THE CYANIDE PROCESS

OF

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THE CYANIDE PROCESS
OF
GOLD EXTRACTION:
A TEXT-BOOK FOR THE USE OF MINING STUDENTS,
METALLURGISTS, AND CYANIDE OPERATORS.

BY

JAMES PARK,
PROFESSOR OF MINING AND DIRECTOR OF OTAGO UNIVERSITY SCHOOL OF MINES;
FELLOW OF THE GEOLOGICAL SOCIETY OF LONDON; MEMBER OF THE AMERICAN
INSTITUTE OF MINING ENGINEERS; MEMBER OF THE INSTITUTE OF
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With Frontispiece, Plates and Illustrations.

[Authorized Text-book, Australian Schools of Mines]

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PREFACE TO THE FOURTH ENGLISH EDITION.

The exhaustion of the last edition has enabled the author to revise the text and add much new matter relating to recent improvements in cyanide practice in different parts of the globe.

The principles underlying the treatment of silicious silver ore carrying more or less gold is dealt with more fully than in former editions. The weak point in the present treatment of these ores is the low extraction of the silver. At the current price of silver this is perhaps not of much moment with ores in which the chief value lies in the gold contents; but in the case of low-grade silver ores carrying from 1 to 3 dwt. of gold, and from 10 to 20 oz. of silver, per ton, the subject is one of increasing concern to the mine owner. The difference between an extraction of 50 per cent. and 85 per cent. of the silver may represent the sole margin of profit in a low-grade venture.

The dissolution of gold in a clean pyritic ore seldom presents serious chemical difficulties. From such an ore the bulk of the gold is extracted by plate-amalgamation, leaving only the more finely-divided gold to be extracted by cyanide from the tailings. The treatment of argentiferous gold ores, such as those of New Zealand, South Dakota, and Mexico, is a much more difficult matter, involving not only the direct cyaniding of all the crushed material, but the solving of chemical problems not met with in the treatment of ordinary gold ores.

The dissolution of silver sulphides liberates a corresponding
amount of sulphur, which at once forms alkaline sulphides and
sulphocyanides, both causing loss of cyanide and free oxygen,
the former being especially injurious in that it retards the
progress of dissolution and prevents a satisfactory extraction
of the silver.

The use of a desulphurizing agent will be found to give
increased extractions of both the gold and silver, besides effect-
ing a saving of cyanide, and a lessening of the time required for
dissolution. It should, however, be remembered that to obtain
the highest extractions from the majority of dry silver ores fine-
grinding is a necessity; and fine-grinding does not necessarily
mean sliming. It is hopeless to expect a desulphurizer to increase
the extraction when the values are enveloped in grains of silica.
The economic limit of fine-grinding must be determined for each
ore by actual experiment.

For the successful treatment of pyritic concentrates fine-
grinding is a necessity, and here also the use of a desulphurizer
will be beneficial, more especially when followed by thorough
aeration, or the addition of an oxidizing agent.

Fine-grinding by tube-mills is now in successful operation in
Western Australia, New South Wales, New Zealand, and Mexico,
and in the past two years much additional experience has been
gained in the successful running of these machines. The re-
spective merits of grinding pans and tube-mills is still an open
question, and is probably incapable of solution. Each machine
possesses certain distinctive features which will enable it to assert
a superiority in the grinding of an ore suited to its peculiar
capacity. Thus on some Kalgoorlie ores the tube-mill may show
the higher efficiency, while on Rand ore the pan may possess a
decided superiority. This also is a matter for actual experiment.

One of the most notable advances in the metallurgy of gold
in recent years is the introduction of the Rose process for the
refining of gold and impure bullion. Its use should reduce the
cost and labour of refining cyanide bullion.
PREFACE.

Filter-pressing is still in successful operation in Kalgoorlie and New Zealand, but is slow and laborious, and from its intermittent action necessarily costly. Vacuum filter processes are again on their trial, and of these the Moore process is said to be the most promising. What is much needed for the successful treatment of low-grade slimes is a filtering process capable of continuous operation.

JAMES PARK.

Otago University,  
Dunedin, N.Z., 1906.
PREFACE TO THIRD EDITION.

The favourable reception accorded to earlier editions has encouraged me to revise the old matter, and at the same time add much new material, which, for the most part, relates to lead-smelting of gold-slimes, the treatment of sulpho-telluride ores, and filter-press practice. In Western Australia, the adoption of filter-pressing was mainly determined by a combination of peculiar local conditions, namely, the natural tendency of the gold ores and matrix to form slimes, the scarcity of fresh water, and the saline character of the only water available for milling purposes. The density of the brackish waters rendered the settlement of the finer material in the decantation process so slow, laborious, and imperfect as to make the use of filter-presses almost imperative for quick and effective treatment. The slimes, it should be noted, formed the first or primary product of the mills, and were of high value. In South Africa, where the question of filter-pressing versus decantation has been engaging attention for some time, the slimes are in all cases a secondary product and of low-grade value.

Filter-pressing has been adopted with much success by the Waihi Company, in New Zealand, for the treatment of ordinary slimes, but, on the other hand, it should be mentioned that several neighbouring companies are satisfied with the decantation process for the treatment of similar material. It is quite clear that the relative merits of the two processes must, in every case, be determined by exhaustive trials extending over a period of, say, four to
six months, so as to eliminate the element of fortuitous chance in favour of either, and thereby enable a reliable estimate of costs to be prepared. With South African and New Zealand conditions, the solution of the problem is obviously one of cost. In Western Australia, the question of cost is subordinate to that of expediency.

The introduction of lead-smelting of gold-slimes marks a notable advance in cyanide practice, and if the claims of Mr. Tavener, the author of the process, are verified by more comprehensive trials—as indeed seems most probable—lead-smelting will in a short time displace the old smelting and acid-treatment processes, at any rate in all the larger permanent cyanide plants.

There is still much room for special chemical investigation in several directions, and in this respect the researches of a number of accomplished metallurgical chemists in South Africa have added much valuable material to the literature of the cyanide process. The successful regeneration of foul cyanide solutions is still unsolved, and is at present the subject of investigation by several American chemists. The results of their work will be awaited with much interest.

I have to acknowledge my indebtedness to Mr E. G. Banks, M.I.M.M., and Mr Frank B. Allen, M.A., B.Sc., for special contributions on Waihi slime and filter-press practice and Kalgoorlie sulpho-telluride practice respectively; and to the proprietors of the Mining Journal, Mining and Engineering Journal, and Australian Mining Standard for permission to make extracts from articles which appeared in their columns at different times. In these and all other cases, due acknowledgment and reference are made in the text.

JAMES PARK.

University, Dunedin, N.Z.,
September 1908.
PREFACE TO THE FIRST ENGLISH EDITION.

The favourable reception accorded to the Third Edition of this work, published in New Zealand, has enabled the author to again revise and enlarge the text, with a view of placing before his readers, earlier than was anticipated, the latest information available on this progressive branch of metallurgy. The general plan of the original work, which was intended for the use of mining students, metallurgists, and cyanide operators, has been retained in the present edition, which is the first published in England, and into which only such changes have been introduced as the author considered needful. Of late years the application of scientific investigations and methods to the treatment of ores has rendered metallurgy more and more dependent on chemical knowledge, and in no department is this more obvious than in the Cyanide Process of Gold Extraction, which often presents the most perplexing problems, due, in many cases, to the simplest causes.

In the present edition several new illustrations and tables have been added, while the information relating to the treatment of slimes, the analyses of solutions, and cyanide poisoning, has been greatly extended; while, by the adoption of slightly smaller type, and closer setting, the actual number of pages has been reduced, although the text has been considerably enlarged.

Since the revision of these pages, wet-crushing and cyanide treatment have largely superseded dry-crushing and direct
cyaniding in New Zealand, and in every case their installation has been attended with complete success, notwithstanding the large proportion of slimes at some of the mines. The ores are mostly chalcedonic in character, and contain both coarse and fine gold associated with silver sulphide. The silver occurs in constantly varying proportions, requiring varying strengths of solution to obtain adequate extractions, and unremitting care on the part of the metallurgist in charge.

The results obtained in the treatment of these comparatively complex ores are of world-wide interest and have been embodied by the author in a separate appendix.

AUCKLAND. January 1900.

JAMES PARK.
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THE CYANIDE PROCESS FOR THE EXTRACTION OF GOLD AND SILVER.

CHAPTER I.

THE McARTHUR-FORREST PROCESS.

It has long been known that gold and silver are soluble in solutions of alkaline cyanides, but it is only since 1890 that this knowledge has been applied on a commercial scale to the extraction of the precious metals from their ores.

The discovery of the fact that the dilute solution of potassium cyanide is a solvent for natural gold ranks among the most remarkable discoveries of the nineteenth century in metallurgical science; and the widespread and successful application of the fact must mark an epoch in the history of gold extraction for all time.

SCOPE OF THE PROCESS.

The cyanide process can be applied with success to the treatment of free-milling ores in which the gold occurs in fine particles, or of tailings and concentrates resulting from wet-crushing and copper-plate amalgamation, or dry-crushing and pan-amalgamation. It can also be used for the treatment of many so-called refractory ores, especially those in which the gold occurs in such a finely divided form that even amalgamation in pans fails to
recover a satisfactory percentage of the values; or of ores in which the gold is coated with a film of metallic oxide or sulphide, rendering it non-amalgamable, and ores in which the gold is associated with, or entangled in, a highly pyritic matrix.

All the common ores of silver are more or less soluble in dilute solutions of cyanide. Those most readily soluble are the chloride (AgCl) and the sub-sulphide (Ag₃S), and these are fortunately the most abundant; but the rate of dissolution of silver and its ores is much slower than that of gold, and is accompanied by a higher consumption of cyanide.

LIMITATIONS OF THE PROCESS.

The cyanide process cannot be applied with success to the treatment of ores in which the gold occurs even in a fairly coarse condition. When an ore contains a proportion of both fine and coarse gold the cyanide process may be used to extract the fine gold, but a supplementary treatment will have to be used to recover the coarse gold, since the slowness of the dissolution would take too long for a commercial basis of working.

With free-milling ores of the latter class the recovery of the coarse gold is generally effected by copper-plate amalgamation; and, in the case of wet-crushing, this treatment precedes the cyanide leaching, while, in the case of dry-crushing, it follows it.

The experience gained during the use of the cyanide process has shown that solutions of potassium cyanide, even when very dilute, act most energetically on all the sulphide, oxide, and carbonate ores of copper, and also on the sulphides of antimony, bismuth, and cobalt; hence, when any one of these is present, even in small proportion, the treatment of the ore becomes difficult, and sometimes impossible, on account of the great consumption of cyanide. In practice it is found that an unduly large consumption of cyanide is generally accompanied by a low rate of extraction of the gold and silver contained in the ore or tailings.

From the foregoing it is obvious that the process will be most successful in the treatment of ores in which the gold occurs in a very fine state, and in which the quantity of base minerals or metallic salts, destructive to cyanide, is small.

Further, the author ascertained, as the result of many comparative working trials in the N.Z. Government Metallurgical Works at the Thames goldfield, that argentiferous gold ores which were amenable to treatment by the Washoe pan-amalgamation process, in most cases yielded better results by cyanide treatment, even when they contained a small percentage of zinc and lead.
An intelligent knowledge of first principles, backed by experience of working details and working requirements, has led to many ingenious adaptations, the tendency of which has been to widen the scope of the original cyanide process. A notable case in point is the successful treatment of the rich sulpho-telluride ores of Cripple Creek and Kalgoorlie, which affords satisfactory evidence of the progressive trend of this important branch of metallurgy.
CHAPTER II.

THE CHEMISTRY OF THE PROCESS.

When gold is acted on by an aqueous solution of potassium cyanide, a solution is obtained which, when evaporated, yields octahedral crystals having the composition of the auro-potassic cyanide (AuKCy₂), which is a double cyanide of gold and potassium.

The exact reaction which takes place when gold is dissolved by potassium cyanide is not yet well understood, being still a subject of much doubt and uncertainty. According to some authorities, the gold is oxidized before it is dissolved; while others maintain that the cyanide is first oxidized and then acts on the gold.

The reaction suggested by Elsner in 1842 is the one now most generally accepted by chemists. It is represented by the following equation:

\[ 4\text{Au} + 8\text{KCy} + \text{O}_2 + 2\text{H}_2\text{O} = 4\text{AuKCy}_2 + 4\text{KHO}. \]

According to the above equation, an ounce of oxygen is required for every pound of potassium cyanide employed for the dissolution of the gold. This view has received substantial support from the author's experiments in 1891, and from those of Škey* in 1892, and has since been proved experimentally correct by Maclaurin † in his classical paper on the subject.

The valuable researches of Škey and Maclaurin have shown that the rate of dissolution of pure gold, under theoretical conditions, reaches a maximum in passing from dilute to concentrated solutions of potassium cyanide. By actual experiment, it was proved that the maximum rate was reached with a 0·25 per cent. solution of cyanide. On a working scale the maximum varies with the character of the mineral constituents of the ore, and can easily be determined by a series of laboratory experiments.

A weak solution is always more active than a strong one, and

Maclaurin considers that this remarkable fact may be accounted for by supposing that the rate of dissolution of gold is partly dependent on the number of cyanide molecules in a unit volume; and partly on the number of oxygen molecules in the same volume. One of the most important results of his exhaustive experiments was the demonstration of the fact that the solubility of oxygen in cyanide solutions decreases with concentration of the solution.

Weak aqueous solutions of cyanide exert a very marked action on gold and silver when these metals are associated with ores of copper and antimony. This circumstance becomes very prominent during the treatment of cupriferous ores on a large scale.

The cyanides of the alkaline metals are soluble in water, while those of the heavy metals, with the exception of gold and mercury, are insoluble. The insoluble salts are, however, soluble in excess of potassium cyanide.

The use of an oxidizing agent that will readily part with a portion of its oxygen in a cyanide solution forms the essential feature of several new patent cyanide processes. The employment of such an agent serves to accelerate the dissolution of the gold contained in the ore. The artificial aeration of cyanide solutions is undertaken to supply atmospheric oxygen with the same object.

Consumption of Cyanide.—According to Elsner’s equation, about 4.5 lbs. of cyanide should dissolve 100 ounces of gold, but in practice it is found that it takes nearly forty times that quantity. The causes which operate in the practice of the process to effect so large a consumption of cyanide, over that required by Elsner’s simple equation, are at present not fully investigated.

To dissolve 100 oz. of silver would require 7.5 lbs. of cyanide, according to the equation:

$$4\text{Ag} + 8\text{KCy} + \text{O}_2 + 2\text{H}_2\text{O} = 4(\text{AgKCy}_2) + 4\text{KHO}.$$  

For the dissolution of 100 oz. of silver existing as the sub-sulphide (Ag$_2$S), 7.01 lbs. of cyanide would be required by the following equation:

$$\text{Ag}_2\text{S} + 4\text{KCy} = 2(\text{AgKCy}_2) + \text{K}_2\text{S}.$$  

The potassium sulphide resulting from the dissolution of silver sulphide also tends to cause a further loss of cyanide by precipitating gold and silver which will require an excess of free cyanide to redissolve it. It is the need for this excess of cyanide which necessitates the use of comparatively strong solutions in the treatment of argentiferous gold ores.
Potassium cyanide is, chemically, a most active organic compound, possessing the property of forming so large a number of complicated and unexpected combinations in the presence of mineral acids and base metals, that its reactions and behaviour with different classes of ore, and under varying conditions, can only be unravelled by much patient research, both in the laboratory and under working conditions.

During the treatment of ores by the cyanide process, the most puzzling difficulties are continually met with, requiring the constant care and attention of the metallurgist in charge.

Causes of Loss of Cyanide.—Some of the principal and more obvious causes of the enormous loss of cyanide which takes place in the working of the process are as follows:—
1. Loss by absorption in wooden vats or tanks.
2. Loss by decomposition by atmospheric carbon dioxide.
4. Loss by decomposition due to the presence of mineral acids and salts.
5. Loss due to presence of ores soluble in cyanide.
6. Loss when gold exists as amalgam.
7. Loss due to the presence of charcoal in kiln-dried ore.

