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FACTORS INVOLVED IN THE
HEAP LEACHING OF COPPER ORES



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# DEPARTMENT OF COMMERCE - BUREAU OF MINES

# FACTORS INVOLVED IN THE HEAP LEACHING OF COPPER ORES

By John D. Sullivan2

#### INTRODUCTION

The recovery of copper by leaching from relatively low grade ores containing both exidized and sulphide minerals is rapidly gaining in commercial importance. There is an economic limit below which certain ores can not be treated by concentration methods, and material with values below this limit must be discarded, possibly to be treated later, or the copper must be extracted by wet processes.

Heap leaching has been carried out by the Phelps Dodge Corporation, Copper Queen Branch, for several years, and to-day its plant is a large producing unit. The Ohio Copper Co. at Bingham, Utah, has been leaching in place since 1919. Many other leaching operations are also being carried out in various parts of the United States and in foreign countries. The plant at Rio Tinto, Spain, may be considered the father of heap leaching.

In heap leaching the ore is usually heaped as it comes from the mine without any breaking or crushing treatment. Frequently pieces of rock 5 or 6 feet across a face are found in the heaps. Solution is added to the ore at the surface and seeps downward by gravity and is collected in a solution sump. Solution is usually added to one section of a heap and, after a certain quantity has been added, the solution is added to another section. Sometimes weeks or even months elapse between additions of solution to a given section.

Four factors are essential to the successful leaching of any copperbearing ore: (1) A solution that will attack the copper minerals must get into the body of the ore particles, (2) the copper minerals must be dissolved by the solution, (3) the solution containing the copper must find its way out of the voids into the main solution stream, and (4) the copper in solution must be recovered by some means of precipitation.

The United States Bureau of Mines at its Southwest Experiment Station, in cooperation with the department of mining and metallurgy, University of Arizona, has undertaken a study of these fundamental factors involved in the

<sup>1 -</sup> The Bureau of Mines will welcome reprinting of this paper, provided the following footnote acknowledgment is used: "Reprinted from U. S. Bureau of Mines Information Circular 6425."

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leaching of copper ores. So far, the study has been limited to the first three factors, which are those involved in the leaching operation itself. In leaching a given ore the rate of extraction is not instantaneous but goes on slowly, and the factors of penetration, dissolution, and diffusion go on simultaneously and not in successive steps. However, these steps can be studied only by segregating them so as to have only one factor entering at a time. The results thus obtained lend themselves to comparisons that can be used in commercial practice.

This paper presents a resume of the results obtained in the heapleaching studies made at the Southwest Experiment Station, and the general conclusions which have been drawn.

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Former papers have described in detail the experimental procedure and the results obtained in the study of: (1) The ingress of solutions into ores during leaching, 3.4 (2) the dissolution of various copper minerals, 5.6.7.8.9 and (3) the removal of soluble copper from leached ores. 10.11.12

- 3 Sullivan, J. D., Keck, W. E., and Oldright, G. L., Factors Governing the Entry of Solutions into Ores during Leaching; Tech. Paper 441, Bureau of Mines, 1929, 38 pp.
  - 4 Sullivan, J. D., and Ostrea, Enrique, Factors Governing the Entry of Solutions into Ores during Leaching II: Tech. Paper 498. Bureau of Mines, in press.
  - 5 Sullivan, J. D., Dissolution of Various Oxidized Copper Minerals: Rept. of Investigation 2934, Eureau of Mines, 1929, 9 pp.
- 6 Sullivan, J. D., and Oldright, G. L., The Dissolution of Cuprite in Sulphuric Acid and in Ferric Sulphate Solutions: Rept. of Investigation 2967, Bureau of Mines, 1929, 9 pp.
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  7 Sullivan, J. D., Chemistry of Leaching Chalcocite: Tech. Paper 473, Bureau of Mines, 1930, 24 pp.
- 8 Sullivan, J. D., Chemistry of Leaching Bornite: Tech. Paper 486, Bureau of Mines, 1931, 20.pp.
  - 9 Sullivan, J. D., Chemistry of Leaching Covellite: Tech. Paper 487, Bureau of Mines, 1930, 18 pp.
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    Rept. of Investigations, 3073., Bureau of Mines, 1931. 21 pp.

