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A Chemical Study of the Enrichment of Copper Sulfide Ores

A DISSERTATION

Submitted to the Department of Chemistry and to the Committee on Graduate Study of the Leland Stanford Junior University in partial fulfillment of the requirements for the degree of Doctor of Philosophy

BY

JOHN DUSTIN CLARK
CRYSTALS OF CUPRO-CUPRIC SULFITE
× 50
See experiment No. 10.

CHALCOGITE CRYSTAL
× 200
See experiment No. 41.
Since the time recorded in our earliest history metallic ores have been taken from mines and deposits. Undoubtedly the first discovered deposits were found by accident. The mining of the ore probably followed well defined veins and stopped when such were exhausted. The work of the ignorant prospector of to-day is much like the probable simple procedure of the ancients.

Observations which have led to the science of Geology must have made some miners proficient in predicting the occurrence of mineral bearing veins, etc. It is certain that to-day the mining engineer and mining geologist can, as a result of training and observation, "see beneath the ground", and can successfully direct development work carried on to locate valuable metal-bearing bodies.

The engineer and geologist has made use of many of the sciences in conducting his work. Chemistry has always played an important part, and it seems that its field of usefulness has only been entered upon. It appears that the results of purely theoretical and experimental laboratory work in chemistry may be of the most practical importance when given to these mining ex-
perts. This much is certain, that for understanding enrichment in our copper mines much chemical work is necessary, as may be seen when an authority like W. H. Emmons\(^1\) says, (in his publication, “Enrichment of Sulfide Ores”,) “To the chemist this paper is an appeal for more experimental data on the important mineral syntheses involved in the processes.”

**DEFINITIONS OF ENRICHMENT.**

The meaning of *downward* secondary enrichment and a summary of much of the field observation, laboratory collaboration and commercial significance is given in language readily understood by the average man, yet not lacking in scientific information, by Tolman\(^1\) in his article on “Secondary Sulfide Enrichment of Ores”. Here is quoted his brief definition of this *downward* process. “Secondary sulfide enrichment involves three processes: (1) The oxidation of the metallic sulfides; (2) The solution of these chiefly as sulfates, chlorides or bicarbonates; and (3) The precipitation of the metals in solution in the form of secondary sulfides, (a) by the reduction of the sulfates to metallic sulfides by carbonaceous matter, or (b) by precipitation by means of hydrogen sulfide, or (c) by reaction of the metallic salts with unoxidized

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sulfides below the water level, the latter going into solution as sulfates, (or other salts), and the former precipitating as sulfides. In the broadest sense secondary sulfide enrichment includes deposits of any sulfide precipitated in any way out of the descending meteoric waters which owe their metallic content to the leaching of the overlying rock."

A. F. RogersII has recently applied the term upward secondary enrichment to processes by which metal-bearing solutions bring their metals as sulfides from greater, and deposits them at lesser depths, in distinction from processes in which the ore-forming solutions flow downward.

It is believed that the results of the present investigation have shown that upward enrichment without oxidation is a very important process. As will be demonstrated, crystalline copper sulfides, whether in the form of chalcopyrite, bornite, covellite or chalcocite, may be brought into such condition as to migrate considerable distances without the intervention of any process of oxidation or any process converting the copper into electrolyte solution. Such agencies, as will be shown, are those producing the colloidal copper sulfides.

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II—A. F. Rogers, Econ. Geol. 8, p. 781, (1913).
OXIDATION AND SOLUTION OF COPPER MINERALS.

A review of the literature concerning these steps shows that they may take place in one or more of the following ways:

(1) Oxidation of pyrite to produce sulfuric acid and ferric sulfate, which then attack and dissolve chalcocite or other copper minerals.

(2) Thru the greater electrolytic solution tension of some minerals in contact with others.

(3) Solution of a mineral in pure water.

These three are prominently mentioned and are generally accepted as being effective. To these may be added, as is later shown by the author:

(4) Solution or dispersion of the mineral by carbonated water, and that suggested but hardly proved method:

(5) Solution of the metal as a chloride where pyrite and manganese dioxide have a chloride bearing water percolating thru them.

In addition to the processes which have been suggested in the literature it will be shown in this paper that, (from the chemical point of view at least), the enrichment by the highly dispersed copper sulfides, as such, may be, as a result of this work, looked upon as an important factor in the general process of enrichment.
Going into detail concerning these methods of solution we find that the following equations may represent the oxidation of the pyrite, though as Tolman\(^1\) says, "the exact chemical equations are not known and perhaps will never be written":

1. \(\text{FeS}_2 + 4 \text{O} = \text{FeSO}_4 + \text{S}\)
2. \(\text{FeS}_2 + 6 \text{O} = \text{FeSO}_4 + \text{SO}_2\)
3. \(\text{FeS}_2 + 7 \text{O} + \text{H}_2\text{O} = \text{FeSO}_4 + \text{H}_2\text{SO}_4\)
4. \(2 \text{FeSO}_4 + \text{H}_2\text{SO}_4 + 2 \text{O} = \text{Fe}_2(\text{SO}_4)_3 + \text{H}_2\text{O}\)
5. \(2 \text{Fe}_2(\text{SO}_4)_3 + 9 \text{H}_2\text{O} = 2 \text{Fe}_2\text{O}_3 \cdot 3\text{H}_2\text{O} + 6 \text{H}_2\text{SO}_4\)
6. \(\text{FeS}_2 + \text{Fe}_2(\text{SO}_4)_3 = 3 \text{FeSO}_4 + 2 \text{S} \text{ and } 2 \text{S} + 6 \text{Fe}_2\text{SO}_4)_3 + 8 \text{H}_2\text{O} = 12 \text{FeSO}_4 + 8 \text{H}_2\text{SO}_4\)

and according to Austin\(^II\) the ferric sulfate puts the copper into solution as:

\[
\text{Fe}_2(\text{SO}_4)_3 + \text{Cu}_2\text{S} = \text{CuSO}_4 + 2 \text{FeSO}_4 + \text{CuS}
\]

\[
\text{Fe}_2(\text{SO}_4)_3 + \text{CuS} + 3 \text{O} + \text{H}_2\text{O} = \text{CuSO}_4 + 2 \text{FeSO}_4 + \text{H}_2\text{SO}_4
\]

Buehler and Gottschalk\(^III\) have shown that chalcocite in contact with pyrite goes into solution as a sulfate in the presence of air, because of its greater electrolytic

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\(^I\)-Min. and Sci. Press, Jan. 4, 18, and 25, 1913.
solution tension, and that pyrite, as would be expected, is little acted upon.

It may be mentioned in passing that this is not the only possible explanation of this phenomenon, which may be explained entirely upon differences of reaction velocity in the two cases. Of course if definite equilibrium potentials were measured there would be no question as to the validity of the explanation but since all copper sulfide minerals show quite variable composition it would be natural to suppose that the electrolytic solution tensions would be extremely variable with different specimens. Until it is possible to prepare chemically pure or nearly chemically pure minerals of the various types any interpretation based upon the study of electrolytic solution tension must be considered ambiguous.

Weigel\textsuperscript{I} has shown that metallic sulfides are slightly soluble in pure water without air.

Tolman\textsuperscript{II} was of the opinion that copper minerals with an excess of carbon dioxide, in certain cases were carried downward as bicarbonates. The author has made such solutions. Whether the solution is a true electrolytic one of copper bicarbonate or a mere suspension of a copper carbonate is an open question.


\textsuperscript{II}—Suggested in Min. and Sci. Press. Jan. 4, 18, 25, 1913.
Lane believes that the copper in the Michigan mines has been carried downward as chloride solutions and suggests electrolytic migration. That the copper may have been in the chloride form seems wholly tenable judging from the chloride content of the Michigan mine waters.

Lindgren in summarizing Emmon's work mentions the possibility of solution as chlorides where nascent chlorine has played an important role, this chlorine having been produced in accordance with the question:

\[ \text{MnO}_2 + 4 \text{HCl} \rightarrow \text{MnCl}_2 + \text{H}_2\text{O} \]

He gives instances of increased solution of gold when manganese dioxide has been added to a solution containing very dilute hydrochloric acid.

The above five methods of oxidation seem to cover the principles involved. One will find, however, mention of many possibilities of oxidation and solution which combine two or more of the above methods. Considerable work has been done in attempting to discover the order in which different minerals oxidize. No two investigators have succeeded in getting the same results. In all the investigations made the investigators have failed to consider conditions under which the oxidations took place and their results put the chemist

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in mind of the old Geoffroy and Bergmann affinity tables.

The essential factors entering into the problem of oxidation are: ore, metal, country rock, fissuring, porosity, climate, water level, rainfall, topography, geological age and history of the deposit. These have to be considered in the examination of any particular mine.

**PRECIPITATION OF THE METALLIC SULFIDES.**

Various methods have been suggested by which the metals in descending solutions of metallic salts can be precipitated as sulfides. The following tried and untried means seem to cover the possibilities of such precipitations:

(1) Substitution because of solution tension of some mineral precipitant as:

\[
\text{Ag}_2\text{SO}_4 + \text{ZnS} = \text{Ag}_2\text{S} + \text{ZnSO}_4
\]

\[
14 \text{CuSO}_4 + 5 \text{FeS}_2 + 12 \text{H}_2\text{O} = 7 \text{Cu}_2\text{S} + 5 \text{FeSO}_4 + 12 \text{H}_2\text{SO}_4
\]

(2) Substitution in the presence of sulfur dioxide or other reducing agent as:

\[
4 \text{CuSO}_4 + \text{FeS}_2 + \text{SO}_2 + 6 \text{H}_2\text{O} = 2 \text{Cu}_2\text{S} + \text{FeSO}_4 + 6 \text{H}_2\text{SO}_4
\]

(3) Precipitation by means of hydrogen sulfide

(4) Precipitation by means of free sulfur

(5) Neutralization of descending acidic solutions

(6) Precipitation by means of carbon
(7) Precipitation thru loss of a dispersing agent. This paper has much to do with the last feature.

Under the first heading should be noted the work of Schuermann. He established a series of salts in which the sulfides of any one of the metals thereof will be precipitated at the expense of any sulfide lower in the series. The series is undoubtedly correct for the particular specimens used under the conditions which his experiments were performed. What such an order might be under other conditions one cannot say.