Loss by Absorption in Vats.—This is especially noticeable in new plants. At the Witwatersrand Goldfields, the loss from this cause is said by Mr. C. Butters to amount to a pound of cyanide per ton of tailings treated. At the first monthly “clean up” in a new plant, the actual extraction is often twenty or more per cent. below the theoretical, but after a few months it generally rises to within three to six per cent. of the extraction as determined by assay.

With iron or steel vats there is no appreciable loss by absorption.

Loss due to Decomposition by Atmospheric Carbon Dioxide.—The carbonic acid gas of the atmosphere decomposes potassium cyanide with the formation of potassium carbonate, and the liberation of hydrocyanic (prussic) acid, thus:—

\[ 2KCy + CO_2 + H_2O = K_2CO_3 + 2HCy. \]

The prussic acid thus liberated would be neutralized by any caustic alkali present in the cyanide solution.

The loss due to CO_2 is considerably greater during agitation than with ordinary percolation. The artificial aeration of cyanide solutions also increases the loss of available cyanide due to this cause, except there is an excess of free alkali present to unite with the HCy.

Mechanical Loss in Residues, and by Dilution during Washing.—During washing there is an inability to extract the
whole of the cyanide from the residual tailings. The dilution of the cyanide solutions also occasions a loss of cyanide in washing. A large quantity of dilute cyanide solution is formed, a portion only of which can be utilized to make up fresh solutions.

Loss by Decomposition due to Mineral Acids and Salts.—The metallic minerals most commonly found associated with gold in quartz veins are iron pyrites, copper pyrites, zinc blende, galena, and antimonite. By far the most common and abundant of these is iron pyrites.

It has been shown by Skey and others that clean fresh iron pyrites is not acted on by working solutions of cyanide. The decomposition products of this mineral, however, act most destructively on cyanide, and the obvious conclusion to be drawn from this is that the treatment of pyritic tailings, or concentrates, by cyanide should be undertaken with as little delay as possible; more especially when the pyrites occurs in the marcasite form, which is much more prone to oxidation than the cubical or isometric form.

In the shallow parts of mines, the pyrites is generally oxidized to ferric oxide, which does not act chemically on cyanide, but causes a mechanical loss through the formation, both in wet and dry crushing, of extremely fine slimes, which are very absorbent and retentive of cyanide solutions.

Iron pyrites \((\text{FeS}_2)\) is decomposed by atmospheric oxygen in the presence of moisture into the soluble ferrous sulphate and free sulphuric acid, according to the following equation:—

\[
\text{FeS}_2 + \text{H}_2\text{O} + 7\text{O} = \text{FeSO}_4 + \text{H}_2\text{SO}_4.
\]

In the kiln-drying of ores to be dry-crushed, the heat to which the ore, often in large pieces, is subjected, is not very uniform, especially in large kilns. With pyritic ores the sulphides are decomposed at certain temperatures into oxides and soluble sulphates; and at higher temperatures the latter salts are converted into oxides.

The steam generated from the moisture in the fuel and in the ore itself assists these reactions. In the kiln, where the temperature is high, reducing gases are evolved, and these may impede the oxidation of the sulphide, causing the formation of lower sulphides and basic sulphates, which are insoluble in water, but react on cyanide.

For these reasons, and also for economy, ores intended for dry treatment by cyanide should never be kiln-dried. In the case of developed mines, with an assured and steady output, there is nothing to justify the practice, but much to condemn it.

As we have seen, the atmospheric oxidation of pyrites results in
the production of free sulphuric acid and ferrous sulphate. This
ferrous sulphate may in turn be decomposed, by the action of the air,
into insoluble basic sulphates. Thus, partly oxidized pyritic ores or
tailings may contain free sulphuric acid, soluble ferrous sulphate,
insoluble basic sulphates, and probably also traces of other basic
salts of complex and variable composition, all of which react upon
solutions of potassium cyanide, thereby causing a loss of cyanide.

The reactions which are most likely to take place in acid ores
or tailings in the presence of cyanide are:—

(a.) The liberation of hydrocyanic acid.
(b.) The formation of ferro- and ferri cyanides.

The free acids in the ore react on the cyanide as shown by the
equation:—

\[ 2KCy + H_2SO_4 = 2HCy + K_2SO_4. \]

Feldtmann considers it possible for the hydrocyanic acid thus
liberated to diffuse itself through the ore and dissolve appreciable
quantities of gold.* For this reason he strongly condemns the
practice of washing acid tailings in the leaching vats, as these must
always contain a residual portion of cyanide from which prussic
acid would be liberated. Any gold dissolved by this gas would
be carried away in the water or alkaline wash; and to avoid this
possible source of loss, which he thinks may account for the
mysterious discrepancy sometimes found between the assay and
the actual extraction, he recommends the system of washing in one
vat and leaching in another. On the other hand, Skey, when dis-
cussing this subject with the author, stated that hydrocyanic acid
was not a solvent for gold. It is obvious, however, that this
liberated HCy in the presence of any residual alkali in the vat,
would form an alkaline cyanide capable of dissolving and removing
gold, and Feldtmann probably had this combination in his mind
at the time of writing.

Of the iron salts, the one of most common occurrence in pyritic
ores or tailings is the soluble ferrous sulphate (FeSO_4), which
reacts with potassium cyanide to form potassium ferro-cyanide
and sulphate, thus:—

\[ FeSO_4 + 6KCy = K_4FeCy_6 + K_2SO_4. \]

The potassium ferro-cyanide thus formed is, in its turn, reacted
on by any excess of ferrous sulphate still present with the produc-
tion of Prussian blue according to the equation:—

\[ 3K_4FeCy_6 + 6FeSO_4 + 3O = Fe_2O_3 + 6K_2SO_4 + Fe_4(FeCy_6)_3. \]

* Feldtmann, *Notes on Gold Extraction*, p. 5.
A blue colour in the solution, on the surface of the tailings, or in the seams of the staves of the vats, indicates a large consumption and loss of cyanide due to imperfect washing and neutralization of the acidity in the preliminary treatment.

A white scum or precipitate is sometimes seen on the surface of the solutions when they are coming off acid. This precipitate turns into Prussian blue by exposure to the air and light.

The normal ferric sulphate $\text{Fe}_2(\text{SO}_4)_3$ is insoluble in water, and cannot be removed by ordinary water-washing. It reacts with potassium cyanide, causing a loss of cyanogen due to the liberation of prussic acid and the formation of the ferric hydrate, as shown by the two following equations:

\[
\text{Fe}_2(\text{SO}_4)_3 + 6\text{KCy} = \text{Fe}_2\text{Cy}_6 + 3\text{K}_2\text{SO}_4
\]

and

\[
\text{Fe}_2\text{Cy}_6 + 6\text{H}_2\text{O} = \text{Fe}_2(\text{HO})_6 + 6\text{HCy}.
\]

It is probable that in most partially oxidized pyritic ores and tailings the ferrous and ferric sulphates exist together, the former in large excess. In this case the decomposition of the cyanide would result in the production of ferrous cyanide and potassium sulphate, thus:

\[
12\text{KCy} + 3\text{FeSO}_4 + \text{Fe}_2(\text{SO}_4)_3 = \text{Fe}_3(\text{FeCy}_6)_2 + 6\text{K}_2\text{SO}_4.
\]

In the case of earthy pyritic ores, the weathering or oxidation of the metallic sulphides would result in the production of sulphates of magnesia, lime, or alumina. The action of these sulphates is not very clear, but they most likely react on cyanide with the liberation of prussic acid, accompanied by the formation of the hydrated oxide of the metals in question, which would be precipitated as an insoluble incrustation in the solution pipes.

The above reactions clearly emphasize the necessity of a most careful preliminary alkaline treatment of pyritic material, in order to avoid undue loss of cyanide, and ensure satisfactory results.

All the iron salts and earthy sulphates can be rendered innocuous by the application of an alkali before treatment with the cyanide. By this means all the soluble iron salts are precipitated as ferrous hydrate, which rapidly oxidizes to ferric hydrate; while the basic ones soon oxidize in the presence of the alkali. It is important to remember that the alkali should be applied before, and not with the cyanide solutions, as these iron salts will destroy the cyanide as much in a strongly alkaline as in a nearly neutral solution. When the tailings contain free acid only, alkali and cyanide should be applied together.
Loss due to Presence of Ores Soluble in Cyanide.—
The sulphide, oxide, and carbonate ores of copper, and the sul-
phides of antimony and bismuth, are acted on by potassium
cyanide both in weak and strong solutions, and thereby cause a
loss of cyanide in proportion to their abundance in the ore. In
the treatment of an ore containing as little as 0·25 per cent. of
copper the consumption of cyanide will be doubled.

It is during the treatment of cupriferous ores that the selective
action of weak cyanide solutions becomes most apparent. An ore
may contain sufficient copper to decompose a 1 per cent. solution
of cyanide and give a low extraction of gold, whereas a 0·35 per
cent. solution would dissolve proportionately less copper, and give
a fairly satisfactory extraction of the gold. But the same results
would be obtained even in the absence of copper, for it has already
been shown that the rate of dissolution of gold reaches a maxi-
num in passing from dilute to strong solutions. Hence a 0·35
per cent. solution should extract more gold than a 1 per cent.
solution, the weaker solution being nearer the strength at which
a maximum rate of dissolution occurs, and which has been proved
experimentally to be a 0·25 per cent. on pure gold.

The cyanide treatment of ores, and zinc-precipitation of the
gold, have shown the existence of copper in ores in which no trace
of that metal could be detected, even by the most rigid chemical
examination on large samples. An instance of this came under
the notice of the author, at the Crown mines at Karangahake.
The ore being treated there consisted of almost pure white quartz,
free from all metallic impurities; nevertheless, a portion of the
zinc in the precipitation boxes was often coated with a film of
bright metallic copper. The copper could not be derived from an
outside source, or from any of the mechanical fittings in the mill
or cyanide plant, and Mr. James Napier, the metallurgist and
chemist in charge, was of the opinion that it existed in the ore in
an infinitesimally small quantity, and only became manifest on
the zinc turnings after the treatment of hundreds of tons of ore.

Copper pyrites is oxidized to the soluble sulphate at low tem-
peratures, and this salt requires a greater heat to decompose it
than iron pyrites. It is, therefore, probable that a portion, at least,
of this mineral present in an ore, being dried in kilns, preparatory
to dry-crushing and direct cyanide treatment, would be sulphatized,
and thereby cause an appreciable loss of cyanide in a manner
similar to that caused by the decomposition products of iron pyrites.

Malachite and azurite, the green and blue carbonate of copper,
are both readily soluble in dilute solutions of cyanide, with the
production of copper-potassic cyanide and liberation of prussic
acid.

Antimonite, the grey sesqui-sulphide of antimony, is also readily
acted on by weak cyanide solutions. It is frequently met with in the gold-bearing ores of the Thames and Reefton goldfields. The presence of a small percentage of antimonite in the large accumulation of tailings at Boatman's Creek, near Reefton, is said to have caused all attempts to treat them to end in failure, chiefly owing to the large consumption of cyanide and the low rate of extraction.

**Loss of Cyanide when Gold exists as Amalgam.**—It is well known to most millmen that a considerable portion of the gold in tailings, resulting from copper-plate amalgamation or pan-amalgamation, exists in the form of amalgam. When such tailings have to be treated the cyanide has to dissolve the mercury as well as the gold, thus causing a larger consumption of the solvent than would be necessary if the gold existed in a free state.

According to Gmelin, mercury is not dissolved or acted on by potassium cyanide; but the practical working of the cyanide process has shown that his conclusion is contrary to actual experience.

Besides using up available cyanide, it combines with oxygen during its dissolution, thereby reducing the supply of free oxygen in the solution. On the other hand, mercury has a stronger affinity for sulphur than for cyanogen, hence when dissolved it will act as a desulphurizer.

The loss of mercury in mills using plate-amalgamation varies with the character of the ore, being less with clean than with acid ores. With clean ores the loss may vary from 0.01 lb. to 0.04 lb. per ton of ore crushed, and with acid ores may reach 0.08 lb.

In the treatment of argentiferous gold ores by pan-amalgamation the loss often amounts to 0.4 lb. of mercury per ton of ore treated.

The mercury generally occurs in the tailings in the form of amalgam in a very fine state of subdivision, and is dissolved by the cyanide, together with the associated gold and silver. It is precipitated with the bullion in the zinc precipitation boxes. When the zinc slimes are oxidized the greater portion of the mercury is volatilized.

**Loss of Cyanide due to Iron.**—Cast-iron or steel, mostly in the form of fine particles, is present in all pulverized ores. It is introduced at all stages of the crushing and pulverizing by the wear and tear of the working faces of the crushing or grinding machines.

The amount of iron so introduced in the majority of cases greatly exceeds the amount of gold and silver in the ore, and being soluble in dilute aqueous solutions of cyanide, must cause a loss of available cyanogen.
The Cyanide Process.

According to the equation representing the dissolution of iron in potassium cyanide, 1 lb. of iron requires theoretically 7 lbs. of KCy to effect its dissolution, in practice probably not less than 100 lbs.; whereas the same weight of gold requires theoretically only 0.62 lb., but in practice from 25 to 40 lbs., according to the physical condition of the gold and the character of its mineral associates.

If the iron existed in as finely divided a state as the gold and were equally soluble, the iron would be extracted in a much greater proportion than the gold. The consumption of cyanide would also be relatively greater. But, fortunately for the metallurgist, the gold exists in a more finely divided state than the bulk of the iron, and is five times more soluble, two factors which favour the dissolution of the gold in preference to the iron.

That a loss of cyanide due to the dissolution of iron in the pulverized ore does take place is believed by many metallurgists, but whether the amount is small or great has not yet been determined.

Besides using up available cyanide, the dissolution of the iron will remove oxygen from the solution, and thereby retard the dissolution of the gold.

When crushing with cyanide solution in the mortars, as practised in New Zealand and South Dakota, the molecule of cyanide is only momentarily in contact with flat surfaces of iron. The loss of cyanide due to this cause is hardly appreciable.

Loss of Cyanide due to the presence of Charcoal in Ore.—It has long been known to chemists that charcoal possesses the property of decomposing auro-potassic cyanide solutions, and charcoal precipitation has been successfully substituted for zinc precipitation in Victoria. The author has found that a very large bulk of charcoal and prolonged contact with the solutions are required to effect complete precipitation of the bullion. Charcoal also, after some hours' contact, acts on potassium cyanide solutions, thus causing a loss of cyanogen. For these reasons it seems improbable that charcoal will ever supersede zinc as a precipitant of gold from cyanide solutions.

It is obvious from the above that the presence of charcoal in ores which have been kiln-dried preparatory to cyanide treatment may cause a loss of both cyanide and gold during the leaching.

At the Kapai-Vermont Cyanide Works, at Kuaotunu, New Zealand, the metallurgist, Mr. J. A. Walker, stated that it paid to employ labour to pick the lumps of charcoal and partially-carbonized wood out of the ore before it passes to the ball-mill to be pulverized. This is a difficulty which could easily be avoided by drying the ore in a revolving or other mechanical
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dryer, in which the heated air and gases only come in contact with the ore.

The Action of Cyanide on Metallic Sulphides.—This was the subject of an investigation by the late Mr. William Skey, analyst to the New Zealand Government. The results were communicated to the Government early in 1895, and published in an interesting paper in the Annual Report of the Mines Department for 1895.

Referring to the oft-repeated statement that very dilute solutions of potassium cyanide have no effect on copper sulphides, it will be seen that Mr. Skey's results entirely disprove this. They are as follows:

1. Chalcopyrites, crushed, then thoroughly well washed, was afterwards kept in a cyanide solution of 0.03 per cent. for one hour. The filtered solution contained a very perceptible quantity of copper as cyanide, also traces of sulphur and oxidized compounds of sulphur.

2. Copper-glance, crushed and washed, was then subjected to the cyanide solution of 0.03 per cent. for one hour. The filtered solution gave the same reaction as in the case of chalcopyrites.

3. Covellite (sub-sulphide of copper).—As previously shown, this sulphide is very easily attacked by weak solutions of cyanide, sulpho-cyanide of copper resulting.* The 0.03 per cent. solution dissolves the mineral as a sulpho-cyanide and cyanide of copper.

The carbonates and silicates of copper, as they naturally occur—even, that is, the dense form—are also decomposed by cyanide of this strength.