### INGRESS OF SOLUTIONS INTO ORES

In leaching an ore, the solution must first permeate the ore in order to come into contact with the copper-bearing minerals. There are, in general, two classes of voids in rocks: (1) Fractures and cleavage planes and (2) pores of capillary size or smaller. The crevices and cleavage planes are usually of such a nature that they are open along one side or at both ends. For a solution to enter these openings it must mechanically push the gas out of the voids. As these openings are not in general in a horizontal position, there may be a larger hydrostatic head at one end of the opening than at the other. It is not particularly difficult for solutions to enter large voids. In the second class of voids, the pores may extend through the rock or they may be connected by a network to one another so that they are open at both ends, but pores open at one end only are also present. Even in small pores openat both ends, if solution penetrates simultaneously from both ends, the pores act as though closed at one end. In the case of a pore closed at one end, solution can penetrate only by replacing the gas within the pore. A solution will not penetrate unless the pressure within the pore is less or equal to the outside pressure. The gas in the closed part of the pore must either dissolve in the penetrating liquid or diffuse through it, if penetration takes place.

The movement of liquids in rocks through supercapillary openings (circular openings greater than 0.508 mm. in diameter, or sheet openings greater than 0.254 mm. between walls) is probably governed by the ordinary laws of hydrostatics. Poiseuille's law probably governs the movement of liquids in capillary openings (circular openings 0.0002 to 0.508 mm. in diameter, or sheet openings 0.0001 to 0.254 mm. between walls). Flow through subcapillary openings would be infinitely slow, as the attraction of solid molecules extends from wall to wall, and no liquid would be free to move, as it is attracted to the walls as a film.

A description has been given elsewhere of the apparatus and procedure used for determining: (1) The rate and volume of penetration of solutions into ores, (2) the total volume of voids within particles of ores, and (3) the density of ores.

Table 1 shows the rate at which water penetrated into various sizes of a typical porphyry ore of the Southwest. The data show that the time needed for a given percentage penetration increases markedly with increases in size of particles.

<sup>13 -</sup> Sullivan, J. D., Oldright, G. L., and Keck, W. E., Method for Measuring Voids in Porous Materials: Rept. of Investigations 3047, Bureau of Mines, 1930, 8 pp.

Table 1. - Rate of penetration of distilled water into various sizes of a perplyry ore; temperature, 30° C.

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Time, hours	1 - 850 1500 12 45	Voids filled per cent					
	-1+3-inch pieces	2-inch pieces	3-inch pieces	4-inch pieces			
1/60	66.1	31.4	20.4	16.8			
1/30	69.3	37.2	26.5	21.8			
1/10	75.7	45.7	34.2	28.1			
1/4	80.5	54.2	44.7	33.7			
1/2	83.6	59.5	53.8	37.5			
1	86.4	63.6	62.0	42.0			
2	89.2	68.4	69.2	46.9			
3	89.8	71.6	70.4	50.2			
5	91.0	75.0	71.5	53.3			
$23\frac{1}{2}$	95.3	80.5	75.7	62.2			
48	96.8	82.0	77.2	66.3			
84	98.1	83.0	78.5	69.1			
108	98.1	83.5	79.4	71.2			
336	THE STATE OF THE S	34 4 86.02	83.5	81.3			
360	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		83.7	82.0			
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Surface Tension.—The general opinion has been that solutions entered the pores of rocks by capillarity. If capillarity is the governing factor, then by changing the surface tension of the penetrating liquid, the rate of entry of solution should also be changed. Tests were made in which the surface tension of water was lowered from approximately 75 dynes to about 25 dynes per centimeter by adding enough sodium cleate to make a saturated solution, but the rate of entry of solution into the ore was practically identical with that of pure water and

Solubility of Ges in Voids.—As surface tension has little or no effect upon the rate of penetration of solution into ores, the rate must be governed primarily by some other factor, which is indicated to be the solubility in the penetrating solution of the gas or gases within the voids of the ore. The solubility of sulphur dioxide in water is 3,937 cubic centimeters in 100 cubic centimeters of water at 20°C., whereas the solubility of air 16 is 1.8 cubic centimeters in 100 cubic centimeters of water at the same temperature. Data in

<sup>14 -</sup> Harkins, W. D., Davies, E. C. H., Clark, G. L., Surface Energy, VI: Jour. Am. Chem. Soc., vol. 39, 1917, p. 587.

