

## **SART for Copper Control in Cyanide Heap Leaching**

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### **ABSTRACT**

Copper cyanide is a common component of cyanide-treatable precious metal ores. Copper level in production heaps can be predicted from laboratory column tests, but the exact correlation is not necessarily intuitive. Generally heap leach operators like to keep copper levels in solution below 300-500 ppm and may note problems with gold recovery and cyanide consumption when copper concentrations exceed this amount. There are several methods of copper removal from cyanide solutions including ion-exchange; direct electrowinning; Acidification, Volatilization and Recovery AVR); and sulfide precipitation such as the Sulfidization, Acidification, Recycling, and Thickening process (SART). SART involves acidification with addition of soluble sulfide, separation of the resulting copper sulfide precipitate, and addition of lime to reestablish alkalinity prior to returning the solution to the leaching process, recovering both copper and cyanide as valuable products. In principle SART is very simple. Yet some SART plants which have been built may have been unnecessarily complex. This paper explores the basics of SART and makes the case for a simple plant design as applied to the heap leaching circuit.

### **1. Introduction**

Many current precious-metal heap leach operations do not require control of copper in the leach solution. Historically many copper-laden precious metal deposits were dismissed as economically and technically unattractive but with the significant increase in precious metal prices in recent years, it is now more common to consider such deposits. Consequently, there are now several projects which have installed systems to control copper levels in the cyanide leach process. The most common of these is the SART process. Without SART or some other system of copper removal, field process solutions will stabilize at substantially higher copper content than laboratory tests would indicate, leading to a variety of economic issues.

## **2. Issues with Copper in Precious Metal Ores**

The presence of cyanide-leachable copper in a large enough amount in a gold-bearing ore can be significantly detrimental to the economics of a gold project for several reasons:

- Copper, when dissolved with adequate free cyanide at the typical heap-leach/mill operating pH of 10-11, predominantly forms the  $\text{Cu}(\text{CN})_3^{2-}$  complex, and will bind at least 2.3 kg of sodium cyanide for every kg Cu leached. This use of cyanide reduces gold leaching rate and represents a significant consumption and inventory of cyanide in the leach circuit. The strategy of using a very low level of free cyanide may result in selective leaching of gold from some ores. However, if free cyanide is low enough or non-existent, the  $\text{Cu}(\text{CN})_2^-$  complex or insoluble  $\text{CuCN}$  may form, and gold may not leach at all.
- Copper affects accurate free/ or gold leachable cyanide concentration analysis, complicating accurate cyanide dosing and cyanide control.
- Copper competes with gold for adsorption on activated carbon in the normal adsorption/stripping circuit, particularly at low cyanide concentrations relative to copper (e.g. CN/Cu ratios of <4) [Fleming and Nicol 1984]. This can effectively reduce the gold loading capacity of the carbon, increasing the plant size and carbon inventory and thus the cost of the adsorption circuit.
- Copper-loaded carbon can also result in significant copper reporting to dorè bars along with gold and silver, increasing refining costs.
- The copper-bound cyanide inventory in the heap can create environmental issues with heap closure that may require more extensive washing and possibly even cyanide destruction treatment, adding significant back-end cost to the project.

Nearly all copper oxide minerals show significant solubility in cyanide [Marsden and House 1992, Hedley and Tabachnick 1958]. Many sulfide copper minerals also show significant solubility, although less so than with the oxides. The overall cyanide soluble copper fraction of a particular ore is best estimated through bottle roll and/or column test work, as the exact mineralogical composition cannot be conveniently determined.

In an operating cyanide heap leach, copper slowly leaches from old ore on the heap, and unchecked can build up in solution to levels that begin to affect gold recovery and cyanide consumption, in turn affecting project economics. While each operation varies, a good general guideline is that if lab work indicates that field process solutions will stabilize higher than 500 ppm copper, gold recovery might be affected and copper control should be considered. Without some form of copper removal or treatment, the copper concentration in a typical heap leach process solution (grams copper per litre) as a rule of thumb will build up to between three and six times the amount of leached copper (in grams per tonne ore) as determined in long-term laboratory column leach tests. This means that low-grade gold ores (especially those below 1 g/tonne Au) showing as little as 100 g/tonne leachable copper might be in need of copper treatment.