These results take in all the compounds of copper that are of general occurrence at the gold-mines, and show that potassic cyanide, however weak, will generally decompose copper ores, when in contact with them.

Mr. Skey's results upon certain other ores that are frequently associated with gold in reefs are as follows:

Stibnite (sulphide of antimony).—This ore is generally supposed to be unaffected by potassic cyanide, but was found to be very easily and largely affected by this salt. The sulphur of this mineral, like that present in certain copper sulphides, shows a great tendency to combine with potassium to form that most objectionable salt—sulphide of potassium—the rest of the sulphur combining to form sulpho-cyanogen.

Galena (sulphide of lead).—This mineral is slowly attacked by cyanides, but all its sulphur combines with cyanogen to form the

harmless compound, sulphi-cyanogen, while the lead comes into solution combined with that radical.

_Zinc blende_ (sulphide of zinc).—Hardly affected by cyanide of any strength; it is commonly held to be perfectly neutral thereto; but a strip of silver, buried in the crushed ore, was blackened in one hour, showing that a slight degree of decomposition had taken place.

_Iron-pyrites_ (bi-sulphide of iron).—Almost, if not quite, unaffected by cyanide solutions of any strength.

_Sulphur_ (free).—The Government Geologist states that he met with sulphur in a free state in tailings at Boatman's Creek, in the Reefton district, as a product of the decomposition of stibnite.

It is asserted by Wiggers* to be insoluble in potassic cyanide solutions, but it is very easily soluble therein if only the air, with which the latter is generally charged, be driven off, say, by boiling water.† As in the cases of the solution of copper, this means a loss of solvent power for gold and silver.

Cold solutions were used throughout Mr. Skey's experiments.

The formation of alkaline sulphides during the cyanide process, as applied to ores containing stibnite and copper-sulphides, is, undoubtedly, the chief cause of the loss of gold that so often occurs in that process.

With oxidized ores, as malachite or dioptase, a loss of available cyanide will necessarily occur; but with these sulphides, in addition to loss of cyanide, there will be a loss of gold, and a still greater loss of silver, in proportion to the quantities present. This loss is brought about by the sulphur—that is, the alkaline sulphide—sulphurizing these metals to form sulphides with them,‡ the sulphide film so formed upon the metal preventing, or greatly retarding, the proper action of the cyanide solution.

That gold does combine, and very readily, with the sulphur of both the alkaline sulphide and of hydrogen sulphide, Skey has already shown.§

It is to precipitate the sulphur that gets into the cyanide in the cyanide process that Mr. McArthur has proposed to use, or does use (as per patent), a soluble lead salt dissolved in the cyanide.

The problem for the chemist at the cyanide works is to find a practical method, whereby all the sulphur of antimonial and cuprous sulphides can be made to combine with the cyanogen, rather than with the potassium of the cyanide.

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The following results obtained by Skey show how extremely objectionable alkaline sulphides are, when present in the cyanide solution.

A rather strong solution of the cyanide, containing a small proportion of sulphur, was placed over a strip of gold coupled with a piece of copper-glance (sulphide of copper), but no solution of gold was perceived; however, on substituting chalcopyrites for the copper-glance, the gold was rapidly removed.

This experiment shows that the gold was sulphurized at the outset by the alkaline sulphide present in the cyanide, and that it required connecting with a substance of a strongly negative kind in order to effect the decomposition of the auriferous sulphide so formed.

Further experiments of a different kind showed that while pure 1 per cent. cyanide solution dissolved a given weight of gold in ten minutes, a solution of the salt of the same strength, but containing \( \frac{1}{100000} \) part of sulphur (as a sulphide),* required two hours to dissolve the same weight of gold. The speeds were as 12 to 1 in favour of the pure cyanide.

The following results show to what extent even a gentle sulphurizing, or flouring of the gold, interferes with its solution:

- Gold sulphurized 60 seconds in \( K_2S \), dissolved in cyanide in 62 minutes.
- Gold sulphurized 54 seconds in \( K_2S \), dissolved in cyanide in 50 minutes.
- Gold sulphurized 1 second in \( K_2S \), dissolved in cyanide in 36 minutes.
- Gold, clean, dissolved in cyanide in 12 minutes.

The gold was well washed from adherent potassic sulphide before being placed in the cyanide. Making clean gold the unit, the approximate times of dissolution are 1:3:4:5.

**Loss of Cyanide due to Antimonite or Stibnite.**

Antimony sulphide is often present in gold ores, but is not directly soluble in KCy. It is very soluble in caustic alkali, and then decomposes cyanide. For this reason it is advisable to avoid excess of caustic alkali in the treatment of ores containing antimony. The antimony is deposited on the zinc, and is partially volatilized during the roasting of the gold-zinc slimes.

At the Bidi and Bau mines in Upper Sarawak in Borneo, the gold ores contain about 1 per cent. of antimony as stibnite; nevertheless a satisfactory extraction of the gold is obtained at a reasonable consumption of cyanide. The ore is only coarse-crushed, consequently the stibnite passes into the vats in comparatively

large pieces, which expose only a small surface to the action of the solution.

**Loss due to Arsenic.**—Native arsenic is found in the gold ores of Coromandel, in New Zealand, and of Upper Sarawak. It is not acted on by cyanide or alkali, but realgar and orpiment are readily soluble in caustic alkali, and decompose cyanide even in the absence of free alkali. The arsenic is deposited in the zinc, and partially volatilized during the preliminary roasting of gold-zinc slimes.

**Action of Cinnabar.**—Cinnabar is found in the gold ores of Mercur, Utah, and Thames, New Zealand, and some other places. It is not acted on by cyanide. The telluride of mercury, coloradoite, which is found in some Kalgoorlie ores, is readily soluble in cyanide. The soluble salt of mercury is a desulphurizing agent, and by the benefit it confers in this way it partly compensates for the oxygen and free cyanide consumed in its dissolution.

**Action of Tellurium.**—At Cripple Creek and Kalgoorlie much of the rich gold ores are associated with various tellurides. It is doubtful whether these tellurides react directly with cyanide, but the gold is often so intimately entangled in them that they render its extraction slow and difficult. It was to obtain adequate extractions of the gold that fine grinding and sliming was introduced at Kalgoorlie in the raw-treatment of the telluride ores with cyanide and bromo-cyanide.

**Action of Selenium.**—The bullion of the Waihi Company contains about 1 per cent. of selenium. In what form the selenium exists in the ore is unknown, as no trace of selenium can be detected in any part of the mine. The presence of the selenium proves conclusively the soluble character of either selenium or of some of its combinations.

**Action of Cyanide Silver Sulphide.**—In argentiferous gold ores the bulk of the silver outside of the oxidized zone occurs in the form of sulphides, of which the simple sub-sulphide $\text{Ag}_2\text{S}$ is the most common. The dissolution of the sub-sulphide and other sulphides of silver, as we have already seen, liberates sulphur, part of which combines with potassium to form the alkaline sulphide $\text{K}_2\text{S}$, and part with the cyanogen molecule to form sulpho-cyanide, which is only harmful in that it robs the solution of available cyanide.

Alkaline sulphides are powerful reducers, and always cause a loss of available cyanide in the leaching of gold and silver ores.

In the leaching of gold ores the injurious effect of an alkaline sulphide in the solution when present in small amount is not appreciable, as the sulphatizing effect it exerts on the gold is
neutralized by the excess of available cyanide which is always present in the solution.

In leaching silver ores, the dissolution of the silver sulphide, so long as solutions containing free cyanide are applied, continue to liberate sulphur, with the formation of alkaline sulphide and sulpho-cyanide. The alkaline sulphide will precipitate or reduce the dissolved silver as Ag₂S, which will require a fresh supply of cyanide to redissolve it. This reaction will tend to reduce the amount of available cyanide in the leaching solution, and hence retard the dissolution of the remaining silver in the ore. It is manifest that the richer the silver ore, the greater will be the consumption of cyanide.

A high extraction can always be obtained from argentiferous gold ores by ordinary percolation by treating the vat charge first with a strong cyanide solution, say 0·4 or 0·5 per cent. KCy, and then adding lead acetate to the subsequent weak solutions and washes to the amount altogether of, say, 0·516 per ton of ore.

The treatment of silver ores necessitates the use of a larger volume of cyanide solution than that of ordinary gold ores. The artificial aeration of the solutions before use exercises a most beneficial effect by replacing the oxygen removed in the dissolution of the gold and formation of sulpho-cyanides.

**Action of Lead Acetate.**—Lead acetate is a desulphurizer, and when added to a cyanide solution containing an alkaline sulphide or sulpho-cyanide removes the sulphur as PbS.

In the treatment of gold ores containing metallic sulphides soluble in cyanide, such as those of silver or copper, or sulphides soluble in caustic alkali in presence of cyanide, such as those of arsenic or antimony, it will add considerably to the efficiency of the solutions to use a small amount of lead acetate as a desulphurizer.

The action of the lead acetate is beneficial, whether the treatment is effected by agitation or percolation. But it must be clearly understood that perfect aeration of the solutions is necessary to ensure the highest extractions.

**Action of Mercuric Chloride.**—Mercury has a greater affinity for sulphur than for cyanogen, hence it is a desulphurizer, and removes sulphur from the alkaline sulphides and sulpho-cyanides (thio-cyanates) by formation of the sulphide of mercury which is insoluble in cyanide.

Mercuric chloride performs the same function as lead acetate, but is more expensive and dissolves more slowly in water or cyanide solution.

**Action of Sulpho-Cyanides.**—It has been held by some metallurgists that the presence of these in working solutions of
cyanide is injurious, retarding the dissolution of the gold. As the result of much research, Godfrey Doveton, formerly of Camp Bird Mills, Ouray, Colorado, informs the author that he has found that a solution containing potassium sulpho-cyanide up to a certain point was more active than a solution of corresponding strength in KCy with sulpho-cyanide present, and that even 2.50 grams of KCyS in 100 c.c. of solution did not influence the extraction unfavourably. The salt alone, in solution in water, is a slow solvent for gold.

Experiments on mill solutions show much the same result, and he has come to the conclusion that the presence of even considerable quantities of sulpho-salt in working solutions should not cause any uneasiness.

The Action of Manganese Oxides on Cyanide.—During the treatment of a parcel of ore from the Komata gold-mine, near Waitekauri, the author found there was an unusual consumption of cyanide. The ore consisted of soft, mullucky, friable quartz, coloured quite black by a large percentage of pyrolusite and wad, and containing a trace of nickel and cobalt.

A series of experiments were afterwards made to determine the cause of the loss, and the results of these at first led the author to the conclusion that the manganese oxides oxidized a portion of the cyanide to cyanate. It is well known that pyrolusite parts with a portion of its oxygen under the influence of heat alone, but more readily so in the presence of an easily oxidizable substance. Further research showed, however, that the loss was mainly due to the cobalt in the ore, which dissolved somewhat readily, thus consuming cyanide. It is interesting to note that the dissolved cobalt was precipitated with the gold and silver on the zinc in the extractor box, and, like copper, was found to interfere with the precipitation of the gold.

The Action of Oxygen-bearing Agents.—It seems probable that in ores containing copper or other base metal soluble in a solution of potassium cyanide, the base metal would, from its preponderance in the ore, necessarily utilize the greater amount of the available oxygen, thereby tending to render the dissolution of the gold slow and imperfect. Hence it is reasonable to conclude that the employment of an oxidizing agent that would supply the deficiency of oxygen in such complex gold-containing ores would be beneficial. That the use of an oxygen-bearing agent is justified in special cases, seems well established by the experience of the author at Kuaotunu in 1893, and the results reported by reliable metallurgical chemists in South Africa, America, and Western Australia.

The Action of Oxygen in Cyanide Solutions.—Accord-
ing to Elsner’s equation, oxygen takes an active part in the chemical dissolution of gold in cyanide solutions, a view strongly supported by many chemists. Others maintain that oxygen takes no active part in the reaction. They believe that the solution of the gold being electro-chemical, the part played by the oxygen is to combine with the film of hydrogen which forms on the surface of the metal as dissolution progresses. By preventing polarization the dissolution of the metal thus goes on uninterrupted.

The necessity for oxygen in the solutions, introduced either by aeration or the addition of an oxidizer, is now recognized by all metallurgists and chemists.

**Action of Bromo-Cyanide.**—Sulman and Teed in 1894 recommended the use of a halogen compound, such as CyBr, in cyanide solutions, to accelerate the dissolution of the gold and silver. The use of cyanogen bromide in the cyanide solutions for the treatment of slimes forms a prominent feature of the Diehl process now in successful operation at Kalgoorlie.

The part played by the halogen salt is unknown, but it probably acts beneficially in performing the part of an oxidizer.

**Influence of Temperature on Dissolution of Gold.**—The chemical dissolution of gold in a solution of potassium cyanide is accelerated by a rise of temperature, on account of the greater freedom with which ionization takes place. But with rise of temperature there is a decrease in the quantity of oxygen absorbed by the solution; hence the useful effect of the increased temperature is neutralized by the diminished amount of oxygen held in the solution.

At ordinary temperatures there is a loss of available cyanide due to the liberation of HCy by atmospheric CO₂, and in the absence of free alkali in the solution to unite with the HCy the loss may be considerable, especially during the circulation or agitation of solutions.

In practice there is a serious loss of available cyanide when the solutions are heated, due to the greater activity of atmospheric CO₂ and the increased dissolution of base metals due to the rise of temperature.

Attempts have been made to cyanide ores in closed vessels artificially heated, but the extra extractions did not compensate for the additional cost.
CHAPTER III.

LABORATORY EXPERIMENTS.

The cyanide process is essentially a chemical one, and a commodious and well-equipped laboratory forms one of the most important and necessary parts of the whole plant.

It is the duty of the metallurgist in charge to determine, by actual experiment, the lowest strength of cyanide solution required to extract an adequate percentage of the gold, and also to devise means of overcoming the problems which are inseparable from the treatment of different classes and grades of ore, with so active a compound as potassium cyanide.

The daily output of ore from a mine is subject to continual change, both as regards physical condition and chemical composition, hence the treatment requires, within certain limits, corresponding modifications, to obtain the maximum extraction at the minimum cost. To obtain these results, the metallurgist must be a trained analytical chemist, full of resource and originality.

The testing and valuing of the ores before, during, and after treatment must be entrusted to a careful and trustworthy assayer. The testing and making up of the working solutions are very simple operations, which may be left to experienced and intelligent foremen who possess a knowledge of arithmetic extending as far as decimals.

With free-milling gold ores the actual working extraction will generally be as high as that obtained in the laboratory, but, in the case of ores containing copper or antimony, too much reliance must not be placed on the laboratory experiments.

The author's experience is that high extraction may be obtained in the laboratory from ores totally unsuited for treatment by the cyanide process on a working scale.

The conditions on the one hand are theoretical, on the other actual, and before adopting the cyanide treatment for a sulphide or other ore, working experiments should be made on parcels ranging from two to five tons, in order to ascertain the consumption of
cyanide and actual extraction. If the working trials are successful the cyanide treatment may be adopted with confidence.

On the other hand, in the case of an ore containing comparatively coarse gold, the laboratory experiments—where the sample is hand-crushed—will give lower results than those obtained in practice in the cyanide works. The author made a number of experiments on an ore from Marlborough, N.Z. The average extraction in the laboratory was under 40 per cent., while the cyanide plant extracted over 60 per cent. At the battery the ore was dry-crushed through a 60-mesh screen, and investigation showed that a large portion of the gold was reduced fine enough to pass through the screen, and thus became amenable to cyanide treatment.

THE ACTUAL EXPERIMENT.

1. Procure six bell-jars, six or eight inches in diameter. In the neck of each jar fit a cork, perforated with one hole. Through the hole pass a short length of glass tube, on the end of which place a few inches of pliable black rubber tubing. On the end of the rubber tubing place a screw-clip, by means of which the rate of percolation of the cyanide solutions can be regulated to a nicety.

2. Now invert the jars, and fix them in a wooden frame, so as to stand upright. In each jar place a thin layer of small rounded pebbles, about the size of French beans; above the pebbles place an inch of coarse sand, and above this, half-an-inch of fine sand. Above the fine sand place a piece of loose scrim, the diameter of the jar. This completes the filter-bed.