<sup>15 -</sup> Seidel, A., Solubilities of Inorganic and Organic Substances: New York, 1917, p. 329.

<sup>16 -</sup> Seidel, A., Page 10 of work cited.

Table 2 show the rate at which distilled water penetrated into various sizes of a typical porphyry ore which had been evacuated and the voids filled with sulphur dioxide. The ore was the same as that used in the tests summarized in Table 1. When the voids were filled with sulphur dioxide, water penetrated more rapidly, especially during the early part of the impregnation. Not only was the rate of penetration faster, but the total volume of penetration was also greater.

Table 2.—Rate of penetration of distilled water into various sizes of a porphyry ore the voids of which were filled with sulphur dioxide; temperature, 30° C.

Time, hours	Voids filled, per cent				
	-1+3/4-inch pieces	2-inch pieces	3-inch pieces	4 inch pieces	
1/60	69.5	45.0		43.3	
1/30	78.0	52.0	50.2	45.8	
1/10	84.6	65.0	66.4	50.1	
1/4	90.0	80.0	79.4	55.4	
1/2	95.3	86.3	85.2	58.2	
Ľ	95.8	89.2	88.5	62.6	
3	96.4	92.0	90.6	66.4	
3	97.1	93.4	91.0	68.5	
5.	97.7	94.0	92.2	71.0	
232	98.8	95.5	94.3	77.2	
18	99.1	96.7	95.4	79.8	
34	99.7	. 97.7	96.0	82.0	
108	100.0	98.8	96.2	83.0	
336		100.0	97.9	88.4	
360			98.1	89.1	
792	p-service.		100.0	98.4	
388				100.0	

Temperature.—Measurements made at 2 to 3.5° C. and at 35° C. showed that the rate of penetration was more rapid at the lower temperature. For a given ore, 95 per cent of the total penetration that took place was attained in 40 hours at 2 to 3.5° C., whereas 50 hours was required at 35° C. As the solubility of gases in water increases with a decrease in temperature, the solution might be expected to penetrate at a faster rate at the lower temperature.

Various Solutions.—There is surprisingly little difference in the rate of penetration of various kinds of solutions into rocks; 5 per cent copper sulphate, 2 per cent sulphuric acid, 2 per cent copper sulphate or ferrous sulphate plus sulphuric acid, 2 per cent ferric sulphate, and distilled water have very nearly the same rates of penetration.

Manner of Penetration. -- It has been found that in most ores the solution enters along fractures and cleavage planes and that from these larger fractures it gradually seeps into the rest of the rock. The first penetration

takes place very rapidly along these crevices and fractures, and the small voids are filled more slowly from these points of initial penetration. Photographs demonstrating this phenomenon can be found elsewhere. 17

### DISSOLUTION OF VARIOUS COPPER MINERALS

Oxidized Minerals.—The oxidized minerals of copper—azurite, malachite, chrysocolla, and tenorite—are soluble in sulphuric acid and in ferric sulphate solutions. Approximately 100 per cent of the total copper is soluble in one hour at 35°C. in sulphuric acid or in acidified ferric sulphate solutions when the size of mineral is minus 100 plus 200 mesh. In 1 or 2 per cent neutral ferric sulphate the dissolution is slower, but in nearly every instance the mineral is completely dissolved in 24 hours or less. The rate of dissolution in 5 per cent ferric sulphate is about the same as in sulphuric acid or in acidified ferric sulphate.