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The problem of slow continuing copper leaching is compounded when multiple lifts are placed on the heap (as is usually the case). Lower lifts contribute to the copper load and consume free cyanide. It is possible to install impermeable liners on top of old lifts, but this is expensive, technically difficult, and usually results in reduced overall gold recovery.

The additional cost of cyanide (assuming \$2.50/kg) attributed to dissolving copper (at an average 2.5 kg NaCN/kg Cu) at a level of 200 ppm, for example, would be \$1.25 per tonne ore. To look at this another way, if this 200 ppm cyanide-leachable copper was present in an ore with a recovery of 0.5 g/tonne gold, the cash-cost of the copper-bound cyanide consumption alone would be \$78/oz Au (it takes about 62 tonnes of ore to produce one ounce of gold at this recovery).

This illustrated cost does not include the cost of cyanide destruction, which if no copper/cyanide recovery treatment is proposed, may be required for some projects at high copper concentrations. As a rule of thumb the cost of cyanide destruction is roughly equal to the cost of purchased cyanide, so the above costs would effectively be doubled if cyanide destruction is required.

For proper heap design for cyanide leaching of copper-bearing gold ores, it is necessary to run long term column tests (60 to 180 days) at two or three different cyanide and/or pH levels. The relationships between gold-silver-copper recoveries, cyanide levels, and leach times will be different for each ore body. Since gold-copper types of ore bodies tend to be emplaced in large acidic volcanic systems, ore characteristics can be variable and more than one sample may need to be tested. It is important that the test program be comprehensive and defined early in the project evaluation process.

### **3. General Methods of Copper Removal from Cyanide Solutions**

Several methods have been proposed to treat copper in precious metal cyanide solutions, detailed descriptions of which can be found elsewhere [Marsden and House 1992, Botz and Acar, MacPhail and Fleming 1998, Fleming 1995, Barter 2001, Ford 2008, Guzman, Briggs and Kidby 1990]. Some of the more established or piloted methods include the Acidification-Volatilization-Recovery (AVR) method, electrowinning, and sulfide precipitation (e.g. MNR and SART). Ion-exchange processes (e.g. AuGMENT and Vitrokele) can be used as pre-concentration steps in combination with these recovery methods. Table 1 below provides a brief description of these processes.

