Comparative Study between D2EHPA and Cyanex 272 as Extractants of Mn(II) from a Leached Liquor Produced by Reductive Leaching of a Pyrolusite Ore with SO2

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Received June 29, 2012; revised July 27, 2012; accepted August 9, 2012

ABSTRACT
This paper compares the efficiency of D2EHPA and Cyanex 272 to extract the divalent manganese cation from an actual leaching solution obtained by reductive leaching of a low-grade pyrolusite ore, using SO2 gas. The studied variables were the extractant concentration in the organic phase, the pH of the leached liquor and the volume ratio between organic phase and leached liquor. It was observed that D2EHPA is a better extractant than Cyanex 272. Therefore, with the best experimental conditions found, both extractants reach a manganese recovery around 95% with five extraction stages conducted under the following conditions: 25 °C, O/A = 2, 10% volume of extractant concentration in the organic phase, pH of the leached liquor before the mixing between 8 and 8.5, and one minute of mixing time for each extraction stage.

Keywords: Manganese Solvent Extraction; Manganese Leaching; Manganese; Pyrolusite

1. Introduction
In last decades the manganese demand has increased mainly due to the growth of the steelmaking industry. The most important manganese minerals are pyrolusite (MnO2), rhodochrosite (MnCO3), rhodonite (MnSiO3), black manganite (MnO(OH)), and alabandite (MnS) [1].

According to Zhang et al. [2], after the steelmaking application of manganese, the aluminum industry shows an increasing demand of manganese. A third place in manganese consumption is occupied by the electronic industry, which consumes both electrolytic and chemical grade manganese dioxide (EMD and CMD) in the fabrication of disposable and rechargeable batteries. Currently, the growing market of mobile communications has increased significantly the demand of batteries and consequently of EMD. Notably, the batteries industry demands a concentrate of manganese ore containing above 80% of MnO2 with a total iron oxides content below 1%.

Generally, low grade manganese minerals are processed by hydrometallurgical techniques. There are several methods to extract the metal from the leaching solution, such as ionic exchange, solvent extraction with electrowinning and precipitation. However, the solvent extraction with electrowinning is the most appropriate method to obtain a product of higher value.

The elimination of manganese impurities of nickel and zinc from leach liquors of cobalt is the most common objective of manganese extraction studies [3-7]. However, there are few studies on the solvent extraction method to concentrate a manganese leached solution out of a pregnant solution obtained by SO2 leaching of low-grade pyrolusite ores. The scope of this work was to study the effect of pH of the leached solution, type of extractant and its concentration in the organic phase, number of extraction stages and the ratio between the organic and leached liquor, on the manganese recovery.

2. Review of Previous Work
This section presents a brief review of previous studies that used di-(2-ethylhexyl) phosphoric acid (D2EHPA) and bis(2,4,4-trimethylpentyl) phosphinic acid (Cyanex 272), to extract metals of the same group of manganese. One of the first research works on solvent extraction of manganese is that of Yoshida et al. [8], who studied the use of thienyl trifluoracetone as extractant. Their objectives were focused to determine the formula of the manganese-extractant complex and the distribution ratios. To separate manganese from cobalt of a sulfated solution, Hoh et al. [9] carried out an experimental work with D2EHPA, observing that a high degree of separation is obtained when the pH of the solution is 4.2, O/A = 2 for solvent extraction and they claimed that four extraction...
stages were sufficient for a good separation.

Sato and Nakamura [10] found out that the distribution coefficient is dependent on the concentration of the acid and D2EHPA, suggesting that the extraction is carried out through a cation-exchange mechanism.

Hughes and Biswas [11] used a rotating diffusion technique to extract the manganese of an aqueous solution containing ∼0.5 mol/L Mn(II) (pH between 2.70 and 3.57), with an organic solution whose D2EHPA concentration in n-hexane as solvent was ∼1.13 mol/L. They observed the formation of a complex in the aqueous film, close to the organic-aqueous interface, which was identified to be the rate-limiting step.