When a large number of cyanide experiments are being made, a box divided into three compartments, to hold the three grades of material for the filter-bed, should be kept well replenished and near at hand.

3. Next procure a fair sample of the pulverized ore to be tested, weighing, say, six or eight pounds. Mix thoroughly, and carefully assay to accurately determine the original value.

Check assays should always be made, and if there is a serious discrepancy between the assay and its check, amounting to over 3 per cent. of the value, fresh assays should be made. The assays form the basis of the calculations and final results of the experiments, and hence the greatest accuracy should be aimed at.

When the ore to be tested is from the battery, or mill, it should be placed in a jar in the condition that it comes from the mill, except, of course, when the tests are to determine the degree of fineness, which would give the best economic extraction.
When the ore is hand-pulverized, a separate portion should be reduced to pass through, say, a 30-mesh, 40-mesh, and 60-mesh sieve respectively. Separate tests should be made of each grade, so as to determine to what extent the extraction is affected by the varying fineness of the ore.

4. Introduce into each jar, say, six pounds of the powdered and sampled ore, the value of which has been obtained by careful assay. Mark the jars 1, 2, 3, 4, 5, and 6.

5. In the case of tailings or ores containing iron pyrites, or other base metallic sulphides, the samples in the jars should be washed once or twice with clean water to remove any soluble sulphates. With very acid tailings a very dilute alkaline wash may be applied. (Test for acidity, see Chapter IV.)

Pan off a portion of the sample to ascertain what proportion of the gold is fine and what coarse. If the test is of importance a complete analysis should be made of the sample, so as to be able to adapt the procedure of treatment to the character of the constituents present.

6. No general rule can be laid down as to the strength of the cyanide solutions to be used, as this will depend as much on the character of the sample as on its gold value; but the strength of solution used on a working scale for any class of ore seldom exceeds 0.4 per cent. All the ores of silver, copper, arsenic, and antimony act on and consume cyanide, and when either of these is present a stronger series of solutions will have to be tried than in the case of clean ores. With the latter a useful series of solutions would contain—

0.05%, 0.1%, 0.15%, 0.2%, 0.25%, 0.3% of cyanide.

With pyritic ores or tailings, or those containing copper, antimony, or arsenic compounds, the most instructive series would be—

0.15%, 0.20%, 0.25%, 0.3%, 0.35%, 0.4%.

In the event of all the tests being unsatisfactory, it would be advisable to try both stronger and weaker solutions than those already employed, according as the character of the ore or material may suggest.

It is necessary with every new ore to make a number of laboratory experiments to ascertain the strength of cyanide solution required to extract an adequate proportion of the gold and silver contents.

7. (a) To each jar, already charged with the ore, add sufficient cyanide solution, of say 0.3 per cent. KCy, to stand just level with the surface of the ore. In most cases it will be found that
the weight of cyanide solution required will be about one-third that of the ore.

(b) Allow the solution to stand in contact with the ore from twelve to eighteen hours.

(c) Open the stop-cock, or screw-clip, and regulate so that the sands will be drained dry in about four hours. Collect the solution in a suitable vessel.

(d) Allow the sands to stand dry for six hours.

(e) Then run on an equal weight of weak solution, say 0·05 per cent. KCy; allow to stand in contact with sands for six hours, and then drain dry in four hours.

(f) Allow to stand dry for six hours and then apply a second weak wash. Let this weak solution stand in contact with sands six hours and then drain dry in four hours.

(g) After last draining apply a water-wash of same volume as last cyanide solution, regulating the screw-clip so that the draining will take twelve or eighteen hours.

(h) Then apply a second water-wash, which may be allowed to run off as rapidly as possible.

A complete record of the procedure should be kept for each jar, noting the following points: —

(a) Date and place of test.

(b) Description of sand and degree of fineness.

(c) Weight of sample and number of jar.

(d) Assay value of ore. State gold and silver contents.

(e) Weight and strength of strong solution used.

(f) Time in contact of solution with sand.

(g) Time in draining solution.

(h) Compare strength of sump solution with original strength

(i) Compare weight of solution with original weight.

(j) Time allowed for aeration, i.e. standing dry.

(k) Weight and strength of weak solutions.

(l) Time in contact with sands.

(m) Time in draining.

(n) Compare strength of weak sump solution with original strength of weak solution used.

(o) Weight of water-washes and time of draining.

(p) State degree of acidity of sands and nature of water or alkaline wash applied before leaching with strong solution.

(q) State if a desulphurizing agent were used.

Should the subsequent assay of the residues show that the extraction is low, the procedure may be modified in many ways. For example, the strong solution can be run on several times in succession, with aeration between each application, before applying the weak solutions. Or, again, the sample of sand can be first
treated with a weak solution, say 0·05 per cent. KCy, before applying the strong. Furthermore, it must be remembered that with many ores time is the most important factor.

8. Test the strength of the spent solution to ascertain the percentage of cyanide used. The solution and washings are collected and measured together, then tested for cyanide.

The consumption of cyanide can be calculated by determining the strength of the combined solution and washings, and making an allowance for the increase in bulk due to dilution.

On a working scale the consumption of cyanide is generally much less than that shown by the laboratory experiments.

Sometimes the cyanide and different washings are kept separate and evaporated down with the addition of litharge (in the manner described under The Assay of Cyanide Solutions), and the gold actually extracted by each calculated separately. The results afford an instructive lesson on the value of successive washings.

9. Remove the leached and washed residues from the jars, dry, mix thoroughly and again assay. As the residues will probably be excessively low-grade, it will be necessary to take 1000 or 1500 grains for the assay determination. Then calculate the percentage of extraction from each jar by difference, recording the results and assay values as follow:—

<table>
<thead>
<tr>
<th>Original Value.</th>
<th>After Leaching.</th>
<th>Percentage of Recovery.</th>
</tr>
</thead>
<tbody>
<tr>
<td>ozs. dwts. gra.</td>
<td>ozs. dwts. gra.</td>
<td></td>
</tr>
<tr>
<td>Gold, . 2 4 12</td>
<td>0 4 12</td>
<td>89·9</td>
</tr>
<tr>
<td>Silver, . 1 0 4</td>
<td>0 3 6</td>
<td>83·9</td>
</tr>
<tr>
<td>Value, . £9 0 0</td>
<td>£0 18 4</td>
<td>89·8</td>
</tr>
</tbody>
</table>

The calculation is simply a matter of proportion. As an example take the recovery of gold:—

- Original gold, . 2 4 12 per ton.
- After leaching, . 0 4 12 ,,.

Extracted, . 2 0 0 ,,.

Then if two ounces were extracted from 2 ozs. 4 dwt. 12 grs., what would the extraction be from 100?

2 ozs. 4 dwts. 12 grs. : 2 ozs. : : 100 : 89·9 per cent.

\[
\frac{2 \times 100}{2\cdot225} = 89\cdot9 \text{ per cent.}
\]

An easy and expeditious method of calculating the percentages
of extraction in the laboratory-test is to use the weights of the bullion, gold, and silver (in grains or grams) as the basis of computation, instead of the same extended as ounces, dwts., and grains.

*Example:*

<table>
<thead>
<tr>
<th></th>
<th>Original Assay</th>
<th>After Leaching</th>
<th>Extracted</th>
<th>Recovery per cent.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bullion</td>
<td>0.0020</td>
<td>0.002</td>
<td>0.0018</td>
<td>90.0</td>
</tr>
<tr>
<td>Gold</td>
<td>0.0018</td>
<td>0.001</td>
<td>0.0017</td>
<td>94.4</td>
</tr>
<tr>
<td>Silver</td>
<td>0.0002</td>
<td>0.001</td>
<td>0.0001</td>
<td>50.1</td>
</tr>
</tbody>
</table>

*Value, £9 2 0*  
*£0 10 3*  
*£8 11 9*  
*94 3*

*The calculation:*

For Bullion.  
\[
\frac{0.0018 \times 100}{0.002} = 90
\]

For Gold.  
\[
\frac{0.0017 \times 100}{0.0018} = 94.4.
\]

For Silver.  
\[
\frac{0.0001 \times 100}{0.0002} = 50
\]

For Value.  
\[
\frac{8.587 \times 100}{9.1} = 94.3.
\]

10. Compare the results obtained, and adopt the strength which gives the highest extraction.

*Remarks.*—With a series of experiments it will be found that the percentage of extraction, or rate of dissolution of the gold, reaches a maximum with a cyanide solution of a certain strength, and that above and below this strength the rate of extraction rapidly diminishes. The strength of cyanide solution which dissolves the maximum percentage of gold will depend on the character of the ore.

It will generally be found that the finer the ore is pulverized the more rapid will be the rate of dissolution of the gold. This is especially the case with argentiferous gold ores and pyritic concentrates.

*Laboratory Test of Slimes.*—It is necessary with all ores to ascertain the probable economic advantage to be derived from sliming or fine-grinding either all or a portion of the sands.

Slime tests are best conducted in a small agitator barrel or vat about 16 inches deep and 8 inches in diameter. The stirrers are actuated by a small pelton wheel, turbine, electric or other convenient motor.

The original assay value and value of residues is calculated
from the dry weight of slimes. The extraction is found by
difference.

Slimes free from sulphurets may be agitated with very dilute
solutions even as low as 0·10 KCy in the absence of cyanicides;
and in most cases the dissolution of the gold will be effected in
from two to six hours. Slimes containing from 2 to 6 per cent.
of sulphurets require longer agitation and somewhat stronger
solutions, say 0·4 per cent. KCy; while clean concentrates may
require treatment for two or more days, and solutions ranging up
to 5 or more per cent.

In all tests the cyanide must be protected by the use of lime
or caustic alkali (NaOH), but the too free use of the latter must
be avoided in ores containing antimony, arsenic, tellurium, or
selenium.

It adds considerably to the value of the tests if all the
operations are conducted so as to imitate in miniature the treat-
ment on a working scale. This is especially important so far as
concerns thickening the pulp, using the same strength and
volume of solution for agitation, regulating speed of agitator,
etc.

In the treatment of concentrates, or slimes containing a con-
siderable proportion of sulphurets, the artificial aeration of the
pulp or the addition of an oxygen-carrying chemical will help the
dissolution of the gold. The addition of a little acetate of lead
is often beneficial, as it desulphurizes the solutions.

The test is conducted as follows:—
1. Charge the agitator with 10 or 15 lbs. of slime thickened to
about one part dry slime to one part water. Neutralize acidity.
2. Add the cyanide solution, in weight about one-third that of
dry pulp.
3. Agitate until a satisfactory extraction has been obtained.
4. After the agitation has continued two or three hours, test
strength of cyanide in the agitator. If necessary make up to
original strength, and add a little lead acetate if the slimes
contain silver sulphides, or base metallic sulphides soluble in
cyanide or caustic alkali.
5. Allow the pulp to settle, decant off strong cyanide. Then
agitate with a wash of 0·05 per cent. KCy, settle and decant
Finish with two water-washes. Then take sample for assay.
CHAPTER IV.

CONTROL, TESTING, AND ANALYSIS OF SOLUTIONS.

Dissolving Power of Different Cyanides. — The simple cyanides of the alkalis and alkaline earth metals, such as those of potassium, sodium, ammonium, calcium, magnesium, strontium, and barium are solvents for gold and silver, but they do not all possess the same dissolving power, nor are they all equally stable compounds.

In 1896 and 1897 the author* proved by numerous experimental tests that cyanogen, in the absence of a protective alkali, is not a solvent for gold. It is a solvent only when combined with an alkali or alkaline earth.

The relative dissolving power of the cyanides depends on the molecular weight of the salt and the ratio of the valency of the base. For example, the molecular weight of sodium cyanide is 49, and of potassium cyanide 65, and the valency of each being 1, the relative dissolving power will be as 49 is to 65. In other words, 49 parts or lbs. of pure sodium cyanide will dissolve as much gold as 65 parts or lbs. of pure potassium cyanide; or, again, 49 lbs. of 90 per cent. sodium cyanide will have the same dissolving power as 65 lbs. of 90 per cent. potassium cyanide, and so on. Thus we find that 1 lb. of sodium has the same dissolving power as 1·33 lbs. of potassium cyanide. The relative dissolving power of NaCy, and NH₄Cy in terms of KCy, is shown in tabulated form below:—

<table>
<thead>
<tr>
<th>Molecular Weight</th>
<th>Dissolving Power per cent.</th>
</tr>
</thead>
<tbody>
<tr>
<td>KCy 65</td>
<td>100</td>
</tr>
<tr>
<td>NaCy 49</td>
<td>132·7</td>
</tr>
<tr>
<td>NH₄Cy 44</td>
<td>147·7</td>
</tr>
</tbody>
</table>

On account of its greater dissolving power it would be more


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THE CYANIDE PROCESS.

It is economical to use NH₄Cy than NaCy, and NaCy than KCy if all were sold at the same price and were equally stable compounds. The practice, however, is to sell NaCy and NH₄Cy in terms of the potassium cyanide cyanogen equivalent. Thus a consignment of 98 per cent. NaCy will be reported as equivalent to 130 per cent. KCy, and charged for accordingly.

Sodium cyanide is somewhat less stable than KCy, and NH₄Cy much less than NaCy. For this reason it is not found economical to use NH₄Cy in the extraction of gold by percolation or agitation in open vats.

Free Cyanide.—The term free cyanide is used to indicate the cyanogen present in solution as a simple cyanide of the alkalis or alkaline earths.

Dissolving Power of Potassium Zinc Cyanide.

It was long believed by cyanide operators that the cyanide which combines with zinc to form potassium zinc cyanide in the precipitation of gold with zinc was lost, and was of no further use for the recovery of gold. It is now known that this is not the case. Goyder,* Crosse,† and others have shown K₂ZnCy₄ to be a solvent for gold in the absence of free KCy, or a protective alkali.

Crosse experimented on spitzluten concentrates with a solution containing all the cyanide as K₂ZnCy₄, and obtained a satisfactory extraction, but the subsequent precipitation with zinc was a failure.

The extraction amounted to 83.4 per cent., and the precipitation to only 36 per cent. The failure to obtain a satisfactory precipitation was no doubt due to the absence of free KCy in the solution. The addition of KCy at the head of the zinc box would probably have brought about an adequate precipitation.

Feldtmann ‡ says that the “addition of alkali to working solutions which have become weak in alkali brings up the strength by regenerating (i.e. decomposing) the zinc cyanide, so that, as a matter of fact, when the solutions are pretty strongly alkaline they contain no zinc as cyanide, but only the hydrate dissolved in alkali (zincate of potash).”

Goyder and Bettel support this view, but Wells, Ellis, and Clennell consider it erroneous.

Wells§ states definitely that K₂ZnCy₄ is not decomposed by alkali into KCy and potassium zincate.

Clennell* thinks it doubtful whether KCy and potassium zinate can coexist in the same solution. He believes that the zinate only forms when silver nitrate is added to the solution in sufficient amount.

TO TEST THE STRENGTH OF CYANIDE SOLUTIONS.

The testing of cyanide solutions is an operation of great simplicity, and can be performed with accuracy and expedition by any intelligent foreman by a volumetric method of estimation. The standard solutions should always be made up under the personal supervision of the chemist in charge of the works.

Three different volumetric methods may be used for the determination, namely:

1. By standard solution of silver nitrate.
2. By standard solution of mercuric chloride.

By Standard Silver Nitrate Solution.

This is the method generally adopted in cyanide plants. It is a modification of Liebig's volumetric estimation of cyanogen.

The reaction depends on the fact that when a solution of silver nitrate is added to a solution of potassium cyanide, the cyanogen unites with the silver, appearing as a white precipitate (AgCy), which is immediately dissolved by any free KCy, that may still be present, forming a double cyanide of potassium and silver.

This reaction is shown by the equations—

\[
\text{AgNO}_3 + \text{KCy} = \text{AgCy} + \text{KNO}_3;
\]

and

\[
\text{AgCy} + \text{KCy} = \text{AgKCy}_2.
\]

A standard solution of silver nitrate can be made up from the molecular weights of the constituents as follows:

\[
\begin{align*}
\text{AgNO}_3 & \text{ saturates } 2\text{KCy.} \\
169.955 & = 130.04. \\
169.955 & = 13.004.
\end{align*}
\]

With grams, use a decinormal solution; then if 16.955 grams of silver nitrate are dissolved in 1000 c.c. of water, 1 c.c. will be equal to 0.013 grm. of KCy.