**Table 1. Some piloted or established methods of recovering copper and/or cyanide from copper cyanide streams.**

Process Name	State of Technology	Process Description
<i>AVR (Acidification, Volatilization, Recovery)</i>	Commercial	<p>Influent is acidified to pH &lt;2 in aeration tower to dissociate free, WAD, and strong cyanide complexes to metal ions and liberate gaseous HCN; HCN recovered as concentrated NaCN in adsorption tower by contact with caustic/lime; Metal ions removed as hydroxide precipitates.</p> <p><b>Pros:</b> Can treat solutions or pulps (Cyanisorb); Recovers concentrated cyanide, amenable to both mill and heap leach makeup; Removes metals from solution, including specific toxics such as arsenic, antimony, and also soluble sulfur species.</p> <p><b>Cons:</b> Relatively high capital costs, potential issues with gypsum scaling in adsorption tower;</p>
<i>Acidification, Copper Cyanide Precipitation</i>	Piloted	<p>Influent acidified to pH &lt; 3 to remove Cu as CuCN. CuCN filtered and collected, HCN regenerated as NaCN by adding lime or caustic.</p> <p><b>Pros:</b> Simple; high copper removal is possible; partial cyanide recovered.</p> <p><b>Cons:</b> CuCN difficult to dewater/filter, and product likely difficult to sell; maximum possible CN recovery only about 65%; acid and Lime/caustic consumption high.</p>
<i>Sulfide Precipitation: 1) MNR Process 2) SART (Sulfidation, Acidification, Recycling, Thickening).</i>	Commercial	<p>Influents treated with acid and chemical sulphide to precipitate copper sulphide, followed by 1) Direct filtration or collection of precipitate in MNR process or 2) thickening and then filtration of precipitate in SART. In both processes, the effluent is reacted with lime or caustic to regenerate cyanide.</p> <p><b>Pros:</b> SART demonstrated commercially, simple process, relatively low capital and operating costs; Recovers copper and also silver (if present) as a salable product, and also recovers cyanide (&gt;95% is possible). Effluent can be returned directly to heap leaching process.</p> <p><b>Cons:</b> Can only treat clean solutions; NaSH reagent fairly expensive. Copper product may have variable value depending on purity and composition as sulfide ion can precipitate other metals (e.g. Zn, Pb) and contain less desirable anions (e.g. CN, SCN).</p>
<i>Electrowinning</i>	Piloted	<p>DuPont process - copper bearing solution electrowon in divided cell to produce copper metal and liberate free cyanide at the cathode; ion-selective membrane prevents CN oxidation at anode</p> <p><b>Pros:</b> Simple process; copper is recovered as high-value metal and cyanide is recovered.</p> <p><b>Cons:</b> Poor efficiency at low metal concentrations, therefore applicability is limited for most heap leach solutions without a pre-concentration step. For high-extraction rates of copper typically a secondary scavenging process for copper is required.</p>
<i>Ion-Exchange 1) AuGMENT®, 2) Vitrokele</i>	Piloted	<p>Pre-concentration processes for copper - adsorption of WAD/free cyanides onto strong-base resin for pre-concentration of copper cyanide, followed by elution and metal recovery; 1) AuGMENT uses commercially available resin with a combination of AVR and Electrowinning process to recover copper and cyanide; 2) Vitrokele uses a proprietary resin in combination with AVR (and MNR/SART is feasible)</p> <p><b>Pros:</b> 1) high value copper product, plus CN recovered, both at high recovery rates; 2) metal and cyanide can be recovered at high rates, although metals may not be salable via AVR</p> <p><b>Cons:</b> 1, 2) relatively complex processes, related to complex elution-regeneration profiles required for cost-effective operation</p>

Of the above processes, SART is most commonly encountered in operations, and will be discussed in the following sections.

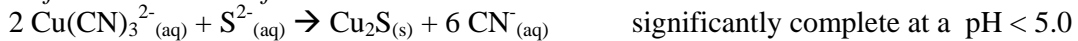
## 4. SART Process for Copper Removal

### 4.1. General Process

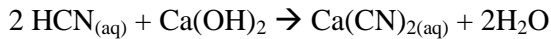
The SART process recovers copper (and other metals such as silver and zinc) as a sulfide precipitate, separates the precipitate from solution, and recovers cyanide through re-neutralization of the effluent. The neutralized solution is recycled to the leaching process.

SART process chemistry is in essence simple and is summarized by the following reactions:

*Sulfidation and Acidification:*



*Neutralization:*



The conventional SART process flowsheet is presented in Figure 1. It is thought by the authors that this traditional approach has resulted in the construction of some SART plants which may have been more complex or expensive than is necessary. The ‘‘SART LITE’’ flowsheet is presented as a possible alternative to the conventional SART circuit in Figure 2. The properly engineered flowsheet needs to fit each specific project so it is recognized that some combination of the two flowsheets might be appropriate in some cases.