A series of shake-out tests was undertaken by Cheng [12] to investigate the fundamentals of the separation of manganese from cobalt and nickel using D2EHPA in kerosene. The author concluded that the separation of manganese from cobalt and nickel by D2EHPA in kerosene was affected by temperature and pH. At pH 3.0, better separation of manganese from cobalt and nickel was achieved at room temperature 23°C. Meanwhile at pH 3.5, better separation of manganese from cobalt was achieved.

Cheng et al. [13] had also used solvent extraction to recover manganese from solutions also containing nickel, cobalt and magnesium, concluding that D2EHPA is a convenient reagent for selective manganese extraction. As well, Pagnanellia [14] studied the use of D2EHPA as extractant and n-heptane as a diluent, analyzing the effect of the extractant concentration in the range from 0.05 to 2.5 M, upon the separation of manganese of both a synthetic and an actual solution employing a volume ratio 1:1 of the organic and aqueous phases.

Zhang and Cheng [15] enounced that solvent extraction plays a vital role in purification and separation processes and that D2EHPA is the cheaper and more widely used organic to extract the manganese when it is present as impurity in a leaching solution. Sousa Junior et al. [16] studied the extraction of manganese sulfate of the MnSO₄·H₂SO₄·H₂O-D2EHPA-isoparaffin system making use of a thermodynamic model based on balance equations of mass and charge. They also found that the equilibrium constant of the extraction was 5.18 × 10⁻⁴. Also, they observed that under the experimental conditions used, one mole of hydrogen ion was released for one mole of metal extracted and that the separation factor was pH dependent.

In an ongoing study, Devi et al. [17] used this regent employing kerosene as solvent to separate manganese and cobalt from sulfated solutions. The authors compared three extractants and concluded that the selectivity order for manganese separation was D2EHPA, PC88 and Cyanex 272. On their part, Salgado et al. [18] studied a hydrometallurgical route based on the liquid-liquid extraction technique using Cyanex 272, for the selective separation of metal values, in particular, zinc and manganese, from spent alkaline batteries. They observed that zinc and manganese are easily separated (pH = 2.0) using 20% (v/v) Cyanex 272 dissolved in Escaid 110 at 50°C.

The single-stage extraction of manganese(II) sulfate with Cyanex 302 in kerosene (O/A = 1) was also studied by Devi and Mishra [19]. They established the mechanism of extraction and stoichiometries of metal-containing extracted species. They also observed that the temperature variation studies showed that the extraction process is exothermic, which was confirmed by the negative value of ΔH.

It is interesting to observe that most of the research work of solvent extraction of manganese was conducted making use of synthetic solutions where the metal is present as impurity. On the contrary, the interest of this work is to compare the performance of D2EHPA and Cyanex 72 reagents in kerosene, to extract Mn(II) out of an actual leached solution. Manganese pregnant liquor was obtained by leaching of a low-grade pyrolusite ore using SO₂ as the reductive reagent; the manganese concentration in this liquor was approximately 0.085 M. Thus, the aim of this study is to share the guidelines and the most appropriate operating conditions to obtain an efficient solvent extraction process.

3. Experimental Part

The elemental analysis of the ore was conducted using X-ray fluorescence (% (w/w): Mn (18.80), Fe (12.81), Si (23.83), Al (1.18), Ba (1.11), Ca (0.59) and O (42). Based on this analysis and on the stoichiometry of the oxidised species, a mineralogical composition was estimated (% (w/w): MnO₂ (28.78), Fe₂O₃ (17.72), SiO₂ (49.33), Al₂O₃ (2.15), BaO (1.19) and CaO (0.79). Notably, these results are in qualitative agreement with the results obtained using X-ray diffraction.

