n^{th} alkaline slurry being comprised between a 2^{nd} alkaline slurry and a $(x-1)^{\text{th}}$ alkaline slurry in an intermediate mechanical activation means to form an intermediate mechanically activated slurry,
- a last mechanical activation step of said last alkaline slurry being said x^{th} mechanical activation step in a x^{th} mechanical activation means to form a last mechanically activated slurry.

10

15

3. Alkaline oxidation process according to any of the claim 1 or claim 2, wherein said refractory sulfide ore is a refractory sulfide ore or concentrate enriched in a metal selected among gold, silver, platinum, palladium, copper, nickel, zinc, cobalt and combinations thereof, particularly pyrite or arsenopyrite containing said metal or combination of metals.

20

4. Alkaline oxidation process according to any of the claims 1 to 3, wherein said oxidizing agent is an oxidizing liquid, an oxidizing powder or an oxidizing gas, such as oxygen, ozone, peroxide, and their mixture.

25

5. Alkaline oxidation process according to any of the claims 2 to 4, wherein said calcium hydroxide of said alkaline liquid phase is obtained by one or more addition of dry quicklime CaO , hydrated lime $\text{Ca}(\text{OH})_2$, or milk of lime, said one or more addition being chosen amongst an addition upstream one n^{th} reactor, in one n^{th} reactor, upstream a x^{th} mechanical activation means, in a x^{th} mechanical activation means or combination thereof.

30

6. Alkaline oxidation process according to any of the claims 2 to 5, wherein each intermediate alkaline oxidation step is fed by a previous alkaline oxidation step, optionally via a mechanical activation step.

7. Alkaline oxidation process according to any of the claims 2 to 6, wherein at least one alkaline oxidation step, preferably more than one alkaline oxidation step and more preferably each alkaline oxidation step of said series of n alkaline oxidation steps is performed at atmospheric pressure.

5 8. Alkaline oxidation process according to any of the claims 2 to 7, wherein at least one alkaline oxidation step, preferably more than one alkaline oxidation step and more preferably each alkaline oxidation step of said series of n alkaline oxidation steps is performed at a temperature comprised between 70 and 100°C, preferably between 80 and 95°C, more preferably
10 between 80°C and 90°C.

9. Alkaline oxidation process according to any of the claims 2 to 8, wherein at least one alkaline oxidation step, preferably more than one alkaline oxidation step and more preferably each alkaline oxidation step of said series of n alkaline oxidation steps is performed with a dissolved oxygen content
15 comprised between 1 and 30 mg/dm³, preferably between 5 and 25 mg/dm³, more preferably higher than 10 mg/dm³ and in particular between 10 and 20 mg/dm³ of liquid phase.

10. Alkaline oxidation process according to claim 9, wherein at least one alkaline oxidation step, preferably more than one alkaline oxidation
20 step and more preferably each alkaline oxidation step of said series of n alkaline oxidation steps is performed with a dissolved oxygen content comprised between 1 and 5 mg/dm³, preferably of 4 mg/dm³ of liquid phase.

11. Alkaline oxidation process according to any of the claims 2 to 10, wherein at least one alkaline oxidation step, preferably more than one alkaline oxidation
25 step and more preferably each alkaline oxidation step of said series of n alkaline oxidation steps is performed at a pH comprised between 10 and 12.5, preferably comprised between 10 and 12 and in particular between 10.5 and 11.5, said pH being controlled with a controlled addition of said alkaline agent.

12. Alkaline oxidation process according to any of the claims 2 to 11, wherein at least one alkaline oxidation step, preferably more than one alkaline oxidation
30 step and more preferably each alkaline oxidation step of said series of n alkaline oxidation steps is performed with solid content in the agitated

reactor comprised between 10 and 70 wt%, preferably, between 20 and 70 wt%, more preferably between 35 and 70 wt%, more particularly between 40 and 65 wt% with respect to the total weight contained in said agitated reactor.

13. Alkaline oxidation process according to any of the claims 2
5 to 12, wherein at least one of said series of x mechanical activation steps is performed in a vertical mill, a vertical stirred mill, a horizontal mill, an attritor, a stirred ball mill or a horizontal stirred mill.