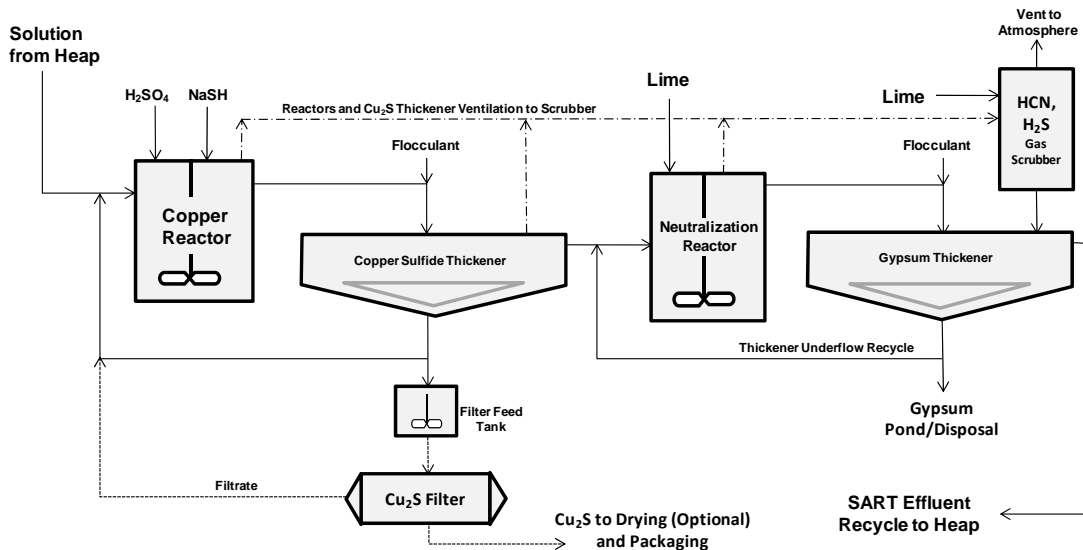


Figure 1. Conventional SART Process Flowsheet.

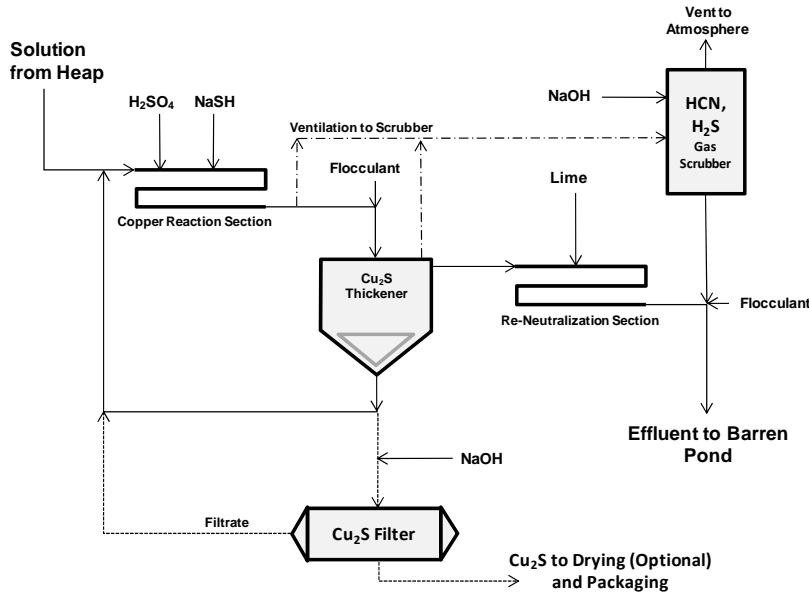


Figure 2. Modified “SART LITE” Process Flowsheet.

#### 4.2. SART Plant Unit Operations

This section will examine some aspects of SART plant design and suggest areas where the design might be simplified and the capital costs reduced. Where equipment design or flow rate examples are used for illustration, we use the basis of a 20000 tonne/day heap leach containing 200 ppm copper cyanide and 100 ppm free cyanide, treated 100% through SART.

##### *Copper Reactor.*

Heap leach solution is typically at a pH of 9.5-11.0. Incoming solution is reacted in a tank to which sulphuric acid is added to maintain a pH of 4.0 to 4.5. NaSH is added to precipitate copper mostly as chalcocite ( $\text{Cu}_2\text{S}$ ). The stoichiometric requirement for NaSH is 0.44 kg NaSH/kg Cu in the effluent (or 0.09 kg NaSH per  $\text{m}^3$  in our example). Typically a slight excess of NaSH is applied to account for other consumers (e.g. silver, oxygen, zinc). Under controlled conditions typical for the SART process gold will not precipitate, but a large percentage of silver will co-precipitate as silver sulfide with the copper sulfide.

Lab tests were run to evaluate the “copper reactor” activities and indicated the following:

- It does not seem to matter to the reaction or to the physical nature of the precipitate if the NaSH is added to an alkaline solution which is then acidified, or vice versa;
- With effective mixing, the reactions take place within a few seconds once the appropriate pH is achieved.
- NaSH reagent is expensive and excess NaSH reacts with cyanide, so the best practice is to use about 95% of the required NaSH and accept a small amount of copper recycle to process.