The work of Palmer and Bastin shows that such substitution occurs, the author assumes because of solution tension. They used many minerals and found them to precipitate gold and silver. Two of their equations are here given:

\[ 2 \text{NiAs} + 5 \text{Ag}_2\text{SO}_4 + 3 \text{H}_2\text{O} \rightleftharpoons 2 \text{NiSO}_4 + \text{As}_2\text{O}_3 + 3 \text{H}_2\text{SO}_4 + 10 \text{Ag} \]
\[ \text{Cu}_2\text{S} + 2 \text{Ag}_2\text{SO}_4 \rightleftharpoons 2 \text{CuSO}_4 + \text{Ag}_2\text{S} + 2 \text{Ag} \]

The first fact to be considered under the heading of substitution in the presence of sulfur dioxide or other reducing agent is that mine waters contain less and less

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I—Suggestion as to the role of colloids is given by P. Krusch, "Primary and secondary ores considered with especial reference to the gel and the rich heavy metal ores", Min. and Sci. Press, 107, pp. 418-423, 1918.


I—Palmer and Bastin, Econ. Geol. 8, No. 2, p. 140, 1913.
oxygen as the depth increases as was shown by Lepsius. Emmons states that the oxygen content and acidity of descending waters diminishes as the depth increases. This is strongly emphasized in his discussion of analyses of mine waters.

Both Tolman and Winchellv, (see also experiments No. 9 and No. 10 in this paper), have produced chalcocite films on pyrite from slightly acid solutions of copper sulfate in the presence of sulfur dioxide. Chalcocite was not precipitated in its absence.

W. H. Emmons suggests that this may be explained by the fact that sulfur dioxide removes any atmospheric oxygen from the solutions. The author will have another conclusion to add to this.

All investigators are firm in the belief that chalcocite is not precipitated in the presence of oxidizing action.

In working up their first paper above mentioned, Buehler and Gottschalk obtained sulfur dioxide where the supply of oxygen was limited.

Consideration of the next method, that of precipita-

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tion by means of hydrogen sulfide formerly raised the question as to whether hydrogen sulfide can form in the zone of precipitation. The writer having established this fact that hydrogen sulfide can form, references to data on this point are omitted.

C. F. Tolman, Jr., and A. F. Rogers hold the opinion that many of our secondary copper deposits are in contact with primary ones and that the latter were brought up by alkaline solutions rich in carbon dioxide and hydrogen sulfide and that the escape or chemical assimilation of hydrogen sulfide has disturbed equilibria sufficiently to allow the precipitation of chalcopyrite from alkaline solution. The writer has shown that such action is possible and has shown the importance of hydrogen sulfide as a dispersing agent.

Cooke\(^I\) was successful in precipitating silver sulfide by means of amorphous sulfur. Vogt\(^II\) and Stokes\(^III\) have shown that amorphous sulfur can readily form in nature.

One of the writer's most important conclusions deals with this action of amorphous sulfur.

Data on precipitation thru neutralization of de-

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\(^III\)-H. N. Stokes, Econ. Geol. 2, pp. 14-23, 1907.
descending acidic solutions are rather numerous. E. C. Sullivan secured a loss in copper content of such solutions by absorption or neutralization by keeping them in contact with powdered orthoclase albite, amphibole, shale, and clay gouge. Grout has secured precipitation by neutralization with rather strong alkalies. His work is largely qualitative. W. H. Emmons makes reference to the work of Ransome, Kemp and Lindgren, who show how limestone can be attacked by descending acid waters and can precipitate malachite and azurite in accordance with these equations:

\[
2 \text{CuSO}_4 + 2 \text{CaCO}_3 + 5 \text{H}_2\text{O} = \text{CuCO}_3\cdot\text{Cu(OH)}_2 + 2 \text{CaSO}_4 + 2 \text{H}_2\text{O} + \text{CO}_2
\]

\[
2 \text{CuSO}_4 + 3 \text{CaCO}_3 + 7 \text{H}_2\text{O} = 2 \text{CuCO}_3\cdot\text{Cu(OH)}_2 + 3 \text{CaSO}_4 + 2 \text{H}_2\text{O} + \text{CO}_2
\]

The writer has secured some data on this subject of neutralization and its effect on precipitation.

Finally Lindgren, Gratton and Gordon show chalcolite in coal, and give reduction by carbon as the cause of its deposition. Their equation is

\[
4 \text{CuSO}_4 + 5 \text{C} + 2 \text{H}_2\text{O} = \text{Cu}_2\text{S} + \text{H}_2\text{SO}_4 + 5 \text{CO}_2
\]

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II—Econ. Geol. 8, p. 407-433, Aug. 1913.

Reference and detail of work done are given under the writer's experiment No. 25.

CHEMICAL OUTLINE OF THIS INVESTIGATION AND REFERENCE TO GEOLOGICAL APPLICATIONS.

The results of this research, from the point of view of the chemist, may be considered as having to deal with two classes of metal-bearing solutions, (A) Solutions in which the copper is in the form of an electrolyte, and (B) Solutions of colloidal copper sulfides.

Taking up the matter covered by the first heading we come at once to oxidation. This was investigated early in the work. Failure to secure the solution of copper as a sulfate in more than traces, thru the agency of dissolved oxygen, (see exp. No. 1, No. 2, No. 3, No. 4 and No. 6) and great success in securing such when alternate drying and moistening were resorted to, (experiment No. 5), fully confirmed the views of Spencer1 that dissolved oxygen probably cannot produce enough copper sulfate to form deposits of any consequence, and that in the zone of leaching we must have sufficient porosity to admit atmospheric air. The conditions under which the writer secured strong oxidation are those of the mining districts of the Southwest, where torrential rainfall is followed by long period of drought.

1—Econ. Geol. 8, p. 631.
This work has shown that carbonate ores may, thru the agency of carbonated meteoric waters, go into solution or suspension and thus may readily contribute to the supply of copper in descending waters. (Experiments No. 7 and No. 8.)

Probably one of the earliest recorded syntheses of chalcocite is that of Winchell\textsuperscript{II} and Tolman. They used acid copper sulfate solution with sulfur dioxide and secured a deposit on pyrite. The author has confirmed this result and has shown the great importance of impurities in the pyrite, (Experiment No. 9), and he has also established the fact that sulfur dioxide plays the role of a strong reducing agent, as thru its use he produced a crystalline cupro-cupric sulfite, (Experiment No. 10). In all solutions in which he used sulfur dioxide the precipitation of copper sulfide was greatly increased. (Experiment No. 20.) It is interesting to note in this connection that in the presence of ferrous sulfate, which generally has been thought to be an almost indispensable agent for the securing of chalcocite precipitation, the chalcocite is not as readily precipitated as in the absence of ferrous sulfate, but that the introduction of sulfur dioxide into such solutions containing ferrous sulfate, greatly increases this deposition. (Experiment No. 20.) Acidity is shown to be less favorable than neutrality or alkalinity. As will be

shown hydrogen sulfide is a powerful dispersing agent and is inimical to the deposition of the sulfides. Sulfide dioxide by its action on hydrogen sulfide, \(2 \text{H}_2\text{S} + \text{SO}_2 = 2 \text{H}_2\text{O} + 3 \text{S}\), tends to overcome this dispersing action and in this manner also favors precipitation.

Doelter is impressed with the fact that practically all substances are colloidal when first precipitated, yet that the crystalline condition seems to be the stable one in nature. He reviews the work and ideas of others and seems to conclude that what we call amorphous substances are very close to the crystalline. He speaks of the fact that the tendency to pass from the colloidal to the crystalline condition is greatly increased by pressure, shock, light and influence of Rontgen rays and radium rays and describes his own very successful work in producing crystalline material by means of shaking, long continued heating and particularly by means of pressure, and concludes that many minerals whose formation was once thought to require high temperatures may easily be formed without such temperatures. The ideas brought out in the paper seem particularly germane to the deposition and formation of massive chalccocite.

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Thru shock the cupro-cupric sulfite crystals mentioned above were produced by the author.

This fact that all precipitations are first colloidal was found to apply particularly to the sulfides of copper, and we now consider such colloidal solutions, our second main (B) topic of investigation. Under this is taken up: 1st, The formation of colloidal amorphous copper sulfides; 2d, The chemical conduct of amorphous copper sulfides, and, 3d, The physical conduct of amorphous copper sulfides.

This work shows clearly that colloidal copper sulfides can be formed from sulfate solutions and from the dispersion of crystalline or massive sulfides.

Taking up the formation of the sulfides from sulfate solution is was found that this could be brought about by any one of four methods.

Hydrogen sulfide, as is well known, is very efficient, and that it is quite possible geologically to have this hydrogen sulfide in the earth was shown in experiments No. 22 and No. 23. Hydrogen sulfide was produced by the action of water on pyrite and it was also produced by the action of very dilute acid on pyrite, chalcopyrite and bornite the relative development of hydrogen sulfide decreasing as the iron content of the minerals decreased.

Amorphous sulfur was found to precipitate a sulfide of copper from copper sulfate solution, this action being very rapid with fresh amorphous sulfur, prepared
by the action of hydrogen sulfide on sulfur dioxide, freed from the gases and suspended in water, yet falling off rapidly as the sulfur aged. Experiment No. 19 shows this action, the curve being very striking. That sulfur should have this action is not at all surprising. The equation, $3 \text{S} + 2 \text{H}_2\text{O} = 2 \text{H}_2\text{S} + \text{SO}_2$ is known to be reversible. Stokes' work readily accounts for the presence of amorphous sulfur in the zone of precipitation.

Several instances of production of amorphous copper sulfides by the action of sulfide minerals, particularly pyrite, on copper sulfate solution are shown in experiments No. 20, No. 21, and No. 35. Formerly this was thought to be the one method of sulfide production.

Finally, amorphous copper sulfides were produced by the action of a thiosulfate solution on one of copper sulfate. This action at first formed the double thiosulfate of sodium and copper which then decomposed to give a mixture of amorphous cupric and cuprous sulfides. Details of this work and references showing this action to be possible geologically are given experiments No. 18 and No. 24.