14. Alkaline oxidation process according to any of the claims 2
10 to 13, wherein said refractory sulfide ore particles fed to said first alkaline oxidation step are obtained by a previous crushing and grinding, advantageously down to having 80% of the particles with a diameter in the range between 25 and 200 μm , preferably between 25 and 150 μm . most preferably between 30 and 120 μm .

15. Alkaline oxidation process according to claim 14, wherein
15 said crushed and ground refractory sulfide ore particles are subjected to mineral flotation to produce a concentrate of refractory sulfide ore particles fed to said first alkaline oxidation step.

16. Device for alkaline oxidation of refractory sulfide ore
particles enriched in a metal to be recovered, comprising:

- 20
- a series of n agitated reactors for alkaline oxidation (CSTR) where n is an integer comprised between 3 and 10, preferably between 4 and 8, more preferably between 5 and 7,
 - a series of x mechanical activation means (MA) where x is an integer equal or lower than n and comprised between 3 and 10, preferably
25 between 4 and 8, more preferably between 5 and 7, and
 - a collection means (B) for feeding a process for recovering said metal,

said device further comprising a lime reagent feeding means connected to at least one agitated reactor or to at least one mechanical activation means or both.

17. Device for alkaline oxidation according to claim 16, wherein
30 at least one agitated reactor, preferably at least 3 agitated reactors, more

preferably each agitated reactor, comprises sparging means for feeding an oxidizing gas.

18. Device for alkaline oxidation according to claim 16 or claim 17, wherein said lime reagent feeding means comprises a further stirred vessel to
5 suspend calcium hydroxide powder in a liquid phase connected to a lime reagent pump.

19. Device for alkaline oxidation according to any of the claims 16 to 18, wherein at least one intermediate n^{th} agitated reactor of said series of n agitated reactors is connected on one side to a previous $(n-1)^{\text{th}}$ agitated
10 reactor, optionally via a mechanical activation means, and on the other side to a following x^{th} mechanical activation means of said series of x mechanical activation means.

20. Device for alkaline oxidation according to any of the claims 16 to 18, wherein at least one intermediate n^{th} agitated reactor of said series of n agitated reactor is connected on one side to a previous $(n-1)^{\text{th}}$ agitated
15 reactor, optionally via a mechanical activation means, and on the other side to a following $(n+1)^{\text{th}}$ agitated reactor of said series of n agitated reactors.

21. Device for alkaline oxidation according to any of the claims 16 to 20, wherein at least one agitated reactor of said series of n agitated reactor
20 comprises heating means.

22. Device for alkaline oxidation according to any of the claims 16 to 21, wherein at least one mechanical activation means of said series of x mechanical activation means is a vertical mill, a vertical stirred mill, a horizontal mill, an attritor, a stirred ball mill or a horizontal stirred mill.