To Make DECINORMAL STANDARD.—Take 16.955 grams of silver nitrate (triple-crystallized if procurable), and dissolve in

one litre (1000 c.c.) of distilled water. In large works, where much testing is going on, it is advisable to dissolve 33.91 grams in two litres; then place in stoppered-bottle and mark.

To Test Solutions of KCy:—

1. Fill a burette with silver nitrate solution.

2. Measure 13 c.c. of cyanide solution to be tested from another burette and transfer to a smaller beaker. To obtain accurate results add a few drops of potassium iodide solution to the beaker and shake.

3. Run in standard AgNO₃ solution cautiously from the burette till the white precipitate formed just ceases to redissolve when the beaker is shaken; that is, when a faint permanent opalescence appears the reaction is complete.

4. Read off number of c.c. of standard solution used, and divide by 10. The result will represent the percentage of available KCy. For example:—

Suppose 13 c.c. of KCy sol. took 14.5 c.c. of AgNO₃, then—

\[
\frac{14.5}{10} = 1.45\% \text{ of KCy.}
\]

Remarks.—Two burettes should always be used; one to measure the cyanide solution, and one for the silver nitrate standard. The gram burette should be graduated to 1.10th gram. Erdmann floats should always be used, so as to obtain the exact reading.

The end reaction is very distinct and easily seen, more particularly against a dark background. The beaker containing the cyanide solution to be tested should be perfectly clean and placed in a good light.

When the solution to be tested is turbid it must be filtered, as the end-point cannot be accurately determined unless the solution is perfectly clear.

The addition of 5 c.c. of a 1 per cent. neutral solution of potassium iodide, or a few drops of a 10 per cent. solution, to the solution to be tested renders the end-reaction more defined, and reduces the danger of over-estimating the KCy due to the presence of alkali or ammonia. Besides, the yellowish tinge of the silver iodide makes the end-point more distinct.

In an interesting paper on "The Titration and Use of Cyanide Solutions," Mr. Walter H. Virgoe, Chief Chemist to the Mexican Gold and Silver Recovery Company, shows that where copper is present in the solution, the use of an iodide indicator fulfils the
further purpose of indicating the point where the silver nitrate has titrated all the free cyanide, and is about to attack the cyanide in chemical combination with the copper, thereby preventing the over-estimation of the free cyanide. As an example, he says that a solution containing 0·3 per cent. of copper and titrating 0·52 per cent. of cyanide with silver nitrate alone may titrate only 0·13 per cent. of KCy correctly if KI be used.

Virgoe finds that in titrating pure solutions of potassium cyanide, the amount of indicator used makes no difference whatever, but when copper is present, even in small proportion, he shows that with different amounts of indicator very dissimilar percentages of cyanide are obtained. For this reason he points out the advisability of using a minimum addition of KI before titration of solutions containing copper.

To Test Very Dilute Solutions.—To test the strength of very dilute cyanide solutions, measure off 130 c.c. of the solution, titrate with silver nitrate, and divide the number of c.c. of standard required by 100, and the result will give the percentage of available KCy, thus:

\[ 130 \text{ c.c. of cyanide solution required } 5 \text{ c.c. of standard, then} \]
\[ 5 \div 109 = 0.05\% \text{ KCy}. \]

To Test Strong Solutions.—In very strong solutions the silver cyanide, AgCy, which forms when the silver nitrate is added from the burette, has a tendency to come down in a more or less granular form. It settles to the bottom of the beaker rapidly, and even when well shaken redissolves slowly and with difficulty. For this reason all solutions containing over 1 per cent. of cyanide should be diluted with water before titration, a practice which will be found to give a more sharply defined end-reaction, and at the same time effect a saving of silver nitrate.

(a.) When using the decinormal standard, measure off 13 c.c. of the strong solution and dilute with water (preferably distilled) to 130 c.c. Then measure off 13 c.c. of this diluted solution and titrate with silver nitrate as described above. Note the number of c.c. of standard solution required to complete the reaction, and this will represent the percentage of KCy in the strong solution, for since the 13 c.c. of dilute solution contained only a tenth of the original 13 c.c. of strong solution, there is hence no need to divide the quantity of silver nitrate by ten.

(b.) Or of the strong cyanide solution measure off any aliquot part of the 13 c.c. required in the regular test. Take, say, 3, 4, or 5 c.c., dilute with distilled water to about ten times the volume, and titrate with silver nitrate. Note the number of c.c. of standard solution required to complete the titration.
Thus if 4 c.c. of the strong cyanide solution were taken, and required 6 c.c. of standard nitrate for titration, 13 c.c. would require 19·5 c.c. of standard; and 19·5 divided by 10 = 1·95% KCy.

**Modified Decinormal Standard.**—This standard is commonly used in South Africa and Australasia. Its use lessens the liability to make mistakes in reading the burette, and obviates calculation after titration, except what can be performed mentally and without an effort.

The standard solution is prepared by dissolving 13·04 grams of the re-crystallized silver salt in 1000 c.c. of distilled water. Every c.c. of this solution is equivalent to 0·01 gram KCy.

In testing, take 10 c.c. of the cyanide solution, titrate with standard solution in the manner described when using a decinormal standard, note number of c.c. required to produce a permanent opalescence, divide this number by 10, and the result will be the percentage of available or "free cyanide."

It should be noted that the number 13·04 bears the same ratio to 10 c.c. of solution to be tested that 16·955 bears to 13 c.c.

Thus:

\[ 13 : 10 : : 16·955 : 13·04. \]

**Testing Very Dilute Solutions.**—(a) For testing the very dilute solutions now used in cyanide practice, measure off 100 c.c. of solution to be tested, titrate with standard, and divide the number of c.c. of standard used by 100. The result will be the percentage of cyanide.

Thus—100 c.c. of a weak cyanide solution required 4 c.c. of silver nitrate to complete titration; then—

\[ \frac{4}{100} = 0·04\% \text{ KCy.} \]

(b) A customary and useful practice is to dissolve 13·04 grams of silver salt in 2000 c.c. of distilled water. Every c.c. of this standard is equivalent to 0·005 per cent. KCy.

Of the cyanide solution to be tested take 50 c.c., titrate with standard, and divide the number of c.c. used for titration by 100. The result is the percentage of cyanide.

Thus—50 c.c. of a very dilute cyanide solution required 4 c.c. of silver nitrate to effect titration; then—

\[ \frac{4}{100} = 0·04\% \text{ KCy.} \]

In this case we use a silver nitrate standard of half the strength used in method (a) and obtain the same results by taking 50 c.c. of cyanide solution instead of 100 c.c.
In small plants it is customary here to make up the silver nitrate standard by dissolving 6.52 grams of the silver salt in 1000 c.c. of water instead of 13.04 grams in 2000 c.c. This gives a standard of the same strength as in (b); therefore measure off 50 c.c. of cyanide solution, titrate, and divide c.c. of standard required by 100 to find percentage of free cyanide.

To Test Cyanide Solutions with Grain Standard Solution.

1. When grain burettes are used, make up a solution of silver nitrate by dissolving 169.55 grains of the salt in 10,000 grains of distilled water.
2. Measure off, from a burette, 130 grains of cyanide solution to be tested. Add, say, 5 c.c. of potassium iodide solution.
3. Fill a burette with the silver nitrate standard.
4. Run silver nitrate into cyanide solution until the white p.p. which at first forms just ceases to redissolve. Note the number of grains required.
5. The number of grains of silver nitrate solution used to titrate, divided by 100, will give the percentage of available KCy.
6. For example, if 56 grains of silver nitrate were used, then—
$$\frac{56}{100} = 0.56\text{ per cent. of KCy in the solution.}$$

By Standard Mercuric Chloride Solution.

(1.) When a solution of mercuric chloride is added to a solution of potassium cyanide, a cyanide of mercury is formed, but is at once dissolved by any excess of KCy present.

When all the free or available KCy has been used, a bluish-white opalescence of HgCy₂ appears if a slight excess of mercuric chloride is added. This permanent opalescence indicates the end of the reaction.

(2.) To Make up Standard Solution.—Use the equation—
$$\text{HgCl}_2 + 2\text{KCy} = \text{HgCy}_2 + 2\text{KCl}.$$  
271 saturates 130.  
27.1 = 13 in a decinormal solution.

From the above molecular weights, dissolve 27.1 grams of mercuric chloride in 1000 c.c. of distilled water; then 1 c.c. will equal 0.013 grm. of KCy. Place in a stoppered-bottle and mark.

(3.) The Actual Determination:—
(a.) Fill a burette with standard mercuric solution.
(b.) From another burette measure off 13 c.c. of the cyanide solution to be tested, and to this add about 3 c.c. of dilute ammonia.

(c.) Now run in standard mercuric solution very cautiously, with constant shaking, until a permanent bluish-white opalescence is produced.

(d.) Note the number of c.c. of standard required to complete the reaction; divide this number by 10, and the result will be the percentage of available KCy present; thus, if—

6·5 c.c. were required to complete titration, then—

$$6.5 \div 10 = 0.65\% \text{ KCy.}$$

Remarks.—With pure substances this reaction is very delicate, but with cyanide solutions, containing much impurity, it is not so reliable as the silver nitrate method. Caustic alkalis do not interfere with the reaction. The author has made a number of simultaneous tests, with working cyanide solutions, by the silver nitrate and mercuric chloride methods, and the results obtained were practically the same throughout.

By Standard Iodine Solution.

(1.) This method depends on the fact that when a solution of iodine is added to one of potassium cyanide, the iodine loses its colour so long as any undecomposed cyanide remains.

(2.) To Make up Standard Iodine Solution.—Use the reaction—

$$2I + KCy = KI + ICy.$$  
254 saturates 65.  
25·4 = 6·5 in a decinormal solution.

Therefore, to make a standard solution, weigh out 25·4 grams of iodine, place in a beaker with 200 c.c. of water, and add sufficient potassium iodine to completely dissolve the iodine with frequent shaking.

When the iodine is dissolved, make up to 1000 c.c. with pure water, and place in a stoppered bottle. Then—

$$1 \text{ c.c.} = 0.0065 \text{ grm. KCy.}$$

(3.) The Actual Determination—

(a.) Fill a burette with the standard iodine.

(b.) From another burette measure off 6·5 c.c. of cyanide solution to be tested, and to this add carbonic acid (20 c.c. of ordinary soda water will do) to convert the
caustic and mono-carbonate alkalis, contained in all commercial cyanide, into bi-carbonates.

(c.) Now run in standard iodine, cautiously and slowly until a slight but permanent yellow colour is produced.

(d.) Read off the number of c.c. of standard employed, divide by 10, and the result will be the percentage of KCy required.

Remarks.—This method does not give reliable results in the presence of sulphides, or when the cyanide solution is muddy or discoloured.

Titration of Working Solution containing Zinc.

In working solutions containing little or no free alkali, the presence of zinc is a disturbing element, as the zinc cyanide (ZnCy₂), which forms as a greyish-white precipitate when the silver nitrate is added, obscures the usual end-reaction. The zinc cyanide comes down in the later stage of the titration, refuses to redissolve, even with prolonged agitation, and renders it impossible for the operator to determine the point at which the permanent precipitate of silver cyanide forms. According to Clennell,* the addition of potassium iodide does not remedy the difficulty to any appreciable extent.

The difficulty of observing the end-point in presence of zinc is not a matter of grave concern to the cyanide operator, as the tests for strength are to some extent relative rather than actual. Discussing this point Clennell says: "As, however, it is generally only necessary to obtain relative results, and a knowledge of the real strength of the working solution in actual free potassium cyanide, or its equivalent, is not essential, the test (that is, the titration with silver nitrate) may be retained with advantage if the following points be strictly observed." The points he emphasizes are: (a) the absolute cleanness of the flask or beaker in which the titration is made; (b) making all tests in precisely the same manner, that is, using the same quantity of cyanide solution, and adding the same quantity of potassium iodide; (c) to add no water either before or during titration; (d) to filter turbid solutions if necessary, but not to add lime to clarify them.

To Test the Strength of Solid Cyanide.—The chief impurities in commercial cyanide are alkaline carbonates, and sometimes alkaline chlorides and sulphides in small quantities.

To test the strength of the solid salt for the free or available cyanide which it contains, proceed as follows:—

(1.) Break a cake in two, and select a piece, say, a pound in weight, showing the whole thickness of the cake in section.
(2.) Reduce this pound to a coarse powder, sample well, and further pulverize to a moderately fine powder.
(3.) Weigh out 1 gram of powdered and sampled salt.
(4.) Dissolve in distilled water and make up to 100 c.c.
(5.) Measure off 13 c.c. of this solution and titrate with deci-normal silver nitrate standard solution from a burette as previously described. Note number of c.c. of standard required to form a permanent p.p.; divide by 10, and this will give the amount of cyanide in 1 gram of the salt.

For example: Suppose 13 c.c. of cyanide solution required 7·5 c.c. of standard, then—

\[
\frac{7.5}{10} = .75 \text{ cyanide in 1 gram;}
\]

which is equivalent to 75 per cent. of KCy in the salt.

In the past few years sodium cyanide has been largely used in this country instead of potassium cyanide. It is sold in terms of its KCy equivalent, and its strength in its aqueous solutions is determined by using the silver nitrate standards prepared for testing solutions of potassium cyanide. Thus the equivalent in potassium cyanide of a pure sample of sodium cyanide would be 132·7 per cent. KCy.

To Make up Cyanide Solutions.—There are two different methods of making up solutions in common use in cyanide plants. In some cases the requisite amount of solid cyanide salt is added to the sump solution; in others, the working strength is made up by adding strong solution from the dissolving tank to the sump solution.

The following exercises will render these methods clear:—

1 lb. of pure KCy dissolved in 100 lbs. of water gives a 1 per cent. solution; therefore, if you have a vat containing 100 cubic feet of water to make up to, say, 0·6 per cent., you would require 37·35 lbs. of pure KCy. *

Thus \( 100 \times 62\frac{1}{4} = 6225 \) lbs. of water,

and if 100 lbs. of water require 0·6 lb. KCy, 6225 lbs. would require—

\[
\frac{100}{6225} = 0.6 : x
\]

\[
\frac{6225 \times 0.6}{100} = 37.35 \text{ lbs.}
\]

* See Constants at end of chapter.
CONTROL, TESTING, AND ANALYSIS OF SOLUTIONS. 37

Commercial cyanide is seldom pure; you would, therefore, have to use a greater quantity to make up the required strength.

Suppose the crude KCy contains 78 per cent. of KCy, then—

\[
\frac{78}{100} = \frac{34.35}{x} = \frac{78 \times 37.35}{100} = 48 \text{ lbs. crude KCy.}
\]

The same form of calculation will do for making up any required quantity of cyanide solution. Suppose 4 ozs. of a 0.5 per cent. solution were required.

Then if 100 ozs. of water require 0.5 oz. of cyanide, how much would 4 ozs. require?

\[
\frac{100}{x} = \frac{4}{0.5} = \frac{4 \times 0.5}{100} = 0.02 \times 480 = 9.6 \text{ grains.}
\]

If you have a 0.2 per cent. solution and you wish to make it up to, say, 0.5 per cent., subtract the 0.2 per cent. already in the solution from 0.5 per cent., leaving 0.3 per cent. required. Then proceed to make up as directed in the preceding paragraph.

EXERCISES.

(1.) I have 4000 lbs. of sump solution containing 0.2 per cent. of available KCy, which I wish to make up to a 0.5 per cent. solution: how much additional KCy will be required?

\[
\frac{100}{x} = \frac{4000}{0.3} = \frac{4000 \times 3}{100} = 12 \text{ lbs. pure KCy.}
\]

If your crude KCy salt contains only 82 per cent. of KCy, then—

\[
\frac{12 \times 100}{82} = 14.6 \text{ lbs. crude KCy required.}
\]

(2.) How many lbs. of solid cyanide salt of 75 per cent. strength should be used to make up 10 tons of a 0.4 per cent. working solution? Ans.—119.46 lbs.

(3.) How many lbs. of solid cyanide salt of 82 per cent. strength should be used to make up 5 tons of a 0.45 per cent. working solution, using a sump solution containing 0.15 per cent. of KCy for making up? Ans.—40.97 lbs.