*Matthew Stewart, Daniel Kappes*

- More dilute Cu feeds produce more finely divided precipitate with poorer settling properties.

In the SART LITE flowsheet, the copper reaction tank has been eliminated. The reactions can take place in the pipe between the process feed pump and the copper thickener, where the large volume of solution will serve to smooth out any concentration gradients.

#### *Copper Sulfide Thickener.*

The copper sulfide formed is very fine and forms very quickly. Recycle of “seed” crystals does not appear to result in larger crystal growth. Recycle of thickener sludge along with flocculent and/or coagulant addition does result in the formation of flocs which settle at least ten times the rate of the unconditioned raw precipitate at dilute copper concentrations (e.g. <200 ppm Cu). The settling issue of unconditioned raw precipitate, and thus the need for recycle and proper flocculent addition, becomes increasingly significant with decreasing Cu concentration in the feed. In general, underflow recycle and flocculent addition are critical design features necessary for economical thickener design.

The operating conditions of the copper sulfide thickener are different from those of a typical mineral plant thickener where ore slurries are thickened. In the SART copper sulfide thickener, the slurry contains only 0.1 to 1.0% solids (including the recycled copper sludge). The incoming stream can be introduced above the settled bed, and separates quickly from the “densified” solids. A small-diameter, tall thickener (high density thickener) will likely be more economic than a traditional thickener, and a design based on de-entrainment to create a thin clear overflow will be more economical than a “traditional” design based on solids settling velocity. This is important because this thickener must be constructed of corrosion-resistant materials, and must be covered to capture HCN. The deep cone of a high rate thickener is also important for inventory control and densification of the small amount of copper sulfide produced.

Therefore in the SART LITE flowsheet we have replaced the conventional thickener with a small diameter, high-density thickener.

#### *Copper Sulfide Neutralization.*

A relatively small volume of copper sulfide slurry comes from the bottom of the thickener about 6 - 7 m<sup>3</sup> per day of a 40% solids slurry from a heap leach processing 20,000 tonnes of ore per day, for example. The slurry should be conditioned (made alkaline to pH 10) with the addition of lime or caustic on its way to the product filters. This is an important step since there is always free or combined cyanide in the precipitate, and filter operations usually result in discharge of air as the cake is dried. If the cake was left acidic, the air would contain dangerously high levels of HCN. The neutralization reaction is rapid and can be done in the pipe leading from the thickeners to the filters. NaOH is the preferred neutralization reagent, since lime would introduce a sulfate precipitate, thus lowering the value of the concentrate. There is no need for a separate “filter feed tank”, since the thickeners provide a large reservoir of thickened sulfide pulp.

*Lime Neutralization Reactor.*

In the traditional SART circuit, a stirred tank is included in which lime is added to the acidic copper-free solution. This is a traditional mineral industry approach, but a tank may not be needed in all cases. Modern in-line pH controls and peristaltic metering pumps permit accurate addition and mixing to ensure the slurry is at a pH of 10.5 or higher. The reaction is rapid and can be done in a pipeline. Therefore in the SART LITE flowsheet we have eliminated this tank. In our example about 0.50 kg lime per m<sup>3</sup> is required for neutralization.

*Gypsum Thickener.*

In traditional SART plants a gypsum thickener is included in the flowsheet to account for the fact that some gypsum may form during the neutralization process (as both calcium and sulphate ions are introduced by acid and lime additions). The gypsum thickener has been eliminated from the SART LITE flowsheet, and although it may be appropriate to include in some flowsheets, its inclusion should not be automatically assumed. The level of lime added following copper recovery is not enough to create a saturated solution of gypsum. While in most cases the solutions are already saturated, the gypsum supersaturates very easily and comes out of solution very slowly. One question to ask is: will enough of the gypsum “behave itself” and stay in the thickener, or does the thickener simply provide a random place where some of the gypsum collects? A more appropriate place to collect gypsum may be in the pregnant or barren ponds, where it has time to crystallize and settle. The ponds are normally quite large and the volume of gypsum created over several years of operation can be collected there without a significant pond capacity increase.