Cuprite.—When cuprite is leached with sulphuric acid in the presence of excess atmospheric oxygen, approximately 100 per cent of the copper in pieces up to 4-mesh in size is converted into the soluble sulphate in 20 days at 35° C. With a minus 3 plus 4 mesh product, approximately 94 per cent is rendered soluble in the same time. For pieces 100 mesh or smaller in size, 24 hours is sufficient for complete dissolution. When cuprite is treated with acidified ferric sulphate, particles up to 3 mesh in size are completely dissolved in 8 days and in 3 days 99 per cent is dissolved. For 100-mesh or smaller particles, one hour is sufficient for complete dissolution. When cuprite is leached with sulphuric acid in the absence of oxygen (as, for example, in closed bottles in an atmosphere of nitrogen) only one-half of the copper is rendered soluble, and one-half of it remains as metallic copper, according to the reaction:

When cuprite is leached, either in the presence or in the absence of oxygen, a layer of metallic copper forms practically as soon as the mineral comes in contact with sulphuric acid. This metallic copper forms a difficulty permeable layer on the surface of the particles that slows down the dissolution. The metallic copper may be converted to copper sulphate by the aid of an oxidizer. Atmospheric oxygen is a fairly good oxidizer but ferric sulphate is a much better one. On particles 100 mesh or smaller in size this metallic coating of copper does not markedly hinder the rate of dissolution, but it is very harmful for larger sizes.

Chalcocite.—Chalcocite is the principal sulphide mineral encountered in leaching. When it is leached with ferric sulphate, the dissolution takes place in two stages, which may be written:

$$\text{Cu}_2\text{S} + \text{Fe}_2(\text{SO}_4)_3 = \text{Cu}_3\text{SO}_4 + 2\text{Fe}_3\text{SO}_4 + \text{Cu}_3\text{S}, \text{ and }$$
 $\text{Cu}_3\text{S} + \text{Fe}_3(\text{SO}_4)_3 = \text{Cu}_3\text{SO}_4 + 2\text{Fe}_3\text{SO}_4 + 3\text{S}_4$ 

<sup>17 -</sup> Sullivan, J. D., Keck, W. E., and Oldright, G. L., Factors Governing the Entry of Solution into Ores during Leaching: Tech. Paper 441, Bureau of Mines, 1929, pp. 35-36.

The first reaction is much more rapid than the second. For particles of mineral 10 mesh or smaller, approximately 50 per cent of the total copper is converted into the soluble sulphate in 24 hours at 35° C., whereas approximately 20 days is required to dissolve the other half. Particles as large as 2 or 3 mesh dissolve at only a slightly slower rate than minerals crushed as fine as minus 150 plus 200 mesh. An oxidizing agent is necessary to dissolve chalcocite. Sulphuric acid in the absence of oxygen or water even in the presence of oxygen have practically no dissolving effect upon the mineral. Sulphuric acid in the presence of excess atmospheric oxygen attacks the mineral, but does so more slowly than ferric sulphate solutions. At 35° C. the rate of dissolution is practically the same in ferric sulphate and in ferric chloride. The rate of dissolution increases with increases in temperature. In 48 hours 50, 59, and 87 per cent of the copper was dissolved from minus 100 plus 200 mesh chalcocite at 23, 35, and 50° C., respectively.

Bornite. -- Data on the rate of dissolution of various sizes of bornite, Cu<sub>5</sub>FeS<sub>4</sub>, in acidified ferric sulphate are given in Table 3.

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Table 3.--Dissolution of various sizes of bornite in a solution containing 1 per cent of iron as ferric sulphate plus 0.5 per cent of sulphuric acid; temperature, 35°C.