(4.) How many lbs. of a 14 per cent. stock cyanide solution
should be used to make up 10 tons of a 0.4 per cent. working solution, using a 0.18 per cent. sump solution for making up.

Solution.—This is easiest determined by "Alligation," the team-mixer's rule of proportion, thus:

\[
\begin{align*}
\text{Strong solution} & 14.00 \times \cdot40 \quad \text{Working solution} \quad \cdot40 \times \cdot18 \\
\text{Working solution} & \quad \cdot40 \\
\text{Sum} & \quad \cdot18 \\
\text{Proportion of loss in strong solution} & = 13.60 + \cdot22 = \{ \text{Proportion of gain in weak} \\
\text{Neglecting decimals, then—} & \\
1360 + 22 & = 1382.
\end{align*}
\]

Here we have 1382 parts or lbs. of the required mixture, containing 22 of strong and 1360 of weak solution; therefore, if 22 lbs. of the strong solution give 1382 lbs. of the required mixture, how many lbs. will be required for 10 tons of the mixture?

\[
\begin{align*}
1382 : (10 \times 2240) & : : 22 : x \\
22400 \times 22 & = 356.58.
\end{align*}
\]

The answer is, therefore, 356.58 lbs.

The solution of this problem by a simple algebraic equation is easy, and involves less working. Thus—

Let \( x \) = quantity of 14 per cent. solution required.

Then the quantity of 0.18 per cent. sump solution will be—

\[10 - x.\]

Hence—

\[
\frac{(10 - x) \cdot18}{100} + \frac{x \cdot14}{100} = 10 \times \frac{\cdot4}{100}
\]

\[
1.8 - \cdot18x + \cdot14x = 4 \\
13.82x = 2.2 \times 2240 \\
x = 356.58 \text{ lbs.}
\]

(5.) How many lbs. of a 22 per cent. stock solution should be used to make up 9 tons of 0.5 per cent. working solution, using a 0.12 sump solution for making up? Ans.—350.1 lbs.

(6.) How many lbs. of a 12 per cent. stock solution should be used to make up 10 tons of a 0.6 per cent. working solution, using 0.15 per cent. sump solution for making up? Before drawing from the stock solution, first utilize 4 tons of a 0.8 solution already in the solution vat. Ans.—359 lbs.

Solution of No. 6.—First find out how much of the working
solution can be made up from the 4 tons of 0·8 per cent. cyanide solution, thus:

\[
\begin{align*}
\text{Strong solution} & \quad 0.80 \quad \times \quad 0.60 \quad \text{Working solution} \\
\text{Working solution} & \quad 0.60 \quad \times \quad 0.15 \quad \text{Sump solution}
\end{align*}
\]

Proportion of loss in \( \text{strong solution} \) = \( \cdot20 \) + \( \cdot45 \) = \{ Proportion of gain in weak. \}

Neglecting decimals, then—

\( 20 + 4.5 = 65 \) of required mixture.

Then, if 45 of the strong (0·8 per cent.) give 65 of the required mixture, the 4 tons already in the solution tank will give 5·77 tons, thus:

\[
45 : 4 : : 65 : x \\
\frac{4 \times 65}{45} = 5.77.
\]

And \( 10 - 5.77 = 4.23 \) tons to be made up from the 12 per cent. stock solution, thus:

\[
\begin{align*}
\text{Strong solution} & \quad 12.00 \quad \times \quad 0.60 \quad \text{Working solution} \\
\text{Working solution} & \quad 0.60 \quad \times \quad 0.15 \quad \text{Sump solution}
\end{align*}
\]

Proportion of loss in \( \text{strong solution} \) = \( 11.40 \) + \( \cdot45 \) = \{ Proportion of gain in weak. \}

Neglecting decimals:

\( 1140 + 45 = 1185 \) of mixture.

Now, if 45 lbs. of the stock solution (12 per cent.) give 1185 lbs. of the required mixture, how much will 4·22 tons require?

\[
1185 : (4.23 \times 2240) : : 45 : x \\
\frac{4.22 \times 2240 \times 45}{1185} = 359.15 \text{ lbs.}
\]

The algebraic solution of this problem is much shorter and easier than the above. It is as follows:

Let \( x = \) quantity of 12 per cent. solution to be added to sump solution.

Then the quantity of 0·15 per cent. sump solution required to make up the 10 tons will be:

\( 10 - (x + 4) \).
Hence—
\[
\left[ 10 - (x + 4) \right] \cdot \frac{15}{100} + 4 \cdot \frac{8}{100} + x \cdot \frac{12}{100} = 10 \cdot \frac{6}{100}
\]
\[(6 - x) \cdot 15 + 3 \cdot 2 + 12x = 6
\cdot 9 - 15x + 3 \cdot 2 + 12x = 6
11.85x = 1.9 \times 2240
x = 359.15 \text{ lbs.}
\]

TO DILUTE CYANIDE SOLUTIONS.

(7.) How many tons of a 0.45 per cent. working solution would 6 tons of a 0.8 per cent. solution of cyanide make, using water for dilution? Ans.—10.66 tons.

Solution:
\[
\frac{45}{80} : \frac{80}{6} : x
\]
\[
\frac{80 \times 6}{45} = 10.66.
\]

(8.) How many tons of a 0.4 per cent. working solution of cyanide would 8 tons of a 0.6 per cent. solution make, using a 0.12 per cent. sump solution for dilution? Ans.—13.71 tons.

Solution:

Strong solution \( \cdot 60 \) \( \times \) \( \cdot 40 \) Working solution
Working solution \( \cdot 40 \) \( \times \) \( \cdot 12 \) Sump solution

Proportion of loss in strong solution \( = \frac{\cdot 20 + \cdot 28}{20 + 28} = 48 \) of mixture.

Now, if 28 of the strong solution give 48 of the required mixture, 8 tons will give 13.71 tons, thus—
\[
8 \cdot 48 = 13.71 \text{ tons.}
\]

The algebraic solution is as follows:

Let \( x \) = required quantity of 0.4 per cent. solution.
Then—
\[
x - 8 = \text{quantity of 0.12 per cent. solution to be used.}
\]
Hence—

\[
8 \times \frac{0.6}{100} + (x - 8) \times \frac{12}{100} = x \times \frac{4}{100}
\]

\[
4.8 + \frac{12x - 96}{100} = \frac{4x}{100}
\]

\[
4.8 + 12x - 96 = 4x
\]

\[
28x = 384
\]

\[
x = 13.71 \text{ tons.}
\]

(9.) How many tons of a 0.6 per cent. cyanide solution would 8 tons of a 0.7 per cent. solution make, using a 0.2 per cent. sump solution for dilution? Ans.—10 tons.

THE ASSAY OF CYANIDE SOLUTIONS.

The estimation of the gold contents of cyanide solutions in large works where a great many determinations are made daily must be effected by a method both expeditious and accurate. Method I. can be used for the estimation of both gold and silver; and although somewhat slow, is probably the most reliable of all the methods proposed.

In order to get a bead sufficiently large to give a reliable weighing on the balance, it is necessary to take a relatively large sample of the solution, especially with poor sump solutions. The sample required will range from half a pint to three pints according to the richness. Where the Assay-ton system is preferred, from 20 to 150 A.T. may be taken. In this case 291.6 c.c. = 10 A.T.

Method I.—For Estimation of Gold and Silver.

(a.) Evaporation in Porcelain Dish.

(1.) Measure the sample of solution and evaporate in a large porcelain dish, or in an iron drying-dish enamelled inside. The bottom of the dish is covered with 400 or 600 grains of litharge.

(2.) Evaporate slowly to dryness over a Bunsen flame, or furnace lid. As the evaporation proceeds, rub down the sides of dish so as to collect the whole of the dissolved salts in the bottom. When large samples of solution are being assayed, the evaporation may be allowed to proceed until the bulk has been considerably reduced before adding the litharge; or the litharge may be sprinkled on the surface of the solution at the beginning of the evaporation. Care must be taken to avoid overheating the dish towards the end of the evaporation, as the residue may adhere too
firmly to the bottom and be difficult to remove. Some operators mix a little clean sand with the litharge.

(3.) Scrape the litharge and contents out of the dish with a clean spatula. If necessary rub down the sides with a piece of folded filter-paper moistened with a few drops of dilute nitric acid. Add the filter-paper to the flux.

(4.) The litharge, now containing the dissolved salts of the cyanide solution, is intimately mixed with the following flux, and fused in a clay crucible, as in the ordinary gold and silver fire-assay:—

Litharge, . . . 600 grains (added before evaporation).
Soda, . . . 400 ″
Borax (cover), . . . 100 ″
Glass-powder (or clean quartz sand), . . . 400 ″
Argol, . . . 50 ″

When fused, pour and allow to cool.

(5.) Cupel the lead-button and weigh the resulting bead of bullion. If the ore contains silver, part so as to determine the weight of gold and of silver; then for the pint measurement refer to the tables at the end of the chapter to ascertain the quantity of each per ton of solution.

When the resulting gold is weighed with gram weights, refer to the Gram Table; and when in grains, to the Grain Table (see end of chapter).

Remarks.—When a large number of determinations have to be made, ordinary enameled plates and mugs form efficient evaporating dishes; in this case also the litharge can be stirred into the solution before the evaporation begins.

(b.) Evaporation on Lead Foil.

The cyanide solution is assayed by evaporating a measured portion in a boat of sheet-lead, and then scorifying the residue and cupelling. When the solutions are charged with base metallic cyanides the results are not generally so reliable as when the solution is evaporated with litharge and afterwards fused in a clay crucible. Besides, this method is inconvenient when large volumes of poor solution have to be assayed, as a number of boats must be used for the same test.

This method may be used for comparatively rich solutions, but for everyday cyanide practice the method of evaporation in a dish with litharge is preferable, being the more reliable.
Method II. (Virgoe).—For Estimation of Gold.

(1.) To a litre of solution add excess of weak sulphate of copper solution.
(2.) Acidify with hydrochloric, nitric, or sulphuric acid.
(3.) Filter. The precipitate, which is white and flocculent, contains all the gold and silver. The bluish or greenish colour of the filtrate indicates that excess of copper sulphate has been added.
(4.) Wash precipitate, dry and place in a scorifier with metallic lead.

Crosse prefers to fuse the precipitate in a crucible with litharge and to cupel resulting lead button.

Notes on Virgoe’s Method.—In a discussion on his method before the Chemical and Metallurgical Society of South Africa, Mr. Virgoe supplied the following useful particulars:—

“In dealing with ordinary solutions, I add copper sulphate carefully until no further precipitate forms, stirring the solution in the meanwhile, and taking care to add only a very slight excess of copper sulphate, attested by the faint tinge of blue in the filtrate. Allow the precipitate to stand for a minute or two. $H_2SO_4$ is then added till the solution is acid, no more, no less. An addition of sodium sulphate produces in the filtrate a slight additional precipitate, which is found to contain a very small amount of gold. I find the results are very accurate. For instance, 1 litre of solution was found to contain, by careful duplicate evaporation tests, 8·27 milligrams; the copper sulphate method run in duplicate yielded 8·22 and 8·23 milligrams, just a little low, but the test is severe, and amply demonstrates to my mind the correctness of the method if properly run.

“As far as I have observed, there seems to be a tendency, if strong solutions of copper sulphate are added to strong solutions of cyanide, towards the formation of cupric salts; hence I use weak solutions of copper sulphate, and should expect bad results if the precipitation of gold were attempted by my method from a very strong solution of cyanide with a strong solution of copper sulphate.”

Method III.—For Estimation of Gold and Silver.

This is a modification of Professor S. B. Christy’s method by precipitation with cuprous chloride proposed by A. Whitby †:—

(1.) To 700 c.c. of the cyanide solution add 25 c.c. of a 10 per cent. solution of copper sulphate.

(2.) Then add a slight excess of sulphuric or hydrochloric acid 5 to 7 c.c., and boil for some time to remove the bulk of the free cyanide.

(3.) Now add from 10 to 20 c.c. of a 10 per cent. solution of sodium sulphite. The precipitate, which is chiefly cuprous cyanide, carries down with it the greater part of the gold and silver.

(4.) After vigorous shaking for about two minutes, the precipitate is filtered off and fused with a fluid in the ordinary way. Assays can be completed in three hours, and the results are said to compare favourably with the evaporation test. With solutions weak in cyanide, Whitby recommends the addition of some cyanide before making the test.

Method IV. (Crosse).—For Estimation of Gold.

(1.) Measure the sample of cyanide solution, and acidulate with nitric or sulphuric acid to expel most of the hydrocyanic acid.

(2.) Add silver nitrate solution until a precipitate ceases to form. The silver salt should be added a little at a time, and the solution well shaken after each addition. Most of the gold is carried down with the silver cyanide.

(3.) Allow the precipitate to settle; decant off the clear solution; filter and dry the precipitate and mix with 200 grains litharge, 100 grains glass-powder, 100 grains soda, and 48 grains of argol. Fuse, pour, and cupel the lead-botton.

(4.) Extract the bead of bullion from the cupel, flatten, and part without weighing.

(5.) Weigh the resulting gold and calculate the results.

This method cannot, of course, be used for the estimation of the silver contained in the solution. Clennell states that it gives fairly good results when the solutions are not too low in gold. It is preferred by some metallurgists, as large volumes of solution may be taken for assay, thereby effecting a saving of time.

Method V.—The Estimation of Gold.

In this method the gold is reduced in the acidified solution with zinc. Several modifications in procedure have been suggested. Two of these will be given.

(a) By Durant.*

(1.) Make the solution to be tested strongly acid with sulphuric acid and boil.

(2.) Then add about 30 grams of zinc shavings in quantities of 5 grams at a time, waiting each time till all action ceases before adding a fresh lot of zinc. The solution must be strongly acid, and kept boiling the whole time. After all the zinc has been dissolved there will be a slight residue of lead, carbon, etc.

(3.) Add about 1 gram of precipitated silica to collect the gold.

(4.) Filter through a double filter-paper and wash out the flask well, putting the rinsings on to the filter-paper.

(5.) Dry the paper (folded) on borax in a scorifier at mouth of muffle; char slowly; add granulated lead, or litharge and a reducing agent; scorify and cupel.

This method is reliable and rapid when assaying large volumes of solution. The test should be made in a porcelain basin.

(b) Chiddey.*

(1.) To 4 A.T. of the cyanide solution add 10 c.c. of a 10 per cent. solution of acetate of lead.

(2.) To the solution add 4 grams of zinc shavings. Boil a minute, and then add 20 c.c. of hydrochloric acid.

(3.) When all action has ceased, boil again. Decant off solution and wash lead precipitate with distilled water.

(4.) Transfer spongy lead with a glass rod to a piece of filter-paper; squeeze into a compact lump, and place in a hot cupel.

The mouth of the muffle should contain a piece of dry pine-wood, so that the muffle is filled with flame at the moment of introducing the spongy lead.

The assay takes about 25 minutes. Chiddey claims that the results obtained are somewhat higher than by the evaporation method.

Method VI. (Lindeman).†—For Estimation of Gold.

(1.) Heat 10 A.T. of cyanide solution until quite hot.

(2.) Add ammoniacal copper nitrate until the solution shows a permanent blue colour.

(3.) Then add sulphuric acid slowly to excess. Stir well and filter at once.

(4.) Fold filter-paper, carbonize slowly in a scorifier in muffle; transfer to a crucible; fuse and cupel.

It is claimed that this method checks well with the evaporation assay. The addition of litharge and fluxes should improve the results of the test.

* Alfred Chiddey, Australian Mining Standard, 19th Nov. 1903.
† Maurice Lindeman, Engineering and Mining Journal, New York, July 1904.
ANALYSIS OF CYANIDE SOLUTIONS.

Working solutions of potassium cyanide soon become contaminated with ferro-, ferri-, and sulpho-cyanides and cyanates of the alkaline earths and base metals soluble in KCy, some of which have a beneficial and others an injurious effect. In order, therefore, to be able to modify the treatment to obtain a higher rate of extraction, or to reduce the consumption of cyanide, or effect a better precipitation, it is necessary to know something of the constitution of the working solutions which generally contain, even in the treatment of the purest ore, an assemblage of complicated compounds requiring much technological skill for their estimation.