By the dispersion of crystalline and massive sulfides, the colloidal forms are very readily produced, and most effective for this dispersion is the agent hydrogen sul-

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1—Econ. Geol. 2, 14-23, 1907.
fide. This disperses the crystalline or massive material in acid solution, (Experiment No. 43, No. 26, No. 27 and No. 28), and enormously more effectively in alkaline solutions, (Experiment No. 44, No. 38, No. 39, No. 40 and 41). The dispersion increases in a series chalcopyrite, bornite, covellite and chalcocite as the copper content of the minerals increases, which indicates a strong tendency for the sulfide of copper to migrate away from a sulfide of iron. Thus we see why chalcocite and chalcopyrite are so abundant, the former the one to migrate most and the latter to resist this action the longest.

It seems probable that the hydrogen sulfide forms an unstable compound with the copper sulfide. Linder and Picton\(^1\) have shown that the dispersed material has a slightly larger sulfur content than the undispersed.

Free amorphous sulfur has the effect of dispersing a massive or crystalline sulfide. This is very forcefully shown in Experiment No. 13. In about four months a lump of chalcocite weighing seven grams lost over one gram in weight when sealed in a flask in contact with this amorphous sulfur.

Now, coming to the second part of our main heading (B) we may consider the chemical conduct of amorphous copper sulfides. Probably the most im-

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portant point in this investigation is concerned with the
discovery of the fact that at ordinary temperatures cu-
pric sulfide in contact with water spontaneously decom-
poses to cuprous sulfide and sulfur, \(2 \text{CuS} \rightleftharpoons \text{Cu}_2\text{S} + \text{S}\). (Experiments No. 31, No. 12, No. 15, No. 17,
No. 18, No. 24, No. 26, No. 27, No. 29, No. 30, and
No. 33.)

Fraulein Wassjuchnowa\(^\text{II}\) showed that this action
takes place in the dry state at temperatures above 500
degrees.

From the number of experiments cited one can see
that this transformation took place very readily. It did
not, however, take place as readily in the presence of
ferrous sulfate as in its absence, (Ex. No. 14 and Ex.
No. 16), and in those cases in which a large excess of
free sulfur was used no large amount of \(\text{Cu}_2\text{S}\) was
obtained in the length of time which the material stood.
\(\text{Cu}_2\text{S}\) was also produced by use of sodium arsenite.
This is described and discussed under experiment
No. 32.

Finally coming to the physical conduct of the amor-
phous copper sulfides it is to be noticed that there is a
tendency for them to accrete on certain minerals, and
that they deposit from their dispersed condition when
those agents which favor dispersion have their effects
dlessened. In experiment No. 13 it was found that

\(^{\text{II}-\text{Zeitschr. f. Electrochemie 19, No. 22, p 902, 1913.}}\)
some sulfide had gathered on chalcopyrite and in No. 29 a heavy dense growth was noted on bornite. Because of the relatively very much greater accretion on the bornite it would seem that this mineral may become chalcocite thru continued addition of copper sulfide, (See also Ex. No. 42). The same may be said of chalcopyrite tho in the case of this mineral the action would be very much slower. The fact that it is practically impossible to find a chalcocite which is free from specks of bornite gives additional weight to this view. This again is checked by microscopic evidence produced by Tolman and Ray.  

The author was able to cause copper sulfide to accrete on sphalerite, (Ex. No. 30), and also on chalcocite as Cu₄S the phenomenon in this case being caused by the removal of a dispersing agent, H₂S.

The effect of the removal of this dispersing agent H₂S is well brought out in Experiment No. 33. Here both crystalline, (apparently), and amorphous sulfides were thrown down as the hydrogen sulfide was released. Moreover with a reduction of alkalinity, such as in nature could be accounted for by descending acidic waters mingling with the alkaline solutions, a precipitate of the amorphous sulfide was produced.

With the complete elimination of hydrogen sulfide as is seen in experiment No. 41 and as seen in tendency

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1—Unpublished manuscript.
in experiments No. 38 and No. 40, crystalline chalcocite was produced.

The above discussion throws some light on the question as to why chalcocite and chalcopyrite are the chief copper minerals of our mines.

This investigation was started with the view of determining why such should be the case, and it ends apparently with finding no reason why such should not be the case. All the processes involved in enrichment tend toward the formation of chalcocite. Only under particular conditions, (absence of hydrogen sulfide or presence of sulfur dioxide), would this be crystalline.

It is nevertheless evident that chalcocite, either crystalline or amorphous is the ultimate stable form toward which all copper-bearing minerals tend when in contact with solution. All other copper sulfide minerals are to be looked on as unstable. Of these chalcopyrite shows in all reactions a very low reaction rate which accounts for its apparently greater stability.

**PREFACE TO EXPERIMENTS.**

From the beginning of the work the author acted on no preconceived opinions but attempted to get data on any phase of the subject by means of any experiment which seemed to have a fair chance of being productive, and in all over sixty experiments were tried.

Each experiment was carried on carefully but with
attention to exactness commensurate with the expected results. This may be illustrated by the statement that the cyanide method was used in determining copper in some experiments, while in others gravimetric determinations were made on a very sensitive button balance, and all reagents were made up of selected crystals from C. P. analysed chemicals.

Duplication of the work will undoubtedly give closer figures at some points. It probably will alter no principle established here from experimental basis.

All experiments were carried on in the light and at ordinary laboratory temperature except as otherwise stated.

Experiment No. 1.—Study of oxidation of pyrite and of a pyrite and chalcocite mixture.

In belief that more light could be thrown on the subject of oxidation of pyrite and of pyrite and chalcocite mixtures, if such were in the form of very finely divided material which would be kept in water, this water being always saturated with oxygen, 100 grams of very pure pyrite which had been put thru a 200-mesh sieve were put into a long, narrow separatory funnel and covered with water. A tube was led into this and every third day oxygen was allowed to bubble thru the water for 24 hours.

At the end of the first 12 hours evidence of oxidation was seen. This continued to increase. Ferrous sulfate gathered in solution at the bottom of the cylinder and an "iron hat" formed at the top. This oxidation did not go on as fast as it had been expected to, and in view of other data obtained, (see Experiment No. 5), the
passage of oxygen was discontinued, and the liquid was allowed to pass thru the pyrite and drain into a receptacle once each day. The drainings were poured back into the cylinder daily. *This treatment increased the oxidation.* At the end of 5 months the liquid contained 0.0393 g SO₄ and 0.0147 g H₂SO₄. The balance of the sulfate radical was distributed between ferrous and ferric iron.

A mixture of 50 grams 200 mesh pyrite and 200 mesh chalcocite which had been treated in the same way, except for the passage of oxygen gave at the end of 5½ months a solution which contained merely a trace of copper.

In view of the large amount of copper obtained by Buehler and Gottschalk, the idea suggested itself that as the cylinder was very long a precipitation of the copper sulfide might have taken place in the lower part of this cylinder. This was not investigated.

**Experiment No. 2.—Study of oxidation in the presence of pyrite, manganese dioxide and solution of sodium chloride.**

With the view of securing some data on the idea suggested by Lane, that copper minerals may go into solution in the form of chlorides, 50 grams of 200 mesh pyrite, 50 grams 200 mesh chalcocite and 0.5 gram 200 mesh manganese dioxide were put into a long, narrow separatory cylinder, covered with N/20

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I—A very fine, exceedingly pure specimen furnished by Mr. A. C. Luhrs of Butte, Mont. Polished sections showed only mere traces of bornite.


NaCl and treated with oxygen, etc., exactly as in Experiment No. 1.

At the end of 5½ months only a trace of copper had gone into solution, tho much manganese had gone into solution and had redeposited, evidently as hydrated oxide.

**Experiment No. 3.—Study of solution of chalcocite.**

A mixture of chalcocite and pyrite was treated as described in experiment No. 1. The water was kept saturated with oxygen for two months and then for 3½ months the tube was drained daily. Merely a trace of copper was found in solution.

**Experiment No. 4.—Attempt to secure copper in solution as a bicarbonate.**

In the belief that oxidation would be very rapid if the minerals were very finely divided and if they were kept in water saturated with oxygen, 50 grams 200 mesh chalcocite, 50 grams 200 mesh pyrite and 50 grams of 60 mesh marble were put into a separatory funnel and treated as described in Experiment No. 1. Evidently the marble immediately used up any trace of sulfuric acid that was formed as no copper went into solution.

**Experiment No. 5.—Study of oxidation of chalcocite.**

Spencer\(^1\) suggests that oxidation is most rapid where the ore is merely kept moist, and shows theoretically, at least, how the oxidation requires more oxygen than can be obtained from water solution.

Two glass tubes 12" x 2" were nearly sealed at each end. Thru the holes left open 10 grams of 200 mesh chalcocite was introduced into one, and 10 grams of

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\(^1\) Tolman, "Secondary sulfide enrichment of ores" Min. and Sci. Press. 106, p 180, 1913.

\(^2\) Spencer, Econ. Geol. 8, p 631.
200 mesh chalcocite with 10 grams 200 mesh pyrite was put into the other. The mineral was covered with water and the tubes were set aside. These tubes were revolved once each day, thus the mineral was alternately moistened and dried.

At the end of only 29 days the liquid from the first tube showed a strong trace of copper. The liquid from the second tube showed 0.01896 g copper in solution.

Experiment No. 6.—Attempt to secure oxidation data.

In the belief that oxygen which could be liberated from hydrogen peroxide, would be liberated rather evenly throughout a mixture of powdered chalcocite and pyrite thus rapidly producing copper sulfate, a separatory funnel was prepared as described under experiment No. 1. The minerals acted as catalysts for the decomposition of the hydrogen peroxide, and at the end of a day or two about all the available oxygen had been set free. No trace of copper sulfate was secured.

Experiment No. 7.—Study of solution of malachite, azurite and chrysacolla.

As a result of many observations made in and around mines Professor C. F. Tolman, Jr., is of the opinion that these minerals can migrate downward, probably as bicarbonates.

Twelve gas washing bottles were taken. To each of 3 was added 30 cc N/2 potassium carbonate, to each of 3 others 30 cc potassium carbonate with a little sodium silicate, to each of 3 others 30 cc dilute sodium silicate solution, and to each of 3 others 30 cc of water, thus making 3 sets of 4 liquids in each set. 1 gram 200 mesh malachite was added to each bottle in one set, 1 gram 200 mesh azurite to each bottle in the second set and 1 gram 200 mesh chrysacolla to each in the third.

Carbon dioxide was passed thru each bottle for one
month. At the end of this time it was found that there was a trace of copper in the solution in each bottle, the liquids coming from the bottles containing chrysacolla and water, azurite and N/2 potassium carbonate and azurite and water giving the strongest tests.

