Introduction

The Skorpion process was developed to treat a complex non-sulfide oxide orebody using atmospheric leaching followed by solvent extraction (SX) and electrowinning of zinc. The major mode of impurity transfer was attributed to crud runs in the solvent extraction circuit. Crud may be described as a heterogeneous, partly inorganic, partly organic mass of material which accumulates predominantly at the interface between organic and aqueous solutions. The paper describes the root causes of the crud formation in the solvent extraction circuit and optimization strategies which are being followed to prevent impurity excursions in electrowinning.

Keywords: zinc oxide, solvent extraction, crud formation, colloidal silica.

Process chemistry

Leaching

Leaching of zinc oxide ores by sulphuric acid is generally described in terms of the following reaction equations:

\[ \text{ZnO} \cdot \text{SiO}_2 + \text{H}_2\text{SO}_4 + \text{H}_2\text{O} = \text{ZnSO}_4 + \text{H}_4\text{SiO}_4 \]  \[ [1] \]

\[ \text{Zn}_n\text{SiO}_4 + 2\text{H}_2\text{SO}_4 = 2\text{ZnSO}_4 + \text{Si(OH)}_4 \]  \[ [2] \]

\[ \text{Zn}_n\text{Si}_2\text{O}_5(\text{OH})_2 \cdot \text{H}_2\text{O} + 4\text{H}_2\text{SO}_4 + 4\text{ZnSO}_4 + 2\text{Si(OH)}_4 + 2\text{H}_2\text{O} \]  \[ [3] \]

Leaching is carried out in a series of agitated tanks at about 50°C. The final pH is maintained between 1.8–2.0 to maximize stability of colloidal silica. After leaching the pH is raised to 4.0 to maximize agglomeration and the formation of precipitate.

Dissolved siliceous species, namely, Si(OH)_4 (monosilicic acid) remain in solution under conditions that ensure their solubility is not exceeded. Depending on pH, temperature and dissolved salts concentrations, siliceous species will tend to polymerize to form colloidal silica. These siliceous polymers, sized from 1 nm to 1000 nm, have a massive surface area, but are still small enough to be unaffected by gravity. Colloidal-sized, polysilicic molecules can cause serious problems in a solvent extraction (SX) system.
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by stabilizing emulsions. The formation of colloidal silica is irreversible and follows a dehydration process:

\[ nSi(OH)_4 \rightarrow (SiO(OH))_n H_{2}O + (n-1)H_{2}O \]  \[4\]

**Solvent extraction**

During solvent extraction the organic extractant used is 40% v:v di-2-(ethylhexyl) phosphoric acid (D2EHPA, symbolized as RH) diluted in the solvent ESCAID 100. The extraction of zinc is by cation exchange:

\[ Zn^{2+ \text{(aq)}} + 2RH_{\text{(org)}} = 2H^{+ \text{(aq)}} + ZnR_{2\text{(org)}} \]  \[5\]

The organic phase is stripped of zinc using spent electrolyte:

\[ ZnR_{2\text{(org)}} + 2H^{+ \text{(aq)}} = Zn^{2+ \text{(aq)}} + 2RH_{\text{(org)}} \]  \[6\]

The extraction versus pH curves provide an indication of the selectivity of the organic extractant under different conditions. A typical E-pH curve measured is shown in Figure 1. From reaction (5) it is seen that the acidity of the aqueous phase increases as the extraction proceeds, with two protons released for each metal ion extracted. For typical Skorpion operating conditions of 30 g/l Zn in the PLS and 10 g/l Zn in the raffinate, this corresponds to approximately 30 g/l H\textsubscript{2}SO\textsubscript{4} being generated across the extraction circuit. The general order of selectivity with increasing pH in this system is:

\[ Fe\textsubscript{III} < Zn < Cu < Al\textsubscript{III} < Mn < Cd < Cu < Mg < Co < Ni. \]

**Electrowinning**

The last step in the Skorpion process is the electrodeposition of zinc onto aluminium cathodes from the loaded electrolyte:

\[ 2Zn^{2+} + 2H_{2}O = Zn + 4H^{+} + O_{2} \]  \[7\]

The produced spent electrolyte is sent back to the stripping stage of SX.

**Plant description**

A simplified Skorpion flow sheet of the process is given in Figure 2. A detailed discussion of the circuit is given in.

Following an atmospheric sulfuric acid leach, iron, aluminium and silica are precipitated from the pregnant leach solution in the neutralization section. This neutralized and clarified pregnant liquor solution (PLS) is sent to the SX circuit (Figure 3). Extraction is performed over three stages operating at 40°C with an O:A ratio between 1 to 1.5. Zinc is transferred to an organic phase that contains 40% v:v D2EHPA diluted in kerosene. The acidic raffinate is sent back to leaching. The first two stages of the washing circuit use demineralized water to wash out physically entrained impurity species diluted. Spent electrolyte is employed as a scrub liquor in the third wash stage to ‘crowd off’ co-extracted impurity species from the loaded. The loaded organic phase is stripped of its zinc content by contact with spent electrolyte. Following electrowinning, the end product is SHG zinc cathodes (99.995% Zn).