Flowsheets which may permit elimination of the gypsum thickener include:

- a) Intercept pregnant solution off the heap in a tank, process this solution through SART, and discharge to the pregnant pond. Gypsum will settle out along with the normal sludge which collects in the pond, and the clear solution can be processed through carbon columns (or Merrill Crowe) as usual for gold extraction;
- b) Process pregnant solution via SART on its way to the carbon columns, use anti-scalant in large enough dosages to prevent gypsum precipitation onto the carbon;
- c) Process barren solution after carbon columns via SART and use the barren pond as the settling reservoir for the gypsum;

and, the most interesting but (so far) least evaluated -

- d) Process pregnant solution through SART, but leave it acidic as it goes through the carbon columns. Then raise the solution pH to the alkaline side and discharge it to the barren pond. Carbon gold loading is known to be significantly higher in acidic than in alkaline solutions. Additional research is needed to validate the overall practical loading/stripping process in such a condition.



*Matthew Stewart, Daniel Kappes*

*Gas Scrubbing & Plant Safety.*

Both SART and SART LITE contain a gas scrubber. It is a common misconception that HCN will rapidly gas out of an acidified cyanide solution, but in fact hydrogen cyanide is infinitely soluble in water. Therefore, the tank vent system and the scrubber need to be designed only to ensure a slight negative pressure in the appropriate vessels. In the case of SART the scrubber is venting two reaction tanks and one thickener, whereas in SART LITE it is venting only the (smaller diameter) thickener. With proper design of the thickener, the volume of exhaust air needed for scrubbing HCN is very small (probably 200 CFM, or 400 cu m/hr). A small vent fan and small packed scrubber (1800 mm diameter, 5000 mm high) will provide adequate security for our example SART LITE plant.

It is important to not trivialize the dangers posed by an acidified cyanide solution. Regardless of the amount of ventilation, enclosed freeboard spaces in all vessels should be considered very hazardous. The plant (or the operators) should be fitted with HCN monitors with alarms, and procedures for loss-of-power events and maintenance events need to be rigorously designed and implemented.

*Control of Reagent Feeds.*

The major areas of chemical control within the SART plant are:

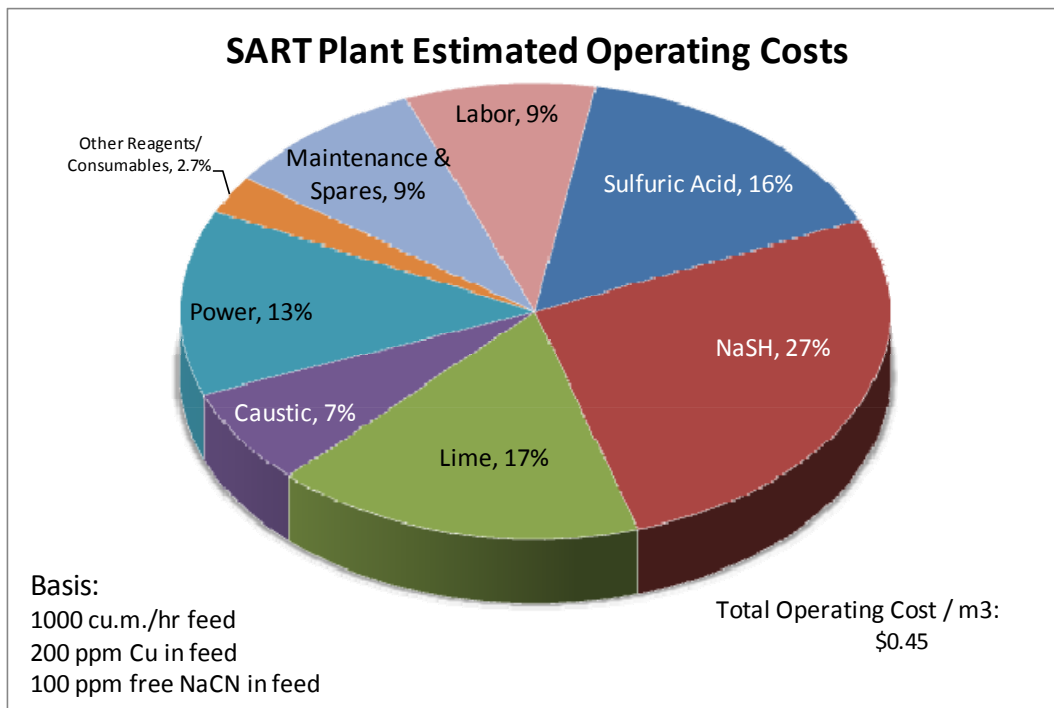
- pH reduction from alkaline conditions to acidic using sulphuric acid. The operating pH range is 4.0-4.5. Excursions above this range reduce copper precipitation, and excursions below this range can increase precipitation of gold and form undesirable copper precipitates such as CuCN and CuSCN.
- Copper precipitation with NaSH. Control of this reagent is very important. NaSH is expensive, and excess NaSH will consume free cyanide to form thiocyanate. The best control philosophy seems to be to use a slight deficiency of NaSH so the discharge of the SART plant still contains a minor amount of copper. Commercial instantaneous on-line analyzers for copper (including cuprous ion) are limited. Fortunately, in most heap leach solutions the copper concentration fed to SART will be fairly consistent as it is averaged over a large volume of solution, so off-line analysis of Cu for NaSH control maybe is satisfactory in some of these cases.
- pH adjustment to alkaline upon solution discharge using lime. The only important consideration in this control is to achieve a pH above 10.5 to fix the regenerated cyanide.
- Flocculent and de-scalant usage. Flocculent will be used in the copper sulfide thickener, and de-scalant may be used at more than one point in the process. These can be evaluated in the lab prior to plant design, but the final use will be determined once the plant is in operation.
- Sodium hydroxide for neutralizing copper sulfide precipitate at a pH of about 10. This control is mainly for safety purposes. Residual cyanide in the copper sulfide sludge fed to the filter presses is very small and will cause minimal if any re-dissolution of copper.