25

Fig. 1

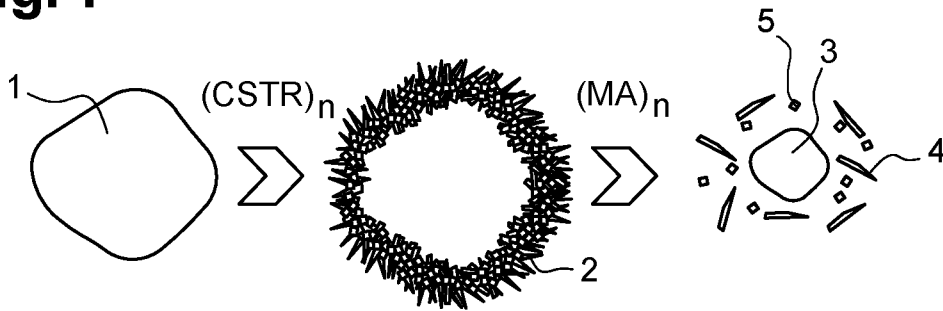


Fig. 2

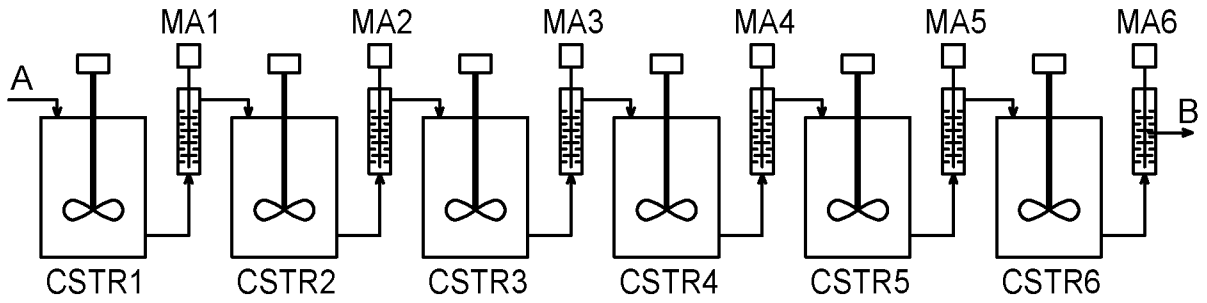


Fig. 3

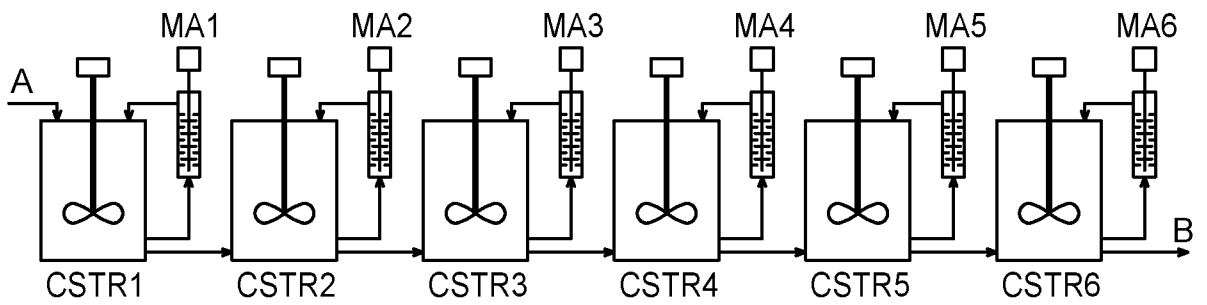


Fig. 4