Experiment No. 8.—Amount of copper going into solution or suspension from azurite and from chrysacolla.

To a gas washing bottle was added 1 gram 200 mesh chrysacolla and 100 cc water, to a second 1 gram 200 mesh azurite and 100 cc water and to a third 1 gram 200 mesh azurite and 100 cc N/2 potassium carbonate.

Carbon dioxide was passed thru each bottle very slowly for 96 days. The liquid in the first bottle was filtered and analysed. It contained 0.0004 g copper.

The liquid in the second contained 0.0007 g copper and the liquid in the third 0.0007 g.

Experiment No. 9.—Synthesis of chalcocite. (Winchell-Tolman experiment).

The experiment as described by these authors was repeated.

A lump of exceedingly pure pyrite was taken and placed into a solution of copper sulfate (1 cc = 0.019665 g Cu) whose acidity with sulfuric acid was N/20. This was then saturated with SO₂ and the whole was sealed in a bottle.

At the end of 18 days a faint trace of dark material appeared on some of the surfaces of the pyrite. At the end of one month one end of the pyrite was strongly coated while the other was as bright as ever. At the end of 5 months a uniform black coating covered the pyrite.

Experiment No. 10.—Attempted synthesis of chalcocite. (Winchell-Tolman experiment modified).

Because the tarnishing of the pyrite was so long delayed in Experiment No. 9 it was believed that impurities were necessary for the formation of chalcocite, and as the analysis of the pyrite used by Winchell and Tolman showed the presence of zinc, a crystal of sphalerite was wired to a piece of very pure pyrite with platinum wire. This couple was then put into a solution such as was used in No. 9 and the flask was sealed.

At the end of 46 days no marked change could be seen. On the 47th day the flask was accidentally subjected to great jarring and on the 48th day small ruby red crystals were seen. These were prismatic, had high relief, were pleochroic and had parallel extinction.

As they gave all the tests for cupro-cupric sulfite as described by Segerbloom in his “Table of Properties” they were undoubtedly cupro-cupric sulfite.

This experiment confirms the belief of Winchell and Spencer that $SO_2$ plays the role of a strong reducing agent. It also tends to confirm the opinion of Lindgren that pyrite does not precipitate $Cu_2S$ or $CuS$ while zinc blende is present. Neither sulfide formed after months of standing.

Experiment No. 11.—Effect of jarring on crystallization.

The experiment as described under No. 10 was repeated. The sealed bottle was allowed to stand for nearly double the length of time which the bottle stood in No. 10. No crystals appeared. The bottle was then

subjected to considerable jarring and crystals appeared within a few hours.

Experiment No. 12.—Production of cuprous sulfide.

H$_2$S was passed thru a solution of CuSO$_4$ until all of the copper had precipitated as CuS. The liquid was filtered off and the precipitate was surrounded with H$_2$S water, and sealed in a flask with an atmosphere of H$_2$S.

At the end of 24 hours a ring of what was, from its color, covellite streaked the flask at the surface of the precipitate. At the end of a few days this covellite was not noticeable.

The flask was opened at the end of 130 days. The residue showed an abundant separation of sulfur when examined with the microscope.

This residue was washed, dried, extracted with CS$_2$ and 0.7173 g was taken for analysis. This gave 0.1723 g sulfur thus leaving 0.5450 g copper.

0.7173 g Cu$_2$S contains 0.5729 g Cu.

On the basis of the copper content this residue contained 70.99% Cu$_2$S.

Experiment No. 13.—To observe transference, substitution or growth.

A lump of chalcocite weighing 7.6332 g and a lump of chalcopyrite weighing 3.2458 g were sealed in a flask with 50 cc copper sulfate solution, (1 cc = 0.039331 g Cu), 50 cc ferrous sulfate solution, (1 cc = 0.026243 g Fe), and a large excess of amorphous sulfur, (2 days old) and an atmosphere of carbon dioxide.

A decided growth of black material was observed on and in contact with the chalcocite at the end of two weeks.

The flask was opened at the end of 137 days. The
The chalcopyrite lump remained attached to the bottom of the flask. This lump was covered with a black velvety film which stood out in ridges on the lump. The film, (very loosely attached), was removed and the lump was weighed. It weighed 6.5745 g—a loss of 1.0587 g.

This film and the general residue in the flask were analysed. Both contained iron which made exact results as to Cu₂S and to CuS of little value. The fact that the film had a copper content of 69.03% Cu, while the general residue contained 64.73% showed the greater copper content to be nearer the lump of chalcopyrite.

It seems probable that in the presence of the excess of amorphous sulfur that the sulfide was mostly CuS tho some Cu₂S was probably present in the film.

The chalcopyrite lump was loose and came out of the flask readily. It weighed 3.2473 g—a gain of 0.0015 g. A deposit of black sulfide hung tenaciously to the lump and could not be washed off. This would account for the gain, and would indicate that chalcocite can grow on or at the expense of chalcopyrite, but not as readily as on or at the expense of bornite. (See Experiment No. 29).

This experiment was repeated with amorphous sulfur which was one day old, and similar results were obtained. It was also repeated with freshly prepared amorphous sulfur. The final results were much as given above.

Experiment No. 14.—To observe transference, substitution or growth.

As described under No. 13, a lump of chalcocite and a lump of chalcopyrite were put into a solution of copper sulfate mixed with one of ferrous sulfate and sealed in an atmosphere of CO₂.
At the end of 76 days the chalcocite showed a loss of 0.0817 g. The weight of the chalcopyrite was unchanged.

This agrees with the results obtained by Emmons and Grout. They secured no chalcocite deposition by using ferrous sulfate as a reducing agent.

Experiment No. 15.—Production of cuprous sulfide.

Cupric sulfide was washed free from electrolyte and was added to water which had been saturated with H₂S. The whole was sealed in a flask with an atmosphere of H₂S.

It was very noticeable that in the absence of an electrolyte the CuS remained colloidal and dispersed, and for 52 days so remained—no clear liquid appearing above the precipitate. On the 53rd day it was noticed that a clear layer had appeared. In a few days the precipitated was well compacted and lay on the bottom of the flask.

At the end of 75 days the flask was opened. The black precipitate was seen with the microscope to be a mass of well-compacted particles of copper sulfide which were mixed with particles of sulfur.

This precipitate was washed with water and alcohol and then partly extracted with CS₂. The CS₂ gave much sulfur upon evaporation.

0.3921 g of the residue was analysed for Cu and for S. It gave 0.1266 g S and 0.2655 g Cu. 0.3921 g Cu₂S yields 0.3178 g Cu and the same weight of CuS yields 0.2606 g Cu. On the basis of the copper content of the residue there was present 8.57% Cu₂S. It probably contained a larger percentage.

Experiment No. 16.—Attempt to produce cuprous sulfide.

100 cc of copper sulfate solution and 100 cc of ferrous sulfate solution (1 cc equalling 0.33931 g Cu and 1 cc equalling 0.02624 g Fe respectively) were mixed, saturated with H₂S and were sealed in a flask with an atmosphere of H₂S.

At the end of 9 days microscopic particles of sulfur could be seen on the side of the flask. At the end of 120 days the contents of the flask were filtered and the residue was analysed. The residue was found to contain both cuprous and cupric sulfides.

When the filtrate was neutralized with NaOH a black precipitate of copper sulfide came down with the Fe(OH)₂.

Experiment No. 17.—Production of cuprous sulfide.

100 cc of a solution of copper sulfate, (1 cc = 0.039331 Cu), was saturated with SO₂, then with H₂S, then with SO₂ and the whole sealed in a flask with an atmosphere of SO₂.

At the end of 11 days the flask broke inwardly. The material as transferred to a new flask and given a new atmosphere of SO₂ and was sealed.

At the end of 58 days a lump raised above the surface of the precipitate to the height of one-half inch and caused the precipitate to appear as if a lump of ore were embedded in it.

This lump continued to grow, and another came. From the first lump some elevated points, suggestive of horns, struck well into the solution. When the flask was finally opened the lumps being very fragile were destroyed. One of the “horns” remained whole and under the microscope was seen to be a large prism. The residue in the flask consisted of black copper sulfide and sulfur.
When, at the end of 136 days, the flask was opened, the odor of SO₂ could not be observed.

The residue was analysed after having been washed, extracted with CS₂, etc. 0.7065 g gave 0.1714 g S thus leaving 0.5351 g Cu. 0.7065 g of Cu₂S contains 0.5643 g Cu while the same amount of CuS contains 0.4695 g Cu. On the basis of the copper content the residue was 69.16% Cu₂S.

The clear supernatant liquid in the flask was filtered and a lump of zinc was added to the filtrate. A very delicate brownish black cloud gathered in the vicinity of the zinc. This was considered to be a sulfide of Cu, (it gave a strong Cu test), which had been held in solution or suspension by H₂S, (derived probably from S), and that when this H₂S was removed by the partial pressure of the H from the zinc, or the sulfide was precipitated by the electrolytic action of the zinc, this cloud appeared.

Experiment No. 18.—Production of cuprous sulfide by the use of sodium thiosulfate.

The rapid precipitation of Cu by amorphous S led to the suggestion of attempting to get this S from the decomposition of a thiosulfate as Hillebrandel seems to have found thiosulfates in deep mine waters, and as Stokes[II] had shown that this is produced when alkaline solutions act on pyrite.

Three flasks were taken and to each was added 50 cc CuSO₄ solution, (1 cc = 0.039331 g Cu). To the first was added 5 cc N/2 Na₂ S₂ O₃, to the second 25 cc and to the third 50 cc of this thiosulfate solution. Each

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flask was given an atmosphere of CO₂ and was sealed.

In each flask the double thiosulfate of sodium and copper formed at once and then began to decompose. The contents of the first flask were black in 2 days, in the second flask in 5 days and in the third at the end of several weeks.

The formation and decomposition of the double thiosulfate of sodium and copper, which according to DutoitIII has the copper in the cuprous condition, goes in several steps or stages. When the two solutions are mixed tiny yellow prisms are formed. These on standing produced long yellow needles which begin to splinter, even going so far as to be a mass of radiating splinters, and these splinters decompose to a black sulfide, decomposition first taking place at the end of the splinters.

At the end of 71 days the residue in the second flask was washed with water and with alcohol, and was extracted with CS₂ and analysed. 0.5035 g of the residue gave 0.1563 g S thus leaving 0.3472 g Cu.