- Sodium hydroxide for HCN gas scrubbing.

## 5. SART Plant Economics

### 5.1. SART Estimated Operating Costs

For illustrative purposes, a breakdown of SART plant operating costs is provided below. In this example the design is based on an arbitrary 20000 m<sup>3</sup>/day influent (833 m<sup>3</sup>/hr) containing 200 ppm Cu and 100 ppm free NaCN, which is the same basis for which the design considerations in Section 4 were treated (i.e. a medium sized heap leach of 20000 tonnes/day containing 200 g Cu/tonne ore leached, treated 100% through the SART circuit).



**Figure 3. Breakdown estimate of SART Plant Operating Costs.**

Reagent costs used are \$1.00/kg NaSH, \$0.125/kg acid, and \$0.15/kg lime, assuming average transportation costs (i.e. reasonably good access to site). Power cost assumed \$0.10/kWh. Labor costs reflect a fairly high amount of plant automation (automatic control of reagents and flows) and thus minimal staffing.

As shown in Figure 3, total cost of the SART treatment in our example is US\$0.45 per m<sup>3</sup>. At 200 ppm copper recovered, (200 grams Cu / m<sup>3</sup>), the plant breaks even at a net realization cost (for copper in the precip) of about US\$3.60 per kg copper (which includes a \$1/kg transport-smelting-refining charge for the Cu) with no credit given to cyanide recovery.

Matthew Stewart, Daniel Kappes

In terms of cyanide saved, at a price of \$2.50/kg NaCN, the plant in our example needs only to recover about 50% of the influent copper-bound cyanide to cover operating costs. A properly designed SART plant should routinely recover 80-95% of copper-bound cyanide.

With high prices for copper (\$7/kg assumed in this case) and costs for cyanide in recent times, a SART plant may actually be an ancillary profit center for a project.

Table 2 illustrates the potential savings/benefits of SART in different terms of cash-cost of Au produced, at different copper grades at a gold grade of 0.5 g/tonne on a recovered basis, using the same assumptions as above (for a 1 g/tonne Au grade the costs would simply be halved).