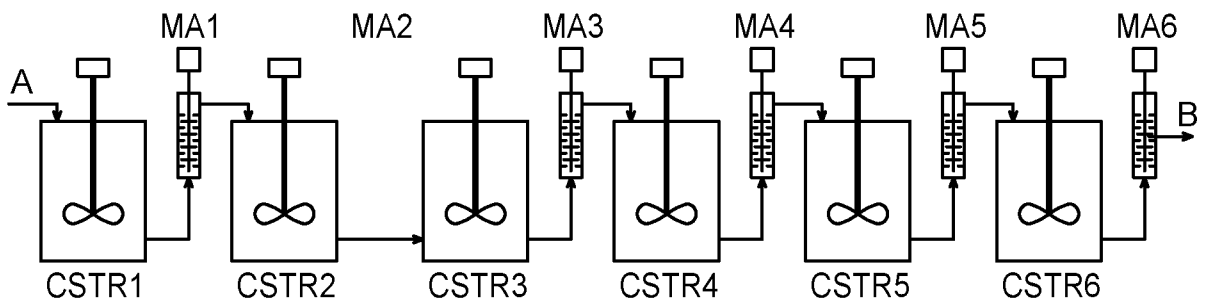


Fig. 5

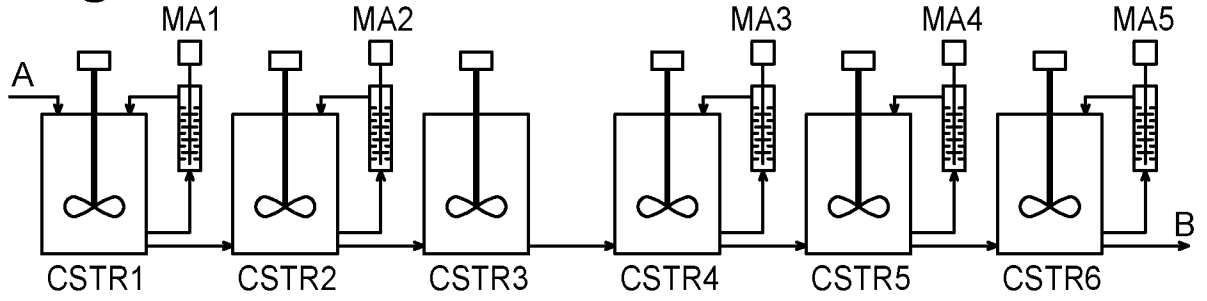


Fig. 6

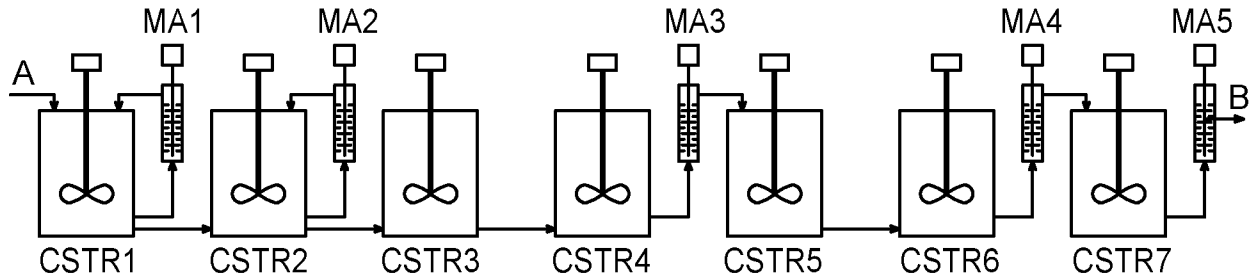
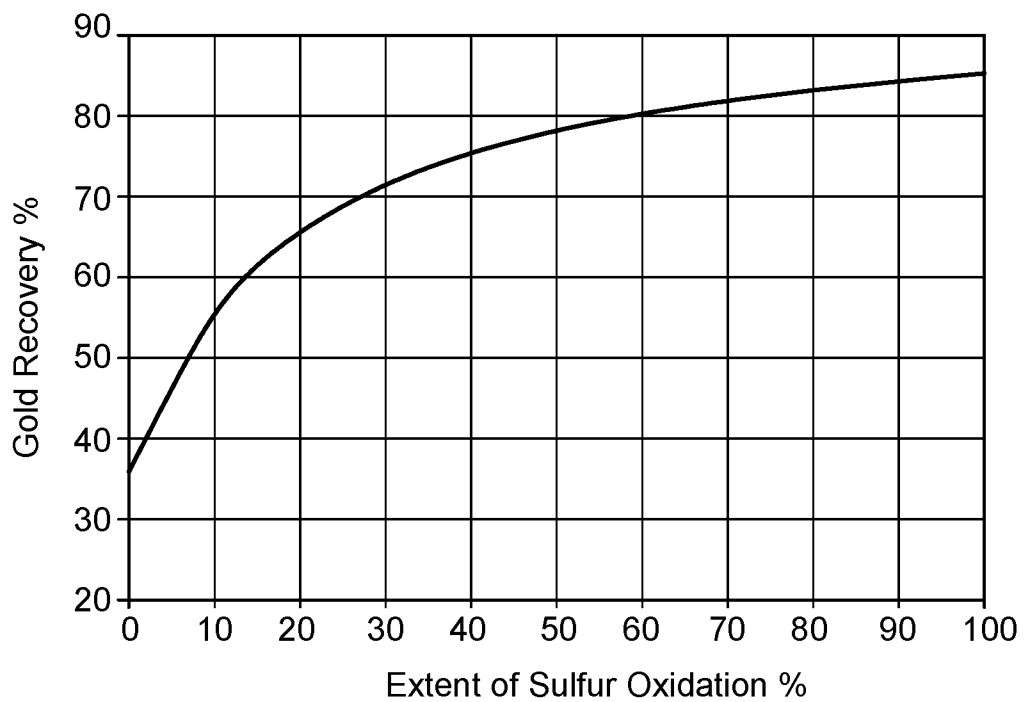