Time	Copper extracted, per cent					
*	-2+3 mesh	-10+28 mesh	-100+200 mesh	-200 mesh		
1 hour	4	19	27	29		
3 hours	7	26	28	30		
8 hours	12	30	31	33		
24 hours	21	38	38	41		
48 hours	32	53	51	56		
3 days	43	61	60	66		
4 days	51	66	65	72		
5 days	58	69	69	77		
7 days	68	74	78	-87		
10 days	78	84	94	97		
14 days	86	97	98	99		
21 days	94	99	99	99		
28 days	97	100	99	100		

The rate of dissolution of bornite is markedly increased by increases in temperature. When minus 100 plus 200 mesh bornite was leached with acidified ferric sulphate 64 per cent of the copper was dissolved in 1 day at 50° C., in 4 days at 35° C., and in 14 days at 23° C. Eighty per cent of the copper was dissolved in 6 hours at boiling temperature. Bornite dissolves more rapidly in ferric chloride than in ferric sulphate. Sulphuric acid plus air attack bornite more slowly than ferric sulphate solutions.

Covellite. -- Table 4 gives data on the rate of dissolution of various sizes of covellite in acidified ferric sulphate.

Table 4.—Dissolution of various sizes of covellite in a solution containing 1 per cent of iron as ferric sulphate plus 0.5 per cent of sulphuric acid

Time Temper	Temperature .	Copper extracted, per cent			
		-3+10 mesh	-10+28 mesh	-100+200 mesh	-200 mesh
1 day 5 days 11 days 24 days 31 days 39 days 47 days	35 35 35 35 <u>1</u> 50 50 50	12 14 16	6 10 11 13 18 21 24	8 16 21 25 44 50 57	13 40 49 56 75 82 87

1 - Temperature maintained at 35° C. for 24 days and then increased to 50° C.

The rate of dissolution of covellite increased with increases in temperature. For a given sample, 81 per cent of the copper was extracted in 14 hours at 98° C., 81 per cent in 22 days at 50° C., and 41 per cent in 24 days at 35° C. The rate of dissolution was more rapid in ferric sulphate than in ferric chloride at 35° C., but the rates were virtually the same at 98° C. Covellite dissolved in sulphuric acid in the presence of excess atmospheric oxygen about half as rapidly as in ferric sulphate.

Chalcopyrite.—Chalcopyrite is frequently found in leaching ores, but it is not appreciably attacked by common solvents at ordinary temperatures. Chalcopyrite is a closely textured mineral, and its rate of dissolution is greatly increased by fine grinding. Data on the rate of dissolution of chalcopyrite are given in Table 5.

Table 5. Data on dissolution of chalcopyrite

Size, mech	Tempera- ture, oc.	Solvent	Copper dis- solved in stated
			time, per cent
-100+200	35	0.5% of iron as Fe2(SO4)3 plus 0.5% of H2SO4	1.8 in 45 days
-100+200	35	1.0% do. 0.5% do.	1.6 in 45 days
<b>-</b> 350	35	2.0% do. 0.5% do.	28.5 in 25 days
-350	35	2.0% do. 0.5% do.	39.2 in 57 days
-100+200	50	1.0% do. 0.5% do.	4.6 in 8 days
-100+200	50	1.0% do. 0.5% do.	6.2 in 14 days
<b>→</b> 350	50	1.0% do. 0.5% do.	32.2 in 8 days
<b>-35</b> 0 .	, 50	1.0% do. 0.5% do.	43.6 in 14 days
<b>-</b> 050	35.	1.0% of iron as FeCl3	32.6 in 25 days
-350	35	1.0% of iron as FeCl3	45.3 in 57 days

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Figure 1. -- Original minus 2 plus 3 mesh bornite
before leaching



Figure 2.--Minus 2 plus 3 mesh bornite after leaching for 14 days at 50° C. with acidified ferric sulphate

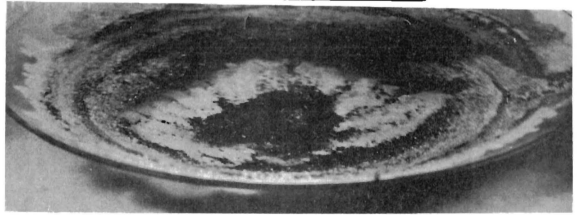


Figure 3.- Residue from leaching minus 2 plus 3 mesh bornite after treatment with carbon bisulphide

The foregoing data show that the rate of dissolution is faster when the mineral is more finely ground, that the rate increases with an increase in temperature, and that the rate of dissolution is more rapid in ferric chloride than in ferric sulphate.