The ore was finely ground to reach a particle size distribution ~38 μm. The manganese dissolution was conducted by reductive leaching of the low-grade pyrolusite ore using SO₂ as the reductant. The leaching was carried out in a two-litre glass cylindrical reactor operated in batch mode. The reactor was properly instrumented to measure and control the temperature, pH, mixing rate and gas flow bubbled into the reactor. Figure 1 schematically illustrates the experimental setup. During the leaching of 50 g of ore, the experimental conditions were 60°C for 120 minutes, mixing to 800 rpm, 1600 mL of pulp and 0.096 mL/s of SO₂ gas. After the leaching, the liquor was separated from the solids by filtration.

The leach liquor contained 90% of Mn and 0.17% of the Fe originally contained in the ore. The poor Fe dissolution was caused by the fact that practically all the Fe was occluded into quartz particles. Thus, the manganese concentration in this liquor was around 0.085 M.
In the solvent extraction stage, the extractants were dissolved in kerosene in order to have the following extractant concentrations: 5%, 10%, 15%, 20% and 25% by volume. Considering that the affinity of the extractants for the metal ions is as follow: Fe^{2+}, Fe^{3+} > Zn^{2+} > Cu^{2+} > Co^{2+} > Ni^{2+} > Mn^{2+} > Mg^{2+} > Ca^{2+}, it was necessary to eliminate the Fe impurities of the leaching solution prior to the extracting stage. To precipitate the impurities of Fe, the pH was increased to 7.5 with NH₄OH 5 M. Initially, the leaching solution contained 4932 ppm of Mn and 639 ppm of Fe, whereas after the precipitation of Fe(OH)₂, the iron content dropped to about 0.03 ppm, with only a small fraction of Mn(II) being precipitated as a collateral effect (5.7%).

The mixing of the two phases and solvent extraction separation was carried out in a separation funnel of 125 mL. Each test consists of five separation stages with 60 s of manual agitation, being careful to apply similar agitation intensity in each test. After each extraction stage, the mixture was let to stand still during 20 minutes, permitting the complete separation of the phases. After that, a sample of the solution was withdrawn to measure the manganese concentration by atomic absorption. Before and after each extraction stage the pH of the aqueous phase was measured and eventually adjusted with a NH₄OH 1 M solution.

The initial tests were focused to evaluate the effect of the extraction time (mixing of phases), that is, the kinetics of the extraction. Once the kinetics and the optimal pH were known, the next stage was to determine the best organic/aqueous phase ratio. The effect of the extractant concentration in the organic phase was also studied, testing the following compositions: 5%, 10%, 15%, 20% and 25% by volume. Each test was conducted by duplicate and varying a parameter at the time, while keeping the others constant.

4. Results

4.1. Determination of the Extraction Time

Figure 2 shows the kinetics of manganese extraction achieved in one mixing stage with D2EHPA, using a ratio of organic to leached liquor (O/A) of 1, and 10% of extractant in kerosene. The results indicate that after three minutes, 13.47% of the manganese is recovered (i.e., extracted). A significant change of pH of the leached liquor was observed, which suggests that the reaction is controlled by the activity of the H⁺ ion in the leached liquor, as it is discussed in the next section.

Figure 3 illustrates the effect of the pH of the leached liquor before the extraction, on the manganese recovery. The results show that above pH = 8, the Mn recovery rises exponentially. However, with both extractants, the organic-leached liquor dispersion is unstable above pH = 8.5 and therefore, it is necessary to operate from pH 8 to 8.5. Apparently, this instability is because, according to Pourbaix diagrams, above pH 8.5 manganese ions may precipitate as hydroxide. Notably, from this figure it is observed that in the range of pH between 5 and 8, the manganese extraction with D2EHPA is about seven times more efficient than Cyanex 272.