Iron and aluminium are also extracted in preference to zinc, but cannot be easily stripped from the organic phase. For this reason, a small portion of the stripped organic is bled for regeneration. In the regeneration step, the organic is brought into contact with a strong hydrochloric solution (6 mol/l HCl) to remove all the iron. The resulting iron chloride solution is neutralized and returned to the leaching section.

The operating and engineering availability of the refinery was significantly increased by decoupling the electrowinning and solvent extraction sections as well as the SX and leach sections to developed additional capacity. An extra thickener was installed to ensure flexibility and low solids transfer into SX.

**Defining the problem**

After more than 15 months of stable operation, the refinery was forced into an unscheduled shut-down in August 2006 by a significant excursion of metal impurities (Cu and Ni) into the electrowinning circuit. After the refinery had restarted, intermittent impurity excursions across SX continued to hinder the performance of the electrowinning circuit and the output of the refinery (Figure 4). A salient feature of Figure 1 is that the intermittent impurity excursions appeared as well-defined spikes against otherwise steady background levels rather than as an overall increase in the impurity background, suggesting a physical rather than a chemical transfer mechanism.

Transfer of PLS across SX may either occur by aqueous
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Figure 2—Simplified process flow sheet of the Skorpion Zinc refinery

Figure 3—Schematic representation Skorpion SX circuit

Figure 4—Evolution of impurities concentrations in loaded electrolyte (1 July - 18 November 2006)
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Entrainment and/or crud, or another mechanism involving some degradation of the extractant or diluent. The contributions of these mechanisms are discussed below.

Effect of aqueous entrainment

Routine aqueous entrainment measurements averaged between 100 and 200 mg/l for the organic and aqueous extraction and washing stages. Mass balance calculations showed that these levels were too low to have caused the observed impurity levels in the loaded electrolyte.

Organic health

Standard shake-out tests on the organic phase confirmed that chemical extraction of metal impurities in this phase was within design limits, excluding chemical transfer as a primary transfer mechanism. Also, co-extraction was not likely to occur in the form of sporadic spikes. Sensitive Fourier Transform infrared spectroscopy (FTIR) and nuclear magnetic Resonance (NMR) spectroscopic analysis of the organic phase could not detect any significant degradation products.

The viscosity of the organic phase was, however, moderately elevated, and minor changes in phase disengagement times were apparent. These alterations were suggestive of a gradual change in the physical properties of the organic phase, but were not expected to impact significantly on the hydraulic behaviour. Detailed chemical analysis showed a build-up of elements such as scandium (100 mg/l) and yttrium (100 mg/l) on the extractant. It is well known that trivalent species such as Fe³⁺ can polymerize D2EHPA and change the viscosity of the organic phase. In the Skorpion Zinc flowsheet a bleed of the organic is treated with 6 m HCl to maintain the organic-phase iron loading at 300 mg/l. Subsequent to the detection of Sc and Y, the runtime on the HCl plant was changed to ensure removal of these species from the organic phase.

Crud transfer

Crud in hydrometallurgical SX can be defined as stable, non-coalescing deposits occurring within the settler of an SX operation. They originate from semi-stable aqueous-organic emulsions formed in SX settlers7,8,9.

The formation of emulsions in the extraction stages was observed prior to and during the metal impurity excursions, and was accompanied by unusually high levels of mobile floating crud and an accumulation of large quantities of bottom crud in the extraction settlers. Elemental analysis of the floating crud indicated the presence of significant amounts of Si (9.85%), Zn (4.1%), Al (1.79%), Ni (0.03%) and Cu (0.06%). The moisture content was 37%. The crud therefore contained a considerable amount of entrained PLS. The most likely cause of impurity transfer was thus the sporadic transfer of floating crud (and with it entrained PLS) from the extraction stages across the washing circuit into the stripping circuit. If intermittent crud runs are the main impurity transfer mechanism, the observed spikes in nickel and copper in the loaded electrolyte can be explained.