0.5035 g Cu₂S contains 0.4022 g Cu and this amount of CuS contains 0.3347 g. On the basis of the copper content this residue was 21.74% Cu₂S.

Experiment No. 19.—The effect of age on the precipitating power of amorphous sulfur.

Early in this work it was noticed that freshly prepared amorphous sulfur precipitated copper sulfide from solutions of copper sulfate much more readily than the older material.

In order to bring out this fact somewhat quantitatively H₂S and SO₂ were brought together to produce amorphous sulfur. This when free from the gases was

Curve showing that as amorphous sulfur ages it can precipitate smaller amounts of copper sulfide from a sulfate solution in a given amount of time.
put into suspension with water and more than enough of this sulfur to precipitate a given amount of copper from a sulfate solution was added to each of seven bottles, the same amount being added to each bottle. A given amount of copper sulfate solution was added to one bottle at once, to a second at the end of 24 hours, to a third at the end of 48 hours, etc. Each bottle was then shaken and allowed to stand for 24 hours when its contents were filtered and analysed.

The results of the experiment are shown in the accompanying curve.

Experiment No. 20.—Precipitating power of pyrite in various solutions.

With a view of ascertaining those conditions most favorable for the precipitation of Cu as a sulfide from CuSO₄ solutions by means of pyrite, and with a view of securing more data on the role of SO₂, a given amount of 200 mesh pyrite was sealed in tubes with an aqueous solution of CuSO₄, an aqueous solution of CuSO₄ saturated with SO₂, in a solution of CuSO₄ and FeSO₄, in a solution of CuSO₄ and FeSO₄ saturated with SO₂, and in an acidified solution of CuSO₄, (H₂SO₄ used) which was isotonic with the aqueous solution which had been saturated with the SO₂. These tubes were shaken once daily and at the end of 47 days their contents were analysed.

The results showed that the amounts of copper precipitated increased with the concentration of the CuSO₄ solution, that great precipitation takes place in aqueous solution, and that this is increased by the presence of SO₂. The results also showed that the precipitation in the presence of FeSO₄ is not as great as in its absence, but that SO₂ increases this precipitation in FeSO₄. They showed that the reducing action of SO₂ and not the action of the H ion is responsible for the increased
precipitation in the presence of this substance. Moreover they showed that neutrality or alkalinity are conditions most favorable for precipitation.

Details are given in the following table:

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<th>Water</th>
<th>Water</th>
<th>N/10</th>
<th>N/10</th>
<th>H₂SO₄</th>
<th>CuSO₄</th>
<th>CuSO₄</th>
<th>H₂SO₄</th>
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Experiment No. 21.—Precipitating power of mono and disulfides and of mono and diarsenides.

With a view of ascertaining whether disulfides and diarsenides would precipitate more copper than the mono compounds, a series of minerals were taken, any

* Probably experimental error.
impurities removed mechanically as far as possible, and the minerals were put thru a 200 mesh sieve. One gram of each mineral was sealed in a tube with an aqueous solution of CuSO₄, each tube shaken daily and at the end of 47 days the contents were analysed.

As a result of this experiment it was concluded that no generality was shown. Details are given in the table.

<table>
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<tr>
<th></th>
<th>Cobaltite</th>
<th>Nacellel</th>
<th>Niobite</th>
<th>Chalcocite</th>
<th>Enargite</th>
<th>Bornite</th>
<th>Galena</th>
<th>Alabamite</th>
<th>Stibnite</th>
<th>Pyrargyrite</th>
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<td>Amt. Cu in 25 cc solution</td>
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<td>0.0916</td>
<td>0.1374</td>
<td>0.1374</td>
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<tr>
<td>Amt. Cu in filtrate</td>
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<tr>
<td>Amt. Cupped</td>
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<td>0.0658</td>
<td>0.0285</td>
<td>0.0969</td>
<td>0.0991</td>
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<td>0.0096</td>
<td>0.0334</td>
<td>0.0316</td>
<td>0.0224</td>
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**Experiment No. 22.—Hydrogen sulfide produced by hydrolytic action of water on pyrite.**

Several one-half gram portions of uniform sized pyrite particles, ranging from those which passed thru the 20 mesh sieve and were retained on the 30 mesh to those which passed thru the 200 mesh sieve were sealed, each portion in a separate tube, with 20 cc of water.

These tubes were kept at about 41 degrees for three months, when the tube containing the 80-90 mesh pyrite was opened, its contents filtered and tested colorimetrically with a standard solution of lead nitrate. The test showed the presence of H₂S in solution. Other tubes were afterward tested and their contents gave like results.
Experiment No. 23.—Production of hydrogen sulfide from pyrite, chalcopyrite and bornite.

Duplicate tubes containing:
(a) 0.5 g 200 mesh pyrite and 15 cc N/10 sulfuric acid
(b) 0.5 g 200 mesh chalcopyrite and 15 cc N/10 sulfuric acid
(c) 0.5 g 200 mesh bornite and 15 cc N/10 sulfuric acid

were sealed and placed in a thermostat at about 41 degrees for 51 days.

The contents of one set of tubes were filtered and the filtrates treated with standard lead nitrate solution. All showed the presence of H₂S in solution, the pyrite having set free the most and the bornite the least.

The duplicate set was then put in an autoclave and heated at 195 degrees for two hours. The tube containing the acid and pyrite exploded, the one containing the acid and chalcopyrite cracked and the one containing the acid and bornite remained intact, tho much pressure was developed by the H₂S inside. This H₂S was readily noticed because of its odor.

Experiment No. 24.—Cuprous sulfide from the double thiosulfate of sodium and copper.

Some of the double thiosulfate was prepared from solutions of copper sulfate and sodium thiosulfate. This was washed and sealed in a flask with an excess of water and an atmosphere of CO₂.

After 8 weeks the material in the flask began to gather into large platy masses which stood up on one edge in the solution. This continued until the material in the flask was well compacted into large platy masses. The flask was opened at the end of 123 days, and the
residue was washed with water, alcohol and was dried and extracted with CS₂ and analysed.

0.5170 g of the residue gave 0.1380 g S thus leaving 0.3790 g Cu. 0.5170 g of Cu₂S contains 0.4130 g Cu, while this weight of CuS contains 0.3437 g Cu. On the basis of the copper content of this residue it contained 50.94% Cu₂S.

Experiment No. 25. — Attempt to precipitate copper sulfide by the use of coal.

Lindgren, Gratton and Gordon think that coal has acted as a reducing agent on copper solutions and in their "Ore Deposits of New Mexico" show a beautifully colored plate of chalcocite embedded in coal.

With a view of securing data on this method of precipitation 2 grams of powdered coal were put into a solution of CuSO₄ and FeSO₄ and the whole was saturated with SO₂ and sealed in a flask. Some action was expected in the presence of these three reducing agents.

The flask was shaken daily and at the end of 122 days was opened and its contents were examined. No deposition of sulfide of copper was observed.

Great care was taken to secure a coal which had no pyrite in it. The results of this experiment seem to indicate that carbon alone will not precipitate the sulfide. Had pyrite been present in the coal some sulfide would undoubtedly have been precipitated.

Experiment No. 26.—Production of cuprous sulfide.

A lump of chalcocite weighing 2.2302 g was put into a solution of copper sulfate, (1 cc = 0.039331 g Cu), and this in a flask was connected with an H₂S generator so that the H₂S was always present in the flask.
The H$_2$S was not passed thru the liquid but was allowed to settle upon it.

At the end of 4 days what appeared to be shining crystalline faces were seen on the lump of chalcocite, and in spite of the precipitation of some CuS from solution the liquid appeared to be more blue than ever. This latter phenomenon was attributed to the reflected color of covellite films.

At the end of 100 days the flask was opened. The chalcocite was found to weigh 1.9923 g—a loss of 0.2379 g. The lump was badly pitted.

A film had gathered around the lump. 0.1083 g of this was taken for analysis. It was not extracted with CS$_2$. It gave 0.0299 g S thus leaving 0.0784 g Cu. 0.1083 g Cu$_2$S contains 0.08648 g Cu while the same weight of CuS contains 0.07197 g Cu. On the basis of the copper content of the film, this contained 43.1% Cu$_2$S.

Hard films of the CuS which had originally formed in the flask could be separated from the general residue. These were washed, extracted and analysed. 0.4718 g gave 0.1437 g S thus leaving 0.3281 g Cu. 0.4718 g Cu$_2$S contains 0.3768 g Cu and the same weight of CuS contains 0.3135 g Cu. On the basis of the copper content this film was 23.07% Cu$_2$S.

The general residue in the flask was treated as described above and analysed. 0.6522 g gave 0.2002 g S and 0.4520 g Cu. Thus the Cu$_2$S content of the general residue was 21.28%.

It is to be noted that the chalcocite lump lost weight, that the film on the lump was richer in Cu$_2$S than any other part of the material in the flask, and that the residue near the film was richer in Cu$_2$S than the general residue. Thus we see the tendency of the Cu$_2$S to give up its Cu to nearby CuS, (the equilibrium point
we do not know), and the tendency of the CuS to drop off its S and become Cu$_2$S, or because of the dispersion of the chalcocite the material nearest to the lump became most contaminated with Cu$_2$S.

Experiment No. 27.—Production of cuprous sulfide.

A lump of pyrite weighing 2.7769 g was placed in a flask and kept under conditions as described under experiment No. 26.

At the end of one week a color and luster resembling covellite was noticeable.

At the end of 91 days the flask was opened. The pyrite weighed 2.7755 g—a loss of 0.0014 g.

The residue was washed, dried, extracted, etc. 0.9059 g gave 0.1994 g S thus leaving 0.7065 g Cu. 0.9059 g Cu$_2$S would give 0.7235 g Cu and the same weight of CuS would give 0.6020 g Cu. On the basis of the copper content this residue contained 86.02% Cu$_2$S.

Experiment No. 28.—Production of cuprous sulfide.

Under conditions as described for Experiment No. 26, 1.8291 g covellite was placed in contact with the sulfate solution and the H$_2$S. It cannot be stated just how long the H$_2$S acted on the solution as its supply was accidentally interrupted.

At the end of 100 days the flask was opened. The covellite weighed 1.8241 g—a loss of 0.0050 g.

The residue when treated as described showed a content of 53.19% Cu$_2$S.