**Table 2. Estimated credit through SART in terms of cash-cost of Au produced.**

Recovered Gold (g/tonne)	Recovered Copper (g/tonne)	SART Operating Cost Plus TSR* (per ounce Au)	Cyanide Savings** (per ounce Au)	Copper Revenue (per ounce Au)	Net SART Credit*** (per ounce Au)
0.5	100	\$26	\$29	\$41	\$44
	200	\$40	\$57	\$82	\$100
	500	\$80	\$179	\$206	\$306

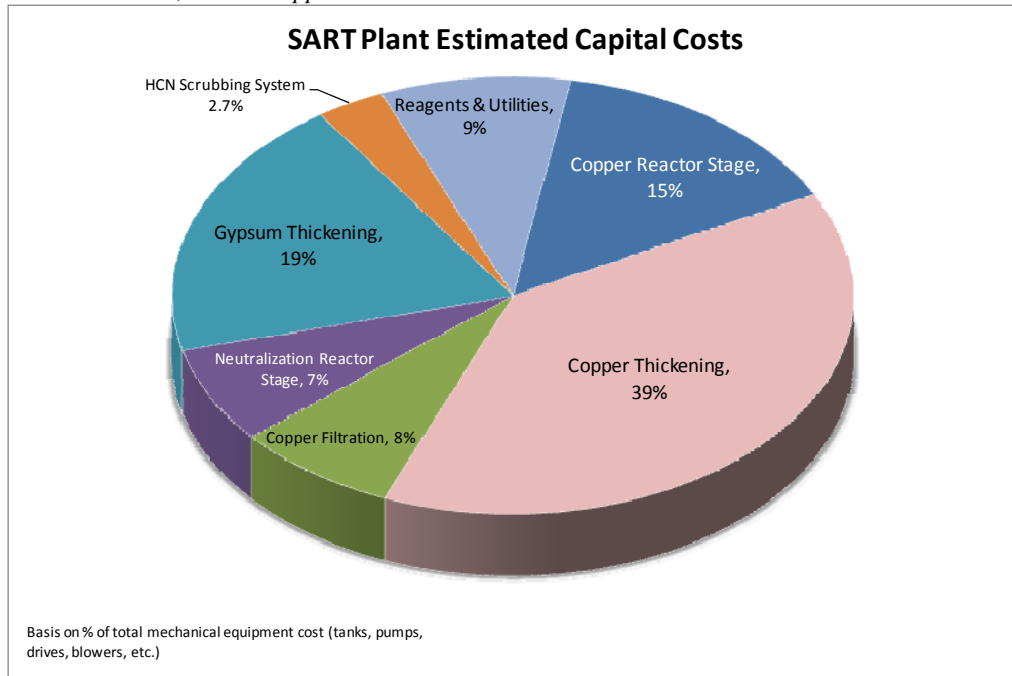
\*TSR - Transport/Smelting/Refining at \$1/kg Cu

\*\*Assumes 80% cyanide recovery in SART

\*\*\*Credit versus no SART treatment of ore

## 5.2. SART Estimated Capital Costs

For illustrative purposes, a breakdown of a SART plant capital costs is also provided. This breakdown is based on the same general assumptions as for estimating the operating costs. A 1000 m<sup>3</sup>/hr plant is assumed. Costing is estimated from a general equipment list that reflects a standard SART plant design. The costs do not include any infrastructure requirements such as water, power supply, etc.



**Figure 4. Breakdown estimate of SART plant capital costs.**

As can be seen from Figure 4, almost 40% of the total mechanical equipment cost comes from the copper thickener alone (tank, rake, associated pumps and equipment), and about 60% of the total cost is from the two thickeners together. This points out that proper copper thickener design and careful consideration to gypsum disposition (e.g. determining if the gypsum thickener is even necessary) are keys to optimizing the plant cost.

When considering the traditional SART flowsheet, an installed turnkey SART plant at the 1000 m<sup>3</sup>/hr size might be expected to cost somewhere between \$750 and \$1500 per daily m<sup>3</sup> treated. A SART LITE plant would cost about 65% of a traditional SART plant. It might not be possible to incorporate all the cost savings of a SART LITE plant, of course, but keeping an open mind to a “value-oriented” engineering approach to the design could in some cases result in a better overall project than to simply follow the conventional approach.

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*Matthew Stewart, Daniel Kappes*

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