Causes of crud formation

The precursor to crud formation is the development of an emulsion10. The emulsion becomes a crud through repeated mixing and continuous inclusion of solids and colloids. Once crud is formed, its dispersion and separation are difficult to achieve under operating conditions. It is generally accepted that the factors contributing to crud formation are the composition and components of the pregnant liquor stream (PLS), the components or degradation products of the organic phase, and the operating equipment or operating conditions within the circuit10. Available information on other systems indicates that the components of pregnant liquor stream with significant influence on crud formation are fine solids, silica, manganese and foreign organic matter10. Considering the chemical composition of the crud, namely Si and Al, it was mooted that crud formation was linked to solids carryover and the presence of colloidal silica. Solids may enter the solvent extraction (SX) circuit with the aqueous feed or precipitate during the process, while colloidal-sized, polysilicic molecules can cause serious problems in a SX system by stabilizing emulsions.

The solids’ content entering SX is monitored by routine measurement of total suspended solids (TSS). The frequency distributions of the TSS values during and after the impurity excursions are shown in Figure 5. The average TSS value during the excursion was 52 mg/l compared with 39 mg/l after the impurity excursions, when a significant fraction (34%) of the TSS measurements was less than 23 mg/l. This must be compared with the targeted value of <10 mg/l to minimize crud formation in SX.

An important observation made during the impurity excursion was a noticeable change in the physical appearance of the crud, from coarse (bottom crud) to voluminous, highly gelatinous type crud (floating crud). In the literature this type of crud has often been associated with fine clay and silica10. The predominance of Si among the contaminants in the floating crud strongly suggested that it was the main contributory factor in the crud formation. Analyses of the crud suggested that it was entering the circuit in at least two different forms: soluble species (colloidal silica) and suspended solids.

Crud transfer was intrinsically linked with nickel excursions in the electrolyte. Bearing in mind that D2EHPA is not selective towards extraction of nickel under the operating conditions. It is generally accepted that the factors contributing to crud formation are the composition and components of the pregnant liquor stream (PLS), the components or degradation products of the organic phase, and the operating equipment or operating conditions within the circuit10. Available information on other systems indicates that the components of pregnant liquor stream with significant influence on crud formation are fine solids, silica, manganese and foreign organic matter10. Considering the chemical composition of the crud, namely Si and Al, it was mooted that crud formation was linked to solids carryover and the presence of colloidal silica. Solids may enter the solvent extraction (SX) circuit with the aqueous feed or precipitate during the process, while colloidal-sized, polysilicic molecules can cause serious problems in a SX system by stabilizing emulsions.
conditions (Figure 1) and that no evidence of organic degradation could be determined, nickel was used in the analysis below as a tracer in crud transfer.

Comparisons of the total silicon concentration during and after the excursion are shown in Figure 6. There was no significant change in the mean of the concentration (60 and 63 mg/l prior to and after the excursion), suggesting that the form of silicon (colloidal or suspended) had a more significant impact on crud formation than the actual concentration. This was also confirmed in laboratory testwork:

Dissolved silica had no notable effect on either the phase disengagement (PD) time or interfacial characteristics. However, colloidal silica (130 mg/L) led to a slightly longer PD time and persistent cloudiness of the aqueous phase, including the interfacial region. It is reasonable to suggest that this could eventually lead to emulsification, a known precursor of crud formation. These analyses clearly demonstrated that while dissolved silica are not a major contribution to crud formation, colloidal silica are contributory.

As already indicated, low pH (<2.0) in the leach section is expected to reduce the selectivity of the leaching process and increase the proportion of amorphous silicates and impurities in the PLS. Considering this, a decision was therefore made to control the pH at 2.0 in order to maximize the stability of colloidal silica.

The frequency distributions of the pH values in the last leach reactor recorded for the periods during and after the impurity excursions are shown in Figure 7. The period after the excursions refers to the time when no impurity excursions were observed and the pH was controlled to 2.0. The average pH in neutralization over the same periods remained the same at 4.2, although the variance decreased after optimization of the pH control.

The significant impact of the leach pH on Ni concentrations in the electrolyte is shown in Figure 8. At the low leach pH, Ni in the electrolyte averaged 120 ppb compared with 80 ppb after the pH control was optimized at pH 2.0. Bearing in mind that the targeted value in electrowinning is 80 ppb Ni, unacceptably high levels of Ni were detected during the impurity excursions. Subsequent pH optimization resulted in far better Ni control and no impurity excursions have been observed since.

Conclusions

The major mechanism of impurity transfer across the SX circuit was attributed to intermittent episodes of aqueous entrainment that were associated with the build-up and release of crud in the extraction stages and transfer to the electrowinning circuit. The predominance of Si among the contaminants in the floating crud left very little doubt that it
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was the main cause of crud formation. Low pH (<2.0) in the leach section reduced the selectivity of the leaching process and increased the proportion of colloidal silica and impurities in the PLS. After adjusting and controlling the leach pH at 2.0, crud transfer episodes have reduced significantly. However, the current TSS values are still above the target of 10 mg/L. Skorpion Zinc is currently evaluating a number of solid-liquid separation technologies, such as pinned bed clarification, to achieve this.

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References