In heap leaching an appreciable amount of the copper in chalcopyrite may be extracted owing to the long period of contact with solutions, and the possible rise in temperature of the ore during weathering.

When chalcocite is leached with ferric sulphate one-half of the copper dissolves rapidly leaving a residue approximating CuS which dissolves much slower; with bornite the copper is attacked preferentially to the iron; while with chalcopyrite the mineral dissolves as chalcopyrite; that is, at any stage in the dissolution(up to an extraction of approximately 45 per cent of the copper which has been the maximum studied), the ratio of Cu:Fe:S in the residue remains the same as in the original mineral. With covellite at any stage in the dissolution the residue has approximately the same ratio of Cu:S as the original mineral.

When large pieces of chalcocite, bornite, or covellite are leached with ferric sulphate the particles do not disintegrate and the residue become a powder. The particles, after leaching, nearly retain their original outline, and very little powder is formed. Apparently the sulphur left behind retains the original form of the mineral particles. When carbon bisulphide is added to the residue, the sulphur dissolves and the particles collapse. Figures 1, 2, and 3 demonstrate this phenomenon. Figure 1 shows minus 2 plus 3 mesh bornite before leaching; Figure 2, the residue after leaching for 14 days at 50° C. with acidified ferric sulphate; Figure 3, the residue after treatment with carbon bisulphide. In the 14-day period of leaching 91 per cent of the copper was dissolved.

#### REMOVAL OF SOLUBLE COPPER FROM LEACHED ORES

In heap leaching, because the solution is added to a given section of the heap and because after a certain quantity of solution has been added the leach solution is added to another section, several months may elapse between additions of solution to a given section. Heap leaching, therefore, results in a process of alternate wetting and drying. When the surface is dried, evaporation pulls solution to the surface where the dissolved salts crystallize as the solutions evaporate, and the next wetting operation removes part of these salts. At the same time that the salts are removed by washing some solution penetrates into the cavities, cleavage planes, and pores within the ore. If all of the moisture within the ore was removed during the drying cycle, all of the copper salts would be precipitated, partly at the surface and partly within the ore. When the ore is washed, the distance that the solution penetrates into the ore is a function of the time of washing.

The alternate wetting and drying tests carried out in the laboratory could not easily be duplicated in heap leaching. In the laboratory the surfaces of the ore may be completely dried, and it is possible to dry the entire sample.

In heap leaching, a complete drying would be practically impossible. As a heap may contain several million tons of ore complete drying, even of the surface, could hardly be expected. Experiments have shown that the copper can be brought to the surface even though the particles of ore are only partly dried.

With a 0.5-hour period of drying and a 0.5-hour period of washing and the cycle repeated until nearly all of the copper was extracted, 80 per cent of the total copper in solution was extracted from 3-inch pieces of ore in 6 hours, whereas 44 hours was required for 5 to 6 inch pieces. This shows the advantage of employing small pieces of ore.

In hear leaching, very short periods of alternate wetting and drying can not be maintained, but laboratory work has shown the advantage gained by keeping the cycles as short as possible. With 3-inch pieces of ore, an extraction of 80 per cent of the water-soluble copper was obtained in 6 hours with a 0.5-hour period of drying and a 0.5-hour period of washing, whereas 25 hours was required for a 6.0-hour period of drying and a 2.0-hour period of washing. Any advocacy of shorter cycles in alternate wetting and drying presupposes that the heaps are porous and well aerated.

The soluble copper can be removed by alternate wetting and drying in approximately 15 to 25 per cent of the time required to remove it by flood washing, provided the washing and drying periods are as close to each other as possible but long enough to permit a fairly thorough drying of the charge and soaking in of the leaching solution. As an example, it took approximately 130 hours to remove 90 per cent of the water—soluble copper by vat washing from the minus 1 plus 3 inch size of a porphyry ore saturated with copper sulphate, whereas only 31 hours was required by alternate wetting and drying when the period of drying was 4.0 hours and the period of washing 0.5 hour.