Experiment No. 29.—Production of cuprous sulfide and growth on bornite.

Under conditions as described under experiment No. 26, 4.2770 g bornite was placed in a flask with the copper sulfate solution and connected with the H$_2$S generator.
At the end of one week the color and luster of covellite was noticeable.

At the end of 94 days the flask was opened. The lump of bornite was found to weigh 5.5569 g—a gain of 0.2799 g. This growth was noticed largely in the gain in weight as the material which had been incorporated with the lump was so added that the appearance of the lump was hardly altered, except that it was a deeper dark blue than usual. This experiment seems to indicate that the copper sulfide tends to concentrate around the bornite. Thus bornite could turn to chalcopyrite thru a gradual increase in its copper content.

On the same basis (e. i., copper content) as stated in Experiment No. 26 the residue in the flask was found to be 16.62% Cu₂S.

Experiment No. 30.—Production of cuprous sulfide.

A crystal of sphalerite weighing 6.7795 g was placed in a flask under conditions as described in experiment No. 26.

At the end of one week the color and luster of covellite was noticeable.

At the end of 93 days the flask was opened. The lump of sphalerite was found to weigh 6.7745 g—a loss of 0.0050 g, yet a black deposit of copper sulfide hung tenaciously to the sphalerite along crystallographic lines. The whole crystal was not covered with the deposit. It seemed that the sphalerite is an excellent precipitant of the sulfides of copper1 yet that in the case of this particular sphalerite crystal only a portion

1—Prof. A. F. Rogers of Stanford University has produced covellite by the simple heating of a solution of CuSO₄ with sphalerite in a bomb furnace. School of Mines Quarterly, 32, p 298, 1911.
of the precipitated sulfide remained on the sphalerite.

The residue in the flask was treated and analysed as has been already described. It showed a Cu₂S content of 29.57%.

Experiment No. 31.—Production of cuprous sulfide.

(This is the experiment in which the spontaneous change of CuS to Cu₂S and S was first noted.)

A solution of CuSO₄ had H₂S passed thru it until all the copper was precipitated as the sulfide. The precipitate was freed from solution by filtration and then this precipitate was placed in a flask with a lump of chalcocite weighing 2.2861 g and the whole was covered with water and sealed in an atmosphere of CO₂.

After 54 days the flask was opened. The lump weighed 2.1404 g—a loss of 0.1457 g. The residue under the microscope was seen to be made up of particles of black sulfide which were intermingled with particles of sulfur. The residue was extracted with CS₂ (which was known to be free from dissolved S) and the extract evaporated. A yellow residue which by the microscope and by chemical test was shown to be sulfur was left upon evaporation of the CS₂.

The residue from which the sulfur had been extracted was analysed for both S and Cu, and upon the basis of its Cu content was shown to be 43.64% Cu₂S. Thus the transformation of CuS to Cu₂S was shown.

Experiment No. 32.—Production of cuprous sulfide.

A suggestion that sodium arsenite can transform CuS to Cu₂S led to the sealing of 15 g CuS and 30 g sodium arsenite in a flask with an atmosphere of CO₂. 200 cc H₂O was present.

After 55 days the flask was opened and its contents

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were analysed. They showed a Cu₂S content of 49.00%.

This transformation was at first attributed to the sodium arsenite but in view of spontaneous changes obtained in other experiments the amount to be attributed to each cause is problematical.

*Experiment No. 33.—Growth of cuprous sulfide on chalcocite.*

With a view of getting some experimental certification of the formation of chalcocite as suggested by Prof. A. F. Rogers¹¹ a lump of chalcocite weighing 4.5240 g was put into a flask containing CuS and a solution of K₂S which had been completely saturated with H₂S. Testing showed that this liquid held considerable Cu in suspension or solution. A stopper with a Bunsen valve was inserted, thus giving the H₂S which held the copper in suspension, a chance to escape. At the end of three weeks crystal faces were seen shining on the chalcocite lump.

At the end of 85 days the flask was opened and the solution displaced by water. The chalcocite lump was washed with hot water until it was free from alkali. One could see that the lump was coated and that this coating had on it tiny shining spots suggestive of crystal faces. The lump was dried. It weighed 4.5874 g—a gain of 0.0634 g. 32.24 milligrams of this coating were taken for analysis. This gave 8.403 mg of S, thus leaving 23.837 mg of Cu. 32.24 mg of an absolutely pure chalcocite would give 25.748 mg Cu.

This sulfur figure was known to be too high. While C. P. chemicals were used a very noticeable amount of iron oxide was detected in the BaSO₄ precipitate. Furthermore, as no extraction by CS₂ was made it is quite

¹¹—A. F. Rogers, Econ. Geol. 8, p. 781, 1913.
possible that a small amount of S may have been in the material deposited on the chalcocite.

The author has no hesitation in saying that he produced a pure chalcocite from this alkaline solution.

The analysis of the residue in the flask showed that it had a Cu₂S content of 57.63%.

Experiment No. 34.—Attempt to secure microscopic evidence of the change of covellite to chalcocite.

H₂S was passed thru a solution of CuSO₄ until all the copper was precipitated as a sulfide, then some 200 mesh chalcocite was added and the whole was sealed in a flask with an atmosphere of CO₂.

A similar set up was made with 200 mesh covellite.

After 23 days both flasks were opened. Their contents showed in each case that the powdered mineral was embedded in the amorphous sulfide. Free sulfur was noted in each case. Both flasks were again sealed with CO₂ atmospheres and both were opened again at the end of 113 days. The appearances of their contents were much as before, except that a much larger amount of free sulfur could be seen.

Experiment No. 35.—Attempt to find the condition of acidity or alkalinity most favorable for the precipitation of copper sulfide from copper sulfate solutions by means of pyrite.

One gram of 200 mesh pyrite was put into each of 11 tubes, a given amount of CuSO₄ was added to each and then such amounts of water and H₂SO₄ were added as to have the acidity of the solutions in the different tubes as follows: N/15, N/20, N/40, N/80, N/100, N/200 and "neutral". To other tubes NaOH and water was added so that after the amount of NaOH had been added which would theoretically throw down all of the
Cu as Cu(OH)$_2$ the alkalinity of the tubes was N/200, N/100, N/80, N/40.

The contents of those tubes containing the alkali were black at the end of 24 hours.

Each tube was shaken once daily and after 31 days the contents were analysed.

It was found that under condition of acidity there was practically no copper sulfide precipitated.

In the case of the tubes containing the alkali the difficulty presented itself of finding out how much of the black material was copper oxide and how much was a sulfide of copper. It seemed reasonable to believe that as the sulfide precipitates in acidity of 1:8 HCl that if the residue were washed with a 1:10 HCl the sulfide would not be dissolved. This procedure was taken. All of the material dissolved, thus this experiment proved that acidity was not favorable to precipitation by means of pyrite but left the question as to alkalinity unsettled. (See also Experiment No. 20.)

Experiment No. 36.—Neutralizing power of different rocks.

As altered rock is always found in veins, etc., showing downward enrichment, and as this alteration is particularly kaolinization, and as experiments No. 20 and 35 indicated that acidity is not favorable to the deposition of the sulfides a series of typical rocks was taken and the neutralizing effect of each rock on acid solution was determined. Each rock was put thru a 200 mesh sieve. 5 grams of each was put into a bottle and covered with 50 cc N/10 sulfuric acid. The bottle containing each was shaken once each day. After 30 days the liquid in each bottle was analysed.
Details are given in the accompanying table:

<table>
<thead>
<tr>
<th>Rock and Locality</th>
<th>Principal minerals in order of abundance</th>
<th>Amt. N/10 H₂SO₄ used</th>
<th>Amt. N/10 H₂SO₄ neutralized</th>
<th>Relative neutralizing power of the rock, granite being taken as 1.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Franciscan Limestone, Back of Stanford University</td>
<td>Calcite, Chalcedony</td>
<td>50 cc</td>
<td>48.5 cc</td>
<td>10.32</td>
</tr>
<tr>
<td>Basalt, Stanford University Quarry</td>
<td>Plagioclase, Augite</td>
<td>50 cc</td>
<td>23.4 cc</td>
<td>4.98</td>
</tr>
<tr>
<td>Diorite, Trinity River, So. Fork, near Low Gap, Cal.</td>
<td>Orthoclase, epidote</td>
<td>50 cc</td>
<td>16.7 cc</td>
<td>3.65</td>
</tr>
<tr>
<td>Franciscan Shale, Tesla, Cal.</td>
<td>Clay, quartz</td>
<td>50 cc</td>
<td>15.1 cc</td>
<td>3.21</td>
</tr>
<tr>
<td>Rhyolite, Alum Rock, San Jose, Cal.</td>
<td>Glass (devitrefined and silicified)</td>
<td>50 cc</td>
<td>11.0 cc</td>
<td>2.34</td>
</tr>
<tr>
<td>Hornblende Andesite, Marysville, Cal.</td>
<td>Glass, hornblende</td>
<td>50 cc</td>
<td>11.0 cc</td>
<td>2.34</td>
</tr>
<tr>
<td>Granite, Santa Lucia, Cal.</td>
<td>Orthoclase, quartz</td>
<td>50 cc</td>
<td>4.7</td>
<td>1.00</td>
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</table>

Experiment No. 37.—Attempt to produce cuprous sulfide.
A lump of pyrite weighing 4.2663 g and some care-
fully washed CuS were put into a solution of N/2 K$_2$CO$_3$ and the whole was sealed in a flask with an atmosphere of CO$_2$.

At the end of 95 days the flask was opened. The pyrite was found to weigh 4.2653 g—a loss of 0.0010 g. The pyrite was bright and shiny.

The residue when examined with the microscope showed that some sulfur had separated. This residue was lost in the process of analysis. Judged from its appearance, however, this residue was made up of CuS with a small amount of Cu$_2$S.

Some iron was observed in the liquid contained in the flask.

Experiment No. 38.—Attempt to produce cuprous sulfide in alkaline solution.

CuS and a fairly concentrated solution of K$_2$S were heated for some time, finally at the boiling temperature of the solution. The liquid was allowed to settle until clear. The clear supernatant solution was placed in a desiccation tube over sulfuric acid, and this tube was evacuated and sealed.