The analysis of cyanide solutions is an object to which many accomplished chemists, mostly in South Africa, have devoted much attention. Many useful schemes have been formulated, the object in most cases having been to devise reliable methods, devoid of too much refinement, yet capable of every-day application in cyanide works.

The "Shaking Test" for Consumption of Cyanide.—This method was much used in the laboratories of the Cassel Cyanide Company, and afforded a rapid and fairly approximate estimate of the consumption of cyanide with different classes of ore. It is useful for comparative purposes, and as a preliminary means of determining the most suitable strengths of cyanide solutions for laboratory experiments.

1. Take 200 grams of the ore and place in a stoppered bottle with, for example, 100 c.c. of a 0·5 per cent. cyanide solution, and shake for twenty minutes.

2. Allow contents of bottle to settle; draw off a portion of the clear solution with a pipette and test for KCy. If it contains 0·2 per cent. of KCy, then 0·3 per cent. has been consumed or decomposed.

3. When much cyanide is used up, test the ore for acidity by Feldtmann's method given below.

To Test Ores and Tailings for Acidity.—

1. Weigh out 224 grams of ore and shake up with 250 c.c. of water in a tall glass-jar or cylinder.

2. Fill a burette with a standard solution of soda, and titrate the ore-solution in the jar until the reaction is neutral to test (litmus) paper.

3. Every c.c. of the soda solution used will represent 0·1 lb. of caustic soda to be added to every ton of ore (or tailings) in a wash before the cyanide treatment.
CONTROL, TESTING, AND ANALYSIS OF SOLUTIONS.

To Make Standard Soda Solution.—Dissolve 10 grams (or 154·3 grains) of caustic soda in 1000 c.c. of pure water, and place in a secure bottle.

Remarks.—During the titration, the litmus paper should be dipped in clean, pure water from time to time to remove the adhering particles of ore so that the reaction may be clearly seen.

Tests for Alkaline Sulphides in Cyanide.—Alkaline sulphides act injuriously in cyanide solutions during leaching, and it is important to detect their presence. They are all soluble in water.

First Test:—To the clear cyanide solution add a little acid. If an alkaline sulphide be present, sulphur will be liberated, imparting a cloudy appearance to the solution.

Second Test:—In the clear solution place a clean, bright silver coin. It will become black and tarnished if a sulphide be present. This and the preceding test will not detect minute quantities.

Third Test:—The most delicate test is by means of the nitro-prussides. These are formed by adding a little nitric acid to a solution of ferro- or ferri-cyanide of potassium.

Add a few drops of a solution of nitro-prusside to the cyanide solution. If an alkaline sulphide is present, even in minute quantity, the solution will assume a brilliant purple colour.

Fourth Test:—When a dilute solution of a soluble lead salt, such as the acetate, is added to a solution containing an alkaline sulphide, a blackish brown precipitate of lead sulphide soon forms.

When the alkaline sulphide exists in the presence of free cyanide, a white precipitate of lead cyanide and carbonate will immediately form on the addition of lead acetate, thus tending to render the lead sulphide precipitate yellowish or nut brown.

THE ANALYSIS OF POTASSIUM CYANIDE.

(FELDTMANN AND BETTEL. *)

From 5 to 10 or more grams of the commercial cyanide are dissolved in water, and the insoluble matter, if any, filtered off.

The solution is agitated with a small quantity of precipitated lead carbonate—of course slightly in excess, and filtered. The precipitate, consisting of lead carbonate and sulphide, is transferred to a flask, and covered with a few c.c.'s of a solution of potassic or sodic cyanide free from sulphides, sulphocyanides, or ferrocyanides. This solution may be prepared from pure potassic or sodic hydrate and a solution of pure distilled hydrocyanic acid.

To the mixture in the beaker add hydrogen peroxide in slight excess— i.e., three or four times as much as is needed to whiten

* Paper read before the Chemical and Met. Society of S.A.
the precipitate. (The hydrogen peroxide for this purpose should be purified by agitation with ether and evaporation of the ether in a water-bath.)

A small quantity—say $\frac{1}{2}$ gram—of manganese peroxide is then added, and the mixture agitated for about two minutes, after which the solution is filtered off, acidified with sulphuric acid, and titrated with $\frac{N}{100}$ potassic permanganate.

1 c.c. of $\frac{N}{100}$ potassic permanganate equals 0·000053 grm. sulphur or 0·000182 grm. potassic sulphide.
The potassic permanganate may be standardized by means of pure potassic sulphocyanide.

1 c.c. = 0·0001618 grm. KCyS.

Estimation of Hydrocyanic Acid.—To 50 c.c. of the solution add a solution of bicarbonate of potash or soda, free from carbonate, or excess of carbonic acid. Titrate as for KCy. Deduct from this the amount of KCy found. The difference equals the amount of HCy present.

1 c.c. AgNO$_3$ = 0·0414 HCy.

Estimation of double Cyanides.—Add excess of pure caustic soda to 50 c.c. of solution, and a few drops of KI solution, then titrate with AgNO$_3$. Deduct KCy and HCy, as found above; the difference is K$_2$Zn(Cy)$_4$ as KCy, less 7·9 per cent.
The KCy as found here is calculated to K$_2$Zn(Cy)$_4$, as under:

KCy $\times$ 0·9493 = K$_2$Zn(Cy)$_4$. Add to this 7·9 per cent. of total, or for every 92·1 parts add 7·9 parts.

Estimation of Ferro- and Sulpho-Cyanides.—When organic matter is present, shake with powdered quicklime and filter.

A burette is filled with the cyanide solution for analysis, and run into 10 or 20 c.c. $\frac{1}{100}$ Normal K$_2$Mn$_2$O$_7$ strongly acidified with H$_2$SO$_4$ until the colour is just discharged.

A solution of ferric sulphate or chloride is acidified with H$_2$SO$_4$ and 50 c.c. of the cyanide solution poured in. After shaking for about half a minute the Prussian blue is separated from the liquid by filtration, and the precipitate washed. The filtrate is next titrated with $\frac{1}{100}$ normal K$_2$Mn$_2$O$_7$.

Let $x$ be c.c. permanganate required to oxidize ferro-cyanide, then $x = y - z$.

(x) 1 c.c. $\frac{1}{100}$ normal K$_2$Mn$_2$O$_7$ = 0·003684 grm. K$_4$Fe(Cy)$_6$.
(y) 1 c.c. $\frac{1}{100}$ normal K$_2$Mn$_2$O$_7$ = 0·0001618 grm. KCyS.
THE ANALYSIS OF CYANIDE WORKING SOLUTIONS.

(By Leonard M. Green, A.R.S.M.*)

The tests used depend essentially on alkalimetric determinations. In Mr. Green’s scheme the following constituents are estimated:—

1. The total cyanide = T.
2. The protective alkali = p.
3. The alkaline or alkaline-earth hydrates = h.
4. Alkaline mono-carbonates = N.
5. The ferrocyanides = S.
6. The zinc = Z.

The methods depend on the facts that:—

1. Potassium ferrocyanide is neutral to phenolphthalein.
2. That 1 c.c. of decinormal potassium ferrocyanide precipitates 0.75 c.c. decinormal zinc from a neutral dilute solution of a zinc salt.
3. That when a dilute neutral solution of a zinc salt is precipitated by adding an excess of sodium carbonate solution, the excess of alkali being afterwards carefully neutralized to phenolphthalein, by the addition of decinormal acid, a precipitate of basic zinc carbonate of almost constant composition is obtained. The precipitate obtained in this way is the normal basic carbonate.
4. That zinc hydrate or carbonate, when treated with an excess of potassium ferrocyanide, forms zinc ferrocyanide and potassium hydrate or carbonate, the alkalinity produced being proportional to the precipitate acted on. This reaction does not immediately proceed to the end, as at first only a portion of the alkalinity is formed; but if this be neutralized with acid a further amount of alkali is formed, and so on to the finish, the reaction taking a little time for completion.

The Actual Analysis.—(1.) The first estimation is that of the total cyanide. This is performed in the usual way by adding to the solution to be tested an excess of caustic soda and a little potassium iodide, and titrating with silver nitrate till a distinct, permanent yellowish cloudiness appears. It must be noted that the end-point is not reached until there is a distinct yellowish cloudiness.

Where much zinc and ferrocyanides are present, a faint, white cloudiness, probably due to the precipitation of zinc-ferrocyanide, is sometimes produced before the true "end-point." This must

* Extracts from Paper read before the Inst. Min. and Met., London.
be disregarded, the true "end-point" occurring when the yellowish cloudiness, due to silver iodide, is permanently formed. This is the only definite end-point in titrating such a solution, and a large excess of sodium hydrate does not appreciably alter it.

(2.) In the second test the alkaline and alkaline-earth hydrates plus half the mono-carbonates, viz., the "protective alkali," is determined. This test is a simple alteration of Cennel's.

Excess of potassium ferrocyanide is added to the solution, and then twice the amount of silver nitrate necessary to indicate the total cyanide, viz., sufficient to precipitate the whole of the cyanide. A slight excess does not matter, as it merely precipitates some chloride, or, if no chlorides are present, some sulphocyanide or ferrocyanide. The zinc all occurs in the precipitate as ferrocyanide, and the alkaline hydrates and carbonates are left in solution.

Phenolphthalein is then added, and the solution titrated with decinormal nitric acid till colourless, or till it acquires the faint greenish yellow tinge produced by the excess of ferrocyanide. The result indicates the "protective alkali."

Usually an identical result is obtained by only adding a little more than the amount of silver nitrate necessary to indicate the total cyanide, but this leaves potassium silver-cyanide in solution, and towards the close of the acid titration there is a slight chance of its being acted on by the acid forming AgCy, and setting free HCy, which would slightly increase the acid required, and obscure the end-point.

(3.) In the third test the alkaline hydrates are estimated. An excess of barium chloride is first added to the solution (sufficient to precipitate the sulphates and carbonates), and then the procedure in the last test is repeated. The result obtained indicates only the alkaline hydrate.

In the event of no hydrates being present, bicarbonates probably exist in solution. They may be estimated by adding a known amount of standard sodium hydrate, and repeating the test, allowing, in calculating the results, for the amount of hydrate added.

(4.) The next estimation is that of the total cyanide + chlorides + sulphocyanides + ferrocyanides + any other salt precipitated by silver nitrate before the precipitation of chromate. This test, though not important in itself, is necessary for the subsequent determination of the zinc.

The amount of acid used in the second test is first added to the solution (viz., sufficient to neutralize the protective alkali). Then one drop of a strong solution of potassium chromate is added, and the solution is titrated with silver nitrate till there is a faint permanent reddish colouration.
Towards the end of the titration, the reddish colouration appears, and only goes again slowly on shaking and standing a few seconds. This is due to the fact that the action of the silver nitrate on the precipitated zinc ferrocyanide is slow, and consequently some silver chromate is temporarily formed; this chromate, however, when well shaken up with the zinc ferrocyanide, is decomposed, and silver ferrocyanide is formed. The end-point therefore only occurs when the reddish colouration is decided and permanent.

(5.) The fifth estimation is that of the zinc — the ferrocyanide. Sufficient sodium carbonate is added to supply enough carbonic acid for the basic zinc precipitate in case there should not be enough already in the solution. Then twice the amount of silver nitrate necessary to indicate the total cyanide is added and the whole is well shaken. All the cyanide is now precipitated as silver cyanide, and any slight excess of silver nitrate will merely precipitate some chloride.

The zinc will have been partially precipitated as ferrocyanide by the ferrocyanide originally present in solution, but the remainder will have been precipitated as a basic carbonate, and there will be an excess of alkali in the solution. This alkalinity is neutralized by the addition of decinormal acid. During this neutralization the basic zinc carbonate gradually acquires its normal composition $3\text{Zn}((\text{HO})_2\text{ZnCO}_3$, and the neutralization must not be hurried.

The colour of the clear solution above the precipitate must not show a trace of pink, even on standing a minute or two, and the contents of the flask should be frequently shaken. In fact, when neutralization is apparently complete, it is best to add another c.c. of acid, shake up well and then add from a burette, a drop or so at a time, a solution of sodium carbonate (roughly $\frac{N}{10}$ is a convenient strength), till the clear solution again just shows a pink colour.

All the zinc which is not precipitated as ferrocyanide is now in the precipitate as basic carbonate, and the solution is neutral to phenolphthalein. If now an excess of potassium ferrocyanide be added, the basic zinc carbonate reacts with the ferrocyanide, forming zinc ferrocyanide and alkalinity proportional to the amount of basic carbonate present.

This alkalinity is then titrated with decinormal acid, the result representing the zinc less what has been precipitated by the ferrocyanide originally present.

Towards the end of the titration the colour is discharged and returns again slightly, so that a little time is necessary for the determination.
(6.) The last determination is of the zinc. Sodium carbonate is added as in the previous test; then silver nitrate solution is run in, in the same amount as was used in test No. 4, and the flask well shaken. The cyanides, chlorides, sulphocyanides and ferrocyanides will then have been precipitated as silver salts, and the whole of the zinc will have been precipitated as a basic carbonate, an excess of alkali remaining in the solution. This is neutralized precisely as in the last test, an excess of ferrocyanide added, and the alkalinity produced titrated with decinormal acid.

This result consequently represents alkalinity proportional to the whole of the zinc. It includes metals acting similarly to zinc, such as cadmium, and copper if present in small quantity.

Summary.

The following is a short summary of the tests described, with the quantities of solution taken, etc.:—

(1.) For Total Cyanide.—Take 50 c.c. of the solution. Add excess of sodium hydrate and a little potassium iodide solution. Titrate with silver nitrate till there is a distinct permanent yellowish cloudiness.

Result = T c.c.

(2.) Protective Alkali.—Take 50 c.c. of the solution. Add excess of potassium ferrocyanide solution. Run in $2T$ c.c. of silver nitrate. Add phenolphthalein and titrate with $\frac{N}{10}$ nitric acid till colourless.

Result = p c.c.

(3.) Alkaline Hydrates.—Take 50 c.c. of solution. Add excess of $\text{BaCl}_2$ solution, and then excess of potassium ferrocyanide. Run in $2T$ c.c. of silver nitrate, add phenolphthalein and titrate with $\frac{N}{10}$ nitric acid till colourless.

Result = h c.c.

(4.) Alkaline Mono-Carbonates.—Take 50 c.c. of the solution. Add $2T$ c.c. of silver nitrate, then p c.c. of $\frac{N}{10}$ nitric acid. Add one drop of a strong solution of potassium chromate, and continue titrating with silver nitrate till there is a faint permanent reddish coloration.

Result (Total $\text{AgNO}_3$ added) = N c.c.
(5.) Ferrocyanides.—Take 50 c.c. of solution. Add about 10 c.c. of sodium carbonate solution (roughly decinormal). Run in 2T c.c. (see Test 1) of silver nitrate, and shake well. Add phenolphthalein, and neutralize with $\frac{N}{10}$ nitric acid till the clear solution is colourless.

Shake well at intervals during the neutralization. Add about 1 c.c. more $\frac{N}{10}$ nitric acid, shake up, and then add a solution of sodium carbonate (roughly $\frac{N}{10}$ is a convenient strength), drop by drop, till the clear solution is just faintly pink.

Add excess of potassium ferrocyanide. The solution becomes strongly alkaline to phenolphthalein.

Titratre with $\frac{N}{10}$ nitric acid till colourless.

Result = S c.c.

(6.) For Zinc.—Take 50 c.c. of solution. Add about 10 c.c. of sodium carbonate solution. Run in N c.c. of silver nitrate (see Test 3), and shake well.

Add phenolphthalein, and neutralize as in the previous test.

Add excess of potassium ferrocyanide. The solution becomes strongly alkaline to phenolphthalein.

Titratre with $\frac{N}{10}$ nitric acid till colourless.

Result = Z c.c.

Notes.—If the standard solutions used are $\frac{N}{10}$ nitric acid, and a solution of silver nitrate containing 13·04 gram per litre, then the following are the factors for the different results:—

(1.) $T \times 0·02\% =$ Total Cyanides estimated as KCy.
(2.) $P \times 0·0112\% =$ Protective Alkali estimated as KHO.
(3.) $H \times 0·0112\% =$ Alkaline hydrates estimated as KHO.
(4.) $P - H \times 0·0276\% =$ Alkali carbonates.
(5.) $Z - S \times 0·0351\% =$ Potassium ferrocyanide.
(6.) $Z \times 0·0081\% =$ Zinc.