Figure 4 compares the pH of the leach liquor before and after the mixing stage with the organic, this last is the pH at the equilibrium. Note that in the case of D2EHPA, the change of pH is larger than that of the Cyanex 272 and this behavior may be the reason of the higher extraction potential of D2EHPA and its larger distribution coefficients. This gradient of pH is caused by the release of hydrogen ions by the extractant which are exchanged by the metal cations to form an organometallic compound, thus causing the leach liquor to become acidified. This transference of H⁺ from the organic to the aqueous phase decreases when the pH of the solution approaches the unity, reducing the extraction efficiency.
Figure 3. Effect of the pH on Mn extraction using Cyanex 272 and D2EHPA. 25°C, 10% extractant in the organic phase, O/A = 2 and 1 minute of mixing time.

Figure 4. Gradient of pH before and after each extraction stage. (a) D2EHPA, (b) Cyanex 272. O/A = 0.5, 10% of extractant in the organic phase. Gray bars: pH before extraction; White bars: pH after extraction.

4.3. Effect of the Organic/Leached Liquor (O/A) Ratio

Figure 5 shows the effect of the organic/leached liquor (O/A) ratio on the kinetics of the extraction. Results indicate that the Mn recovery increases with the increase of the O/A ratio and the time of reaction, with an organic phase composed by 10% of extractant and 90% of kerosene. Evidently, when the organic phase is more concentrated in extractant, the extractant molecules have more probabilities of reacting with the Mn(II), increasing its recovery. In both cases, with D2EHPA and Cyanex 272, the pH of the leached liquor was adjusted at a pH value between 8 and 8.5. Note that contrary to the results obtained with D2EHPA, the results obtained with Cyanex 272 show a more irregular behavior, which is due to difficulties in regulating the pH of the leached liquor.

4.4. Effect of the Extractant Concentration in the Organic Phase

With respect to the effect of the extractant concentration in the solvent, the manganese recovery increases as the
extractant concentration is increased, as shown in Figure 6 (a) for D2EHPA and (b) for Cyanex 272). It is evident that to extract more manganese it is necessary to have enough extractant, otherwise the absence of one of the reagents that participate in the reaction limits the extent of it. However, a high extractant concentration increases the viscosity and decreases the fluidity of the organic phase, reducing the mass transfer rate during the extraction. To avoid these problems, it is necessary to optimize the extractant concentration, without affecting the extraction extent.

4.5. Distribution Coefficients

A common way to evaluate the efficiency in solvent extraction separation is by means of the distribution coefficient (D), which is defined as the ratio of the concentration of the extracted metal in the organic and the concentration in the aqueous phase, in equilibrium with the organic.

Figure 7 compares the distribution coefficients of both extractants, showing that D2EHPA values are larger than those of Cyanex 272. These results reveal the superiority of D2EHPA as manganese extractant over Cyanex 272.

5. Conclusions

D2EHPA is more efficient than Cyanex 272 to extract the Mn(II) from a leach liquor obtained from the reductive leaching of a pyrolusite ore with SO₂. Notably, it is sufficient two or three minutes of intimate contact between organic and aqueous phases to obtain the 100% extraction, i.e., to complete each extraction stage.

Manganese recovery increases as the extractant concentration in kerosene solvent increases. It is demonstrated that using D2EHPA, a ratio of organic/aqueous = 2 is adequate to achieve a manganese recovery of 95% in five extraction stages.

To obtain an efficient solvent extraction, the pH of the leached liquor should be adjusted between 8 and 8.5, although it is suggested a value closer to 8.5. At pH above this value, the organic-leached liquor dispersion becomes unstable affecting the extraction. Apparently this is because at pH higher than 8.5 the Mn(II) is likely to form Mn(OH)₂.

The removal of Fe impurities from the leaching solution was conducted by alkaline precipitation and filtering of the Fe(OH)₂. This cleaning stage does not increase significantly the operating costs because increasing the pH is the first step before the manganese solvent extraction.

6. Acknowledgements

The authors are grateful to CONACYT (Mexico) and to Huajicori S.A. de C.V. for the scholarship and financing
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