At the end of 24 hours a brownish mass had formed. This with the hand lens had the appearance of a lot of tiny needle-like crystals. With high magnification these were seen to be long curved brownish hairs. When viewed with polarized light these hairs failed to show crystalline characteristics. Some sulfur could be seen along with these hairs.

The liquid in the desiccation tube constantly grew more concentrated as the sulfuric acid took up the water vapor and at the end of 113 days these were seen to be masses of amorphous copper sulfide, probably cuprous as other experiments showed.
Experiment No. 39.—Solution and deposition of chalcocite.

Some 200 mesh chalcocite was put into a strong solution of KOH and $\text{H}_2\text{S}$ was passed thru the solution, which was kept cold, until the solution was saturated. The heavy dense chalcocite became colloidal, occupying 4 or 5 times its original volume. This whole solution was heated in an autoclave at 180 degrees for 1½ hours. The solution became deep yellow. This was filtered thru glass wool and was then gently heated on an air bath. $\text{H}_2\text{S}$ came off slowly and finally black lumps gathered on the bottom of the container.

A drop of this filtrate just mentioned was evaporated on a glass slide and its residue under the microscope showed black needles or prisms together with some yellow prisms suggestive of the double thiosulfate of sodium and copper. Some of this same filtrate was neutralized with dilute sulfuric acid whereupon a brownish black precipitate was formed. This gave a strong test for copper.

Experiment No. 40.—Solution and deposition of a copper sulfide derived from covellite, bornite and chalcopyrite.

Experiment No. 39 was repeated with each of the minerals named above.

The filtrate from covellite behaved as did the filtrate obtained in No. 39. A few of the hairs as obtained in Experiment No. 38 were noted on the glass wool filter.

The filtrate from the bornite behaved as did that from the covellite. It did not contain as much copper, however. A larger number of the hairs were obtained from the bornite. These gave a very strong test for copper.
The filtrate from the chalcopyrite gave only a faint test for copper.

It appeared that the \( H_2S \) had dispersed these minerals in the following descending order: chalcocite, covellite, bornite and chalcopyrite and that the alkaline liquid with its \( H_2S \) had dissolved copper from these minerals in this same order.

The dispersion and solution of the chalcocite was much more marked than were these phenomena with the other minerals.

*Experiment No. 41.*—Production of chalcocite crystals, and the effect of hydrogen sulfide as a dispersing agent.

2 grams of 200 mesh chalcocite were put into a tall Nessler tube and covered with 25 cc of a KHS solution. \( H_2S \) was passed thru the solution for an hour.

The same amount of chalcocite was put into a tube and covered with 25 cc of a \( K_2S \) solution. Hydrogen was passed thru this solution for an hour.

The same amount of chalcocite was put into another tube and was covered with 25 cc of a solution of equal parts of \( K_2S \) and KOH. Hydrogen was passed thru this liquid for an hour.

Each solution contained approximately the same amount of potassium. The gases were passed thru the liquids at approximately equal rates.

The material in the first tube showed strong dispersion, that in the second showed \( 1/4 \)th to \( 1/6 \)th as much and that in the third showed scarcely any.

At the end of 10 days it was noted that the material in the second tube had risen, and that a small mass having something of a crystalline appearance floated on the surface. From this hung down a growth resembling a fox's tail.
This growth and “tail” were washed, and the latter examined with a microscope. This “tail” was seen to be made up of a lot of stiff black, non-transparent hairs or rods. One or two were especially large and showed truncated ends. The angles on the ends of one rod were measured. Beginning on the right side of the rod or prism and going anti-clock wise the angles were approximately 37, 51 and 56 degrees. (See figure.)

The floating mass was washed again with water and with absolute alcohol and was then dried. It was found to weight 13 mg. It was analysed with the greatest care and was found to yield 2.57 mg S and 10.40 mg Cu. 13 mg of Cu₂S would yield 10.39 mg Cu. The floating mass was certainly cuprous sulfide. The author assumes that the crystals were of the same composition.

Experiment No. 42.—Elimination of iron from bornite.

Work done simultaneously with this by Tolman and Ray has shown that, from microscopic evidence, we may believe that bornite gives off some of its iron and becomes chalcocite.

With a view of actually securing some data on this idea samples of bornite were analysed for their iron contents. These powdered samples were placed in water in sealed tubes and were heated in an autoclave at a temperature of about 175 degrees, for two hours. The contents of the tubes were filtered and the residues were washed with N/10 sulfuric acid as this acid does not attack bornite to any appreciable extent.

The filtrates and washings gave strong iron tests. Quantitative tests were not made.

Experiment No. 43.—The dispersing power of hydrogen sulfide in acid solution.

To determine the dispersing power (relative), in acid
solution four gas washing bottles were taken. To the first 2 grams of 200 mesh chalcocite was added, to the second 2 grams of 200 mesh covellite, to the third bornite and to the fourth chalcopyrite. 50 cc of N/10 sulfuric acid was added to each bottle. H₂S was passed thru the 4 bottles for a few minutes each day for 4 weeks.

At the end of this time all the minerals were slightly dispersed, this dispersion appearing most in the chalcocite and least in the chalcopyrite.

The contents of the bottles were filtered. All of the filtrates appeared clear, tho that from the chalcocite gave the merest suggestion of being less clear than the others.

Lumps of zinc were added to each filtrate. In the chalcocite filtrate a brownish black cloud appeared at once around the zinc. This cloud did not appear in the others tho they became less clear.

The residues which were left in the tubes containing the filtrates after the zinc had gone into solution were dissolved in nitric acid and this acid was more than neutralized with NH₄OH. All showed traces of copper the test being strongest in the residue from the chalcocite filtrate and least in that from the bornite filtrate. A very noticeable amount of Fe(OH)₃ was observed in the tube containing the residue from the bornite filtrate.

This experiment showed clearly that H₂S disperses these minerals in acid solution and that the dispersion is in a measure proportional to their copper content. It also shows that in acid solution and in contact with H₂S bornite gives up some of its iron.

Dispersion in acid solution is very much less than in alkaline.
Experiment No. 44.—Order of dispersion of minerals in alkaline solution.

Under conditions as described in experiment No. 43 chalcopyrite, bornite, enargite, covellite, and chalcocite were covered with dilute KHS and were treated with H₂S for 4 days. All dispersed. An attempt was made to show that the iron to copper ratio in the dispersed material was less than in the mineral from which the material was dispersed. Because of limited time colorimetric methods were used. These were not suitable because of the iron content of even the best reagents. This experiment did show that these minerals were dispersed in the reverse order as given, and dispersed in such a finely divided form that the material readily went thru a filter. This dispersion was very strong.

SUMMARY.

While the work included in this present paper has been largely exploration along a variety of lines, and while it can scarcely be said that the investigation has been carried in any particular direction to such an extent as to ultimately determine with quantitative accuracy the importance of the various factors studied in respect to the enrichment of sulfide ores, it seems nevertheless true that certain heretofore little understood and in many cases unsuggested processes do enter into the general problem of the chemistry of the enrichment of the sulfide ores of copper. Perhaps the whole results may be summarized in the following general statements:
(1) Enrichment does not necessarily depend upon oxidation and leaching processes and upon the formation of electrolytic solutions of copper, such as the sulfate or chloride, but may perfectly well occur thru the intervention of the colloidal dispersion and solution, and subsequent deposition, either in the amorphous or crystalline form of already existing sulfides of copper. This applies to upward enrichment.

(2) The most effective dispersing agent which is likely to be found in nature is hydrogen sulfide, which may be produced in a variety of ways, notably from any sulfide minerals by contact with dilute acids or even with pure water. It is probably assisted by carbon dioxide.

The spontaneous dispersion of already existing sulfides is least in acid solutions and enormously greater in alkaline solution, from which it is to be expected that such operations will be of greater importance at considerable depth.

Free sulfur may also serve as a dispersing agent, and the same is true of higher sulfur-bearing minerals as pyrite, which acts to all intents and purposes as potential sulfur.

(3) The conditions favorable to the deposition of dispersed copper sulfides are the removal or absorption of the dispersing agent. Thus loss of hydrogen sulfide from a colloidal suspension of sulfides of copper
tends to flocculate and precipitate these sulfides. The condition for the appearance of crystalline rather than amorphous deposits seems to be the fairly complete removal of hydrogen sulfide which is readily accomplished by sulfur dioxide which accounts for the success of the Winchell-Tolman experiments.

(4) There seems to be no doubt but that cuprous sulfide (chalcocite) is the most stable of all of the copper sulfide minerals, in contact with solution, at least, and that there will be a general tendency for all copper sulfide-bearing minerals to eliminate cuprous sulfide, and for all sulfide precipitates from electrolyte solutions to spontaneously go over into the cuprous form. This tendency may possibly be more or less hampered by the presence of an excess of sulfur, such sulfur may be gradually eliminated by solution in alkaline waters, and even by reaction with pure water, and transportation to long distances may be accomplished.

Thus in the following figure, (diagramatic), if AB represents a mass of sulfide-bearing ore, and alkaline waters charged with hydrogen sulfide gradually percolate from B toward A, there will occur first a zone of dispersion (Zone 2). Upon elimination or absorption of hydrogen sulfide, there will follow a zone of amorphous chalcocite enrichment (Zone 3).

Following this with more complete elimination of hydrogen sulfide, accomplished perhaps by the appear-
ance of sulfur dioxide as a result of limited oxygen supply, crystalline chalcocite will appear, (Zone 4).

| Appearance of $SO_2$ thru small supply of $O_2$. Deposit of crystalline chalcocite. | Zone 4 |
| Elimination or absorption of $H_2S$. Deposit of amorphous chalcocite changing to massive. | Zone 3 |
| Copper Sulfide Bearing Zones. |
| Zone of dispersion. | Zone 2 |
| Alkaline water charged with $H_2S$. | Zone 1 |

(5) There seems to be also a curious physical affinity on the part of the chalcocite to draw toward itself any freshly formed cuprous sulfide, and to cause amorphous deposits of the new sulfide on the already present chalcocite. Already present chalcocite may also seemingly exert a sort of induced acceleration on the reaction between electrolytic solution of copper salts and free sulfur, resulting in the ultimate formation of chalcocite. (See Experiment No. 13.)

**APPENDIX.**

After the author had finished the experimentation which led to the conclusions given in this paper, there remained many questions bearing on this work which
still needed explanation. Time was not available for anything like a complete investigation of these questions but the author was able to make some interesting experiments which are suggestive. Since the conclusions as derived from these experiments have only a suggestive value the conclusions and the experiments are here given in this appendix.