Fig. 7



INTERNATIONAL SEARCH REPORT

International application No
PCT/EP2021/069927

A. CLASSIFICATION OF SUBJECT MATTER
 INV. C22B3/12 B02C17/00 C22B11/00 C22B11/08 C22B11/12
 C22B3/02
 ADD.
 According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED
 Minimum documentation searched (classification system followed by classification symbols)
 C22B B04C B02C

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
 EPO-Internal, WPI Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	WO 2006/064350 A1 (GREENHILLS MINING AND IND SERV [ZA]; SINGH ADRIAN [ZA] ET AL.) 22 June 2006 (2006-06-22)	1,3-9, 11-15
A	page 2, line 1 - page 9, penultimate paragraph; figures 1-3	2
X	EP 1 171 641 A1 (MIM HOLDINGS LTD [AU]) 16 January 2002 (2002-01-16)	1,3-15
A	cited in the application paragraphs [0026] - [0042]; figure 2; example 1	2
X	WO 2016/081799 A1 (SMIDTH AS F L [DK]; CHAIKO DAVID J [US]) 26 May 2016 (2016-05-26)	16-22
	page 23, line 5 - page 33, line 10; figures 1-5,12-15	

Further documents are listed in the continuation of Box C. See patent family annex.

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"O" document referring to an oral disclosure, use, exhibition or other means	"&" document member of the same patent family
"P" document published prior to the international filing date but later than the priority date claimed	

Date of the actual completion of the international search 22 October 2021	Date of mailing of the international search report 02/11/2021
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Name and mailing address of the ISA/ European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Fax: (+31-70) 340-3016	Authorized officer Panitz, J
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INTERNATIONAL SEARCH REPORT

Information on patent family members

International application No PCT/EP2021/069927

Patent document cited in search report	Publication date	Patent family member(s)	Publication date	
WO 2006064350	A1	22-06-2006	AP 2137 A	19-07-2010
			AT 394511 T	15-05-2008
			AU 2005315314 A1	22-06-2006
			BR PI0517194 A	30-09-2008
			CA 2590531 A1	22-06-2006
			CN 101111615 A	23-01-2008
			EA 200701261 A1	28-12-2007
			EG 24577 A	09-11-2009
			EP 1831411 A1	12-09-2007
			ES 2307223 T3	16-11-2008
			NZ 556382 A	25-09-2009
			PL 1831411 T3	31-10-2008
			PT 1831411 E	06-08-2008
			UA 93865 C2	25-03-2011
			US 2009293679 A1	03-12-2009
			WO 2006064350 A1	22-06-2006
			ZA 200705658 B	30-04-2008

EP 1171641	A1	16-01-2002	AR 023666 A1	04-09-2002
			AT 301729 T	15-08-2005
			CA 2349940 A1	30-03-2000
			DE 69926661 T2	08-06-2006
			EP 1171641 A1	16-01-2002
			ES 2255299 T3	16-06-2006
			MY 118964 A	28-02-2005
			NZ 511616 A	25-10-2002
			PA 8482201 A1	14-12-2001
			PE 20001188 A1	14-11-2000
			US 6833021 B1	21-12-2004
			WO 0017407 A1	30-03-2000

WO 2016081799	A1	26-05-2016	AU 2015349677 B2	15-06-2017
			AU 2015349747 B2	15-06-2017
			AU 2017221773 A1	21-09-2017
			AU 2017221775 A1	21-09-2017
			CA 2968245 A1	26-05-2016
			CA 2968256 A1	26-05-2016
			CL 2017001285 A1	15-12-2017
			CL 2017001286 A1	15-12-2017
			CN 107106942 A	29-08-2017
			CN 107109521 A	29-08-2017
			FI 127300 B	15-03-2018
			FI 129125 B	30-07-2021
			FI 129230 B	30-09-2021
			MX 365127 B	24-05-2019
			PE 20170798 A1	04-07-2017
			PE 20170836 A1	04-07-2017
			US 11124856 B1	21-09-2021
			US 2017356063 A1	14-12-2017
			WO 2016081799 A1	26-05-2016
WO 2016081908 A1	26-05-2016			
ZA 201703302 B	18-12-2019			