A rapid movement of air past the surface of the ore promotes rapid drying. Anything that interferes with the circulation of air slows down the rate of extraction, thus demonstrating the necessity of having an open heap where free circulation of air is possible. Slime or other material that will coat the surface would also hinder drying. The rate of extraction is also increased by an increase in temperature.

When a rock is saturated with copper sulphate and then dried, the crystallized salt is disseminated throughout the entire rock, especially along the fractures and cleavage planes. A large part of the copper is brought to the surface in the first drying operation. Some of the copper is crystallized in the interior of the rock, especially along larger fractures and cleavage planes. Photographs demonstrating this phenomenon can be found elsewhere. 18

<sup>18</sup> Sullivan, J. D., and Sweet, A. J., Factors Governing Removal of Soluble Copper from Leached Ores: Tech. Faper 453, Bureau of Mines, 1929, p. 22 ff.

## CONOLUSIONS

In leaching, the three factors (1) ingress of solutions, (2) dissolution of minerals, and (3) removal of the soluble copper are interdependent and work simultaneously. The speed of leaching can be no faster than the slowest step. Enough solution can not enter a rock at one time to dissolve the copper minerals completely unless the copper content of the ore is very low. A sample of ore weighing 2,000 grams may soak up about 40 cubic centimeters of solution. If the ore contains 1 per cont of copper, and the solution coming out of the peres contains 20 grams of copper per liter (assumed), 1,000 cubic centimeters of solution or approximately 25 renewals of the solution within the voids would be necessary. Frequent renewals of solution and much time is required to get sufficient solvent into contact with the minerals and to wash out the dissolved copper.

In heap leaching with oxidized ores the slowest step is probably the removal of the soluble salts. If the copper exists as sulphide (chalcopyrite excepted), about 20 days will be required for complete dissolution of the copper, granting contact of mineral and solvent solution during the entire period. In this instance, the slowest step is probably the chemical one or the dissolution of the copper sulphide, granting that conditions found in the laboratory hold in practice.

The rate of dissolution of the copper minerals is a chemical factor; and in heap leaching practice, for a given solvent, the rate can not be markedly changed. Although the rate of dissolution increases with increases in temperature, it is difficult to control temperature in a large heap. Actual temperature measurements within the heaps usually indicate a higher temperature than that of the outside air, which shows that the heat from chemical reactions is fairly well stored within the heap. It may therefore be possible to increase gradually the temperature within the heaps by adding warm solution. In the Southwest the solutions could be warmed by the sun in summer.

Waste smelter gases, if available, might be used with a heap leaching plant. If waste gases are allowed to enter heaps and displace the air present before water is added to a given section of the heap, the solutions should penetrate more rapidly into the interior of the rocks. Also, waste gases would add a reagent that would help to dissolve the copper minerals present.

From a practical standpoint, the method of speeding up the rate of extraction is to crush the ores finer. Crushing the ore particles would not only decrease the time required for saturation but it would also make the ore more amenable to leaching by open cleavage planes and crevices in the ore particles and by shortening the path required for the leaching solutions to come into contact with the mineral particles and for the dissolved copper to be brought to the surface. Crushing to a smaller size than 2 or 3 inches might produce too many fines, which would tend to counteract the advantages of crushing.

Another means of speeding up the extraction is by recirculating solutions over the heap. Frequently mine water containing neither acid nor iron salts is passed over the ore, but owing to the weathering of pyrite and other sulphides there is a considerable concentration of sulphuric acid and salts of bivalent and trivalent iron in the effluent solutions. As chalcocite is amenable to leaching by solutions of salts of trivalent iron, there is good reason for recirculating at least a part of the effluent solutions. To avoid "plugging" the leaching column with flocculent hydrated iron precipitates, the solutions recirculated should contain some free acid.

Heaps should be laid down on prepared footings to prevent the loss of solutions by seepage. This is especially true if pregnant liquors containing copper are recirculated.

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