In this paper it has been shown that copper sulfide is kept colloidal by hydrogen sulfide and with hydrogen sulfide the colloid, (or colloids), migrate very readily. Naturally the question arose as to whether this dispersion and migration would be assisted or retarded by carbon dioxide, which is known to be present in many of the springs which give off hydrogen sulfide, and it is found in some mine gases which are given off from the cracks in the wall rocks of the mines. Quite naturally, too, it was asked whether some solutions would favor dispersion and whether others would tend to flocculate the colloidal copper sulfide. What effect would certain wall substances have on the colloid? Moreover it has been asked very frequently, what becomes of sulfur if such at any time be set free in the process of ore deposition.

One sees in experiments No. 45, No. 46, and No. 47 some interesting facts which suggest possible answers to these questions.

We notice here that those solutions containing small
amounts of sodium and potassium salts are most favorable for the dispersion of the colloidal copper sulfide, and believe that our deep-seated ore solutions actually are rich in potassium. We see that calcium and aluminium salts coagulate the colloidal copper sulfide and notice particularly that even the most insoluble aluminium compound, (dehydrated aluminium oxide), is exceedingly effective as a floculant. We notice that calcium carbonate floculates the colloid but that in the case of this colloidal copper sulfide being precipitated by calcium carbonate the colloid may be slightly dispersed again by means of hydrogen sulfide, some calcium going into solution, altho this dispersion is by no means as great as in the presence of alkali salts.

If one makes but a hasty examination of a good treatise on ore deposits* he will notice many examples of copper sulfide deposits embedded in limestone, some of the limestone having been replaced, and he will find that equally numerous are the data showing copper sulfide deposits in contact with, and embedded in, shale, gouge or other argillaceous material. May not the results of these experiments partially account for such deposits?

As is shown in Experiment No. 46, carbon dioxide is a most excellent agent for dispersing colloidal sulfur,

*—Very strikingly shown in many illustrations in Lindgren, "Mineral Deposits".
Theo, as the author has found, it assists the dispersion of copper sulfide when hydrogen sulfide is present and precipitates the copper sulfide when hydrogen sulfide is absent. We may not think that carbon dioxide has aided in bringing the copper sulfide up to where it is deposited, has caused the colloidal sulfide to flocculate when the hydrogen sulfide has escaped, has been assimilated or has ceased to flow, and then having caused the copper sulfide to deposit, the carbon dioxide has carried on any excess sulfur to be used up in pyritization of the surrounding country rock and perhaps in many other ways?

More thorough investigation of all the lines suggested in this paper will be pushed forward as rapidly as possible.

Experiment No. 45.—Flocculation of colloidal copper sulfide.

Some colloidal copper sulfide was prepared, and by washing was carefully freed from the presence of any electrolyte.

Equal portions of this colloidal material were put into flasks and were treated with dilute solutions of NaCl, KCl, CaCl₂ and AlCl₃ and with powdered CaCO₃ and Al₂O₃ in water as can be seen in detail in the accompanying table.

It was found that the presence of NaCl and KCl increased the amount of the sulfide which was held in suspension, whereas CaCl₂, CaCO₃, AlCl₃, and Al₂O₃ caused the colloid to flocculate. This flocculation was accomplished very rapidly by the Al₂O₃.
A similar set of solutions with the colloidal sulfide were treated with H$_2$S. The hydrogen sulfide increased the amount of material which was held in suspension by water, NaCl solution and KCl solution. Also it carried into suspension some copper sulfide which had been flocculated at CaCO$_3$, thus indicating that the limestone may tend to precipitate copper sulfide from ore solutions having this in suspension, the colloids in the presence of H$_2$S may, to some extent at least, migrate thru limestone. The H$_2$S failed to disperse the colloid which had been precipitated by CaCl$_2$, AlCl$_3$, and Al$_2$O$_3$.

A similar set of solutions with the colloidal copper sulfide were treated with a mixture of H$_2$S and CO$_2$. The results were much as those produced with the H$_2$S alone, except that not as much of the colloid, which had been precipitated by the CaCO$_3$ was again put into suspension, which could be accounted for by the fact that the CO$_2$ acting on the carbonate undoubtedly produced calcium ions.

This experiment showed most strikingly the great tendency of argillaceous material to flocculate colloidal copper sulfide.
<table>
<thead>
<tr>
<th></th>
<th>Colloidal Cus in water</th>
<th>Colloidal Cus in N/300 NaCl</th>
<th>Colloidal Cus in N/300 KCl</th>
<th>Colloidal Cus in N/300 CaCl₂</th>
<th>Colloidal Cus in N/300 CaCO₃</th>
<th>Colloidal Cus in N/300 AlCl₃</th>
<th>Colloidal Cus with 1 gram Al₂O₃</th>
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<td>Suspended colloid</td>
<td>Remained suspended</td>
<td>More in suspension than in</td>
<td>More in suspension than in</td>
<td>Colloid soon flocculated</td>
<td>Colloid soon flocculated</td>
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<td>H₂S caused</td>
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Experiment No. 46.—Flocculation of colloidal sulfur.

Solutions of the substances in Experiment No. 45 were added to colloidal sulfur, and then these solutions containing the colloid were treated with H₂S and with the H₂S and CO₂ mixture exactly as the material was treated in Experiment No. 45.

The results are here tabulated.

<table>
<thead>
<tr>
<th></th>
<th>Colloidal S in water</th>
<th>N/300 NaCl</th>
<th>N/300 KCl</th>
<th>N/300 CaCl₂</th>
<th>with 1 gram CaCO₃</th>
<th>N/300 AlCl₃</th>
<th>with 1 gram Al₂O₃</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Suspended colloid</td>
<td>S</td>
<td>More S</td>
<td>More S</td>
<td>S</td>
<td>S</td>
<td>S</td>
</tr>
<tr>
<td></td>
<td></td>
<td>remained</td>
<td>suspended</td>
<td>suspended</td>
<td>soon</td>
<td>soon</td>
<td>soon</td>
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<td></td>
<td></td>
<td>than in</td>
<td>than in</td>
<td>than in</td>
<td>flocculated</td>
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<td>flocculated</td>
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<td></td>
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<td>1 A</td>
<td>1 A</td>
<td>1 A</td>
<td></td>
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<tr>
<td>B</td>
<td>Liquid saturated with H₂S</td>
<td>S</td>
<td>S</td>
<td>S</td>
<td>S</td>
<td>S</td>
<td>S</td>
</tr>
<tr>
<td></td>
<td>Suspended colloid</td>
<td>flocculated</td>
<td>flocculated</td>
<td>flocculated</td>
<td>remained</td>
<td>remained</td>
<td>remained</td>
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<td></td>
<td></td>
<td>rapidly</td>
<td>slowly</td>
<td>slowly</td>
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</tr>
<tr>
<td>C</td>
<td>Liquid saturated with H₂S and CO₂</td>
<td>Much S</td>
<td>Much S</td>
<td>Much S</td>
<td>S</td>
<td>S</td>
<td>S</td>
</tr>
<tr>
<td></td>
<td>Suspended colloid</td>
<td>remained</td>
<td>remained</td>
<td>remained</td>
<td>dispersed</td>
<td>dispersed</td>
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</table>
Experiment No. 47.—Dispersing and flocculating effects of $H_2S$, $SO_2$, and $CO_2$ on colloidal copper sulfide and on colloidal sulfur, in water and in the presence of $N/1400 K_2CO_3$ and $KCl$.

A colloidal copper sulfide was prepared as described in Experiment No. 45. A colloidal sulfur solution was prepared by the action of $H_2S$ on $SO_2$, the sulfur being suspended in water.

Portions of each colloidal solution were saturated with $H_2S$, $SO_2$, and $CO_2$, and then portions to which $K_2CO_3$ and $KCl$ had been added so that the K ion was $N/1400$, were saturated with the gases.

The results are here tabulated.

### SULFUR

<table>
<thead>
<tr>
<th></th>
<th>Water suspension of the Colloid</th>
<th>$N/1400 K_2CO_3$ and $KCl$ suspension of the Colloid</th>
</tr>
</thead>
<tbody>
<tr>
<td>$H_2S$</td>
<td>$H_2S$ caused the sulfur to flocculate rapidly.</td>
<td>$H_2S$ caused the sulfur to flocculate slowly.</td>
</tr>
<tr>
<td>$SO_2$</td>
<td>$SO_2$ did not flocculate the sulfur rapidly.</td>
<td>$SO_2$ did not flocculate the sulfur rapidly.</td>
</tr>
<tr>
<td>$CO_2$</td>
<td>$CO_2$ practically prevented the flocculation of the sulfur.</td>
<td>$CO_2$ prevented the flocculation of the sulfur.</td>
</tr>
</tbody>
</table>

### COPPER SULFIDE

<table>
<thead>
<tr>
<th></th>
<th>Water suspension of the Colloid</th>
<th>$N/1400 K_2CO_3$ and $KCl$ suspension of the Colloid</th>
</tr>
</thead>
<tbody>
<tr>
<td>$H_2S$</td>
<td>$H_2S$ kept the copper sulfide dispersed for a long time.</td>
<td>$H_2S$ practically prevented any flocculation of the copper sulfide.</td>
</tr>
<tr>
<td>$SO_2$</td>
<td>$SO_2$ caused the copper sulfide to flocculate.</td>
<td>$SO_2$ caused the copper sulfide to flocculate.</td>
</tr>
<tr>
<td>$CO_2$</td>
<td>$CO_2$ caused a very clean cut flocculation of the copper sulfide.</td>
<td>$CO_2$ caused a very clean cut flocculation of the colloid.</td>
</tr>
</tbody>
</table>
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This work was undertaken in the hope of giving more information to the geologist, and in some degree it has been successful.

The amount of success which has attended the efforts of the writer is in no small degree due to the interested suggestion, assistance and oversight given by
Professors C. F. Tolman, Jr., A. F. Rogers and S. W. Young. Professor Tolman indicated geological interpretations of the work, Professor Rogers contributed most willingly many of his finest mineral specimens for experimentation and Professor Young gave direct supervision of the research. To these gentlemen, and particularly to Professor Young, the author acknowledges his deep indebtedness.

JOHN DUSTIN CLARK.

Leland Stanford Junior University,
May 18th, 1914.