The presence of sulpho-cyanides or ferro-cyanides in small quantity does not interfere with these tests. The latter in large quantity does, as precipitates of zinc and silver ferrocyanides rapidly destroy the indicator phenolphthalein although potassium ferrocyanide itself does not appear to do so.
THE CYANIDE PROCESS.

A large excess of sodic carbonate has hardly any effect on the final result of the titration for zinc, but it tends to decrease the sharpness of the end-points, and increases the time taken for the reactions.

THE ESTIMATION OF OXYGEN IN WORKING CYANIDE SOLUTIONS.

(By Andrew F. Crosse.*)

Prepare a solution containing—

20 grams Iodide of Potassium
2 " Nitrate of Potassium.

Dissolve in distilled water and make up to 100 c.c.

The sample of cyanide solution can be taken in an ordinary Winchester quart bottle filled perfectly full.

Syphon off a sample into a separating funnel containing from 250 to 300 c.c., then add 1 c.c. of the solution of iodide and nitrate of potassium, and 3 c.c. of dilute sulphuric acid (1:1). After shaking up and allowing to stand for 15 minutes, titrate in an atmosphere of coal gas with a hyposulphite of sodium solution containing 7·75 grams per litre, 1 c.c. of which is equal to 0·25 m'grams of oxygen for the iodine liberated.

The peculiarity of the reaction is that no iodine is apparently liberated, and the cyanide solution remains colourless. But iodine has been liberated all the same to form a very unstable compound, iodide of cyanogen, and the amount is in exact proportion to the free oxygen plus the quantity to be subsequently allowed for, and can be estimated by the hyposulphite solution, using a starch solution as an indicator. Of course it is necessary to estimate the correction required for the iodine liberated by the reagents, and also in case any nitrates were present in the cyanide solution.

This correction is easily made in the following manner:—

Take about 400 c.c. of the solution under examination, add to it 0·3 gram pure ferrous sulphate and the same weight of caustic lime, shake up well and filter into a flask through which coal gas is passing, the precipitated ferrous hydrate will have absorbed all the free oxygen in the cyanide solution.

A good many experiments were made to test the above method, and the following will be a good example:—

A Winchester quart bottle was partly filled with a sample of an

ordinary working solution, the bottle was well shaken for some time in order to saturate the solution with oxygen.

The determination was made as described, using a pipette containing 292 c.c. of solution, 13·4 c.c. of hyposulphite of soda was required to unite with the iodine liberated.

Then 400 c.c. of the cyanide solution was freed from oxygen and the amount of hyposulphite required was 6·0 c.c.

So that 13·4 - 6·0 leaves 7·4 c.c. of hyposulphite required for the iodine liberated by the free oxygen in 288 c.c. of solution or 1·85 m'grams oxygen in 288 c.c., or 6·4 m'grams oxygen per litre.

In connection with the oxygen determination Mr. Crosse has made some interesting experiments on the amount of oxygen absorbed by sands and spitzluten concentrates.

He says: “I take, say, from 200 to 500 grams of the material under examination and shake it up for some hours in a large bottle full of a cyanide solution of known quantity and saturated with oxygen and then determine the amount absorbed or lost per litre, and so calculate out the amount of oxygen required by each kilogramme or ton of sands. I would, however, remark that though oxygen is being absorbed by certain substances during treatment, such as ferrous sulphides, etc., gold is also being dissolved, but more slowly and to a lesser degree than if no oxygen-absorbing matters were present. If we have ferrous hydrate present, which would almost immediately take up the oxygen, then no gold would be dissolved, but certain substances act slowly, allowing some gold to be dissolved at the same time that oxygen is being absorbed. If this were not the case it would often happen that no gold would be dissolved till several solutions had passed through the sands.”

DETERMINATION OF ZINC AND BASE METALS IN A CYANIDE SOLUTION.*

(By ANDREW F. CROSSE.)

The following is an accurate method for determining copper, iron, zinc and nickel in solutions:—

Take from 500 to 1000 c.c. of solution, acidify with a slight excess of sulphuric acid, add five or six grams of pure acid sulphate of potassium, and evaporate to dryness in a platinum dish and then heat to dull redness in order to melt the mass. The metals are obtained as sulphates and can be separated and estimated in the usual way.

ESTIMATION OF ZINC.

Solutions on the Rand do not usually have more than a trace of copper. The chief metallic ingredient is necessarily zinc, and the estimation of this metal is often useful.

Take 300 c.c. of solution, add about a gram of cyanide of potassium and the same quantity of pure caustic potash or soda, heat nearly to boiling point and then add a slight excess of sulphide of sodium in solution. The zinc will be quickly precipitated as a sulphide, and should be collected on a filter-paper and washed with hot water. Then place the filter-paper in a wide-mouthed bottle of known capacity between 250 and 300 c.c.

This bottle must be provided with a well-fitting india-rubber bung, through which a moderately wide tube is inserted about 8 to 10 inches long.

Then fill up the bottle with a weak solution of pure ferric sulphate containing 5 to 7 per cent. of sulphuric acid, place the bottle or flask in a bowl of cold water and raise the temperature to boiling point. The reason for the glass tube will be apparent as it allows for the expansion of the liquid.

The zinc sulphide will have decomposed and reduced a proportionate amount of ferric sulphate to ferrous sulphate. When nearly cold, filter off the solution through a dry filter-paper, and take half the quantity contained in the bottle and titrate with decinormal permanganate of potassium.

1 c.c. = 0.0325 gram of zinc. Sulphide of zinc is, however, slightly soluble in weak cyanide solutions, but having made various experiments, it was found that one milligram of zinc per 100 c.c. of solution taken, would be the right amount to add as a correction to the results obtained.

Various solutions were prepared containing known quantities of the double cyanide of zinc and potassium with ferro-cyanide and sulpho-cyanide of potassium, and the results obtained were very satisfactory.

NOTES ON THE ESTIMATION OF PROTECTIVE ALKALI IN CYANIDE SOLUTIONS. *

(By Andrew F. Crossel.)

Pure cyanide of potassium becomes rapidly decomposed if brought into contact with air, owing to the presence of carbonic acid. Most South African tailings and slimes contain a certain

small percentage of free sulphuric acid, or basic ferric salts, which decompose cyanide of potassium. We must protect or prevent this decomposition as far as possible. Cyanide of potassium is an expensive material; we should therefore endeavour to prevent its unnecessary decomposition, by the use of some inexpensive alkali—such as lime—and it becomes necessary to have an accurate method for determining the amount of caustic lime or other protective alkali in solution.

Protective alkali is the alkali present which will unite with any acid before decomposition of the cyanide begins.

It is very important in many cases to be able to determine the percentage of this alkali exactly.

In my experiments lime (calcium oxide) is taken as a protective alkali, but the results can be calculated out for caustic potash or caustic soda, if so desired.

If an exact amount of sulphuric acid be added to a solution of pure cyanide of potassium containing the latter in excess, we have the following reaction:—

$$2\text{KCN} + \text{H}_2\text{SO}_4 = \text{K}_2\text{SO}_4 + 2\text{HCN} \quad \text{or} \quad 98 \text{ parts } \text{H}_2\text{SO}_4 \text{ liberate } 54 \text{ parts of hydrocyanic acid.}$$

Now if, instead of $\text{H}_2\text{SO}_4$ we take pure bi-sulphate of potassium, the following reaction takes place:—

$$\text{KHSO}_4 + \text{KCN} = \text{K}_2\text{SO}_4 + \text{HCN} \quad \text{or} \quad 136 \text{ parts } \text{KHSO}_4 \text{ liberate } 27 \text{ parts } \text{HCN} \text{ or one gram } \text{KHSO}_4 \text{ will liberate } 1.985 \text{ gram } \text{HCN} \text{ from a solution containing an excess of cyanide of potassium.}$$

Supposing, however, that the cyanide of potassium solution contains a free alkali, the sulphuric acid is neutralized, and less hydrocyanic acid is liberated, in direct proportion to such alkali present. If there be no protective alkali, I know that in presence of an excess of KCN one gram of KHSO$_4$ liberates 1.985 gram HCN.

If less HCN is liberated it practically amounts to its being retained by the alkali, so that if I take the equation—

$$2\text{HCN} + \text{CaO} = \text{Ca(CN}_2\text{)}_2 + \text{H}_2\text{O}.$$  
54 parts of HCN = 56 parts Calcium Oxide.

The proposition is now as follows:—

Take 500 c.c. of a cyanide solution, add one gram of KHSO$_4$, boil for 45 minutes, and collect the HCN in a flask containing caustic potash, passing the vapour through a Liebig’s condenser.

I estimate the HCN in the ordinary way in the potash solution with nitrate of silver.

Let A = amount of HCN liberated by one gram KHSO$_4$ in presence of excess of cyanide without any alkali.
Let \( B = \) amount found in experiment with alkali.
\[ A - B = C \] \( C \times \frac{4}{24} \) or \( 1.037 = \) amount of calcium oxide present.

I made a series of experiments with various proportions of carefully prepared lime water and cyanide solutions, and obtained very accurate results—as, for instance, the following. I took 300 c.c. distilled water containing two grams of pure cyanide of potassium, and added 200 c.c. of saturated lime water (at 18° Cent.) which contains 0·260 gram calcium oxide. On treating the solution as described I obtained the following results:

I had taken in this case 2 grams \( \text{KHSO}_4 \).

\[
\begin{array}{l}
\text{HCN liberated by 2 grams \( \text{KHSO}_4 \)} = 0.3970 \text{ gram} \\
\text{HCN liberated in experiment} = 0.1509 \text{ , , ,} \\
\end{array}
\]

\[ C = 0.2461 \times 1.037 = 0.2552 \text{ gram} \]

Calcium oxide found 0·255 \( , , , \)

"" taken 0·260 ""

or in percentages 0·051 and 0·052 respectively or only \( \frac{1}{200} \) of a per cent. from the correct amount, or within 2 per cent. of the lime taken. Of course the operation requires care and practice.
GRAM TABLE.

FOR THE ASSAY OF CYANIDE SOLUTIONS.

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<th>If ½-pint of Solution gives Fine Metal;</th>
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GRAIN TABLE.

FOR THE ASSAY OF CYANIDE SOLUTIONS.

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TABLE FOR 8 OZS. = \frac{1}{4} LB. OF SOLUTION.

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<th>If \frac{1}{4} lb. of Solution gives of fine metal grains.</th>
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<tr>
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<td>0 12 23</td>
<td>\phantom{0}5.000</td>
<td>13 16 8</td>
</tr>
</tbody>
</table>

SOME USEFUL CONSTANTS.

The cubic content of a circular vat in feet = Dia² in feet \times 0.7854 \times \text{depth in feet}.

1 cubic foot of water = 62\frac{1}{4} lbs. nearly.

1 ton of water contains about 36 cubic feet.

1 gallon equals 10 lbs.

1 lb. avoirdupois equals 7000 grains.

To convert lbs. troy into lbs. avoirdupois \times 1.2153.

To convert lbs. avoirdupois into lbs. troy \times 0.82286.
CHAPTER V.

THE APPLIANCES AND PLANT FOR CYANIDE EXTRACTION.

The appliances in use are of much the same nature at all cyanide plants, but their size, shape, and arrangement are subject to endless variations, being chiefly affected by local conditions, the character of the material to be treated, and the individual taste or fancy of the metallurgist. In all cases the designer should utilize the natural advantages at his disposal; and, where possible, the solution vat, leaching vats, vacuum-cylinder, storage tanks, zinc extractors, and sumps should be placed on three separate tiers or platforms, so as to permit of the circulation of the solutions by gravitation, as shown in the following diagram:

![Diagram](image-url)


When a vacuum-cylinder is used, a storage tank must be placed below it to receive the solution when the cylinder becomes full. The bottom of the storage tank must be above the extractor.

Whatever arrangement is adopted, the solution has to be pumped from the sumps, either to the solution tank B, or directly
to the leaching vat. In the latter case the cyanide solution is made up to the working strength in the sump, the dissolving tank A being placed on the floor level; or the strength of the sump solution having been ascertained, the required volume is pumped up to the leaching tank and brought up to working strength by dissolving the requisite quantity of cyanide, placed, in a perforated box or tray, at the discharge end of the solution pipe.

Where the height necessary for gravitation cannot be obtained, the need of constructing high platforms, with a correspondingly high building, is obviated by making up the working solution in one of the sumps, with each of which the dissolving tank and solution sump is connected. To do this without inconvenience an extra sump should be provided.

This arrangement is shown in the following diagram:

![Diagram](image)

**Fig. 2.**—A—Dissolving Tank. B—Sump, also used as Solution Tank. C—Leaching Vat. D—Extractor.

The necessary appliances for a successful and well-equipped cyanide plant, where the ore is to be treated by percolation, are as follows:

1. A dissolving tank.
2. Solution vats.
3. Leaching or percolating vats.
4. Vacuum-cylinder and air-pumps if percolation slows down.
5. Storage vats.
6. Zinc precipitation boxes.
7. Sumps.
8. Solution pump.
9. Fresh-water pipes connected with every tank and vat.
10. Line of pipes connecting vacuum-pump and cylinder with leaching vats.
11. Line of pipes connecting solution pump with each sump, leaching vat, and solution tank.
12. Lines of weak and strong solution pipes connecting leaching vats with extractors.
13. Filter vat for washing and drying gold precipitates.
14. Melting and roasting furnace.
15. Assay office and laboratory.

In the case of double treatment, or intermediate filling, additional vats are required. They are generally placed over, or at a higher elevation than the leaching vats.

**Dissolving Tank.**—This is constructed of wood, iron, or steel. When made of wood, the staves are of 2 in. or 2 1/2 in. pine; and when of iron or steel, the plates are from 1/4 in. to 1/2 in. thick, according to the size of the vat, and stiffened with angle-iron and hoops. The size varies from 3 ft. to 6 ft. in diameter, and from 2 1/2 ft. to 4 ft. in depth.

In large cyanide works a perforated tray, to hold the solid cyanide salt, is suspended over the tank by a chain or steel-wire rope, running over a pulley fixed to a beam overhead. The end of the rope or chain passes over a second pulley on the same beam, fixed at a sufficient distance to permit a balance-weight on the end of the rope to clear the side of the tank.

In practice the solid salt is taken out of the original packing case and cleaned, by removing all adhering particles of sawdust or other packing material with a husk-broom. It is then broken into small pieces and placed in the perforated tray, which is allowed to subside into the solution. The rapid dissolution of the cyanide salt can be effected by imparting motion to the tray by pulling at the weighted end of the rope.

The discharge-hole from the dissolving tank should always be placed three or four inches above the bottom, so as to allow a settling space for the impurities contained in the cyanide.

The impurities contained in commercial cyanide consist principally of black carbide of iron, and other insoluble matters, which would, if permitted, obstruct the solution pipes, choke up the filter webs, and thus cause vexatious delays. Besides this, carbide of iron decomposes the potassium cyanide solution of gold; hence its presence in the solutions would tend to cause a loss by precipitating a portion of the gold.

**Solution Vats.**—These are used for making up the cyanide solutions to the working strength. They are open circular tanks from 14 ft. to 20 ft. in diameter, and from 4 ft. to 14 ft. in depth. They are generally constructed of well-seasoned pine. The sides are made of 5 in. or 6 in. planks, 2 1/2 in. or 3 in. thick.

The bottom is constructed of planks 12 in. by 3 in., bolted and dowelled together independently of the sides. The bolts are placed about 3 ft. apart, and are made of 3/4 in. or 1 in. round
iron. The washer and nut are well countersunk into the plank, and after being tightened, the whole is plugged up with a block of wood to prevent leakage around the bolt. The dowels consist of \( \frac{3}{4} \) in. round iron, and are 6 in. or 8 in. long. They are placed about 12 in. apart where the planks come near the circumference.

The bottom is rebated into the side planks, which are kept tight by round iron hoops, \( \frac{3}{4} \) in. to 1\( \frac{1}{2} \) in. in diameter, having three or more cast-iron turn-buckles on each hoop. One, and sometimes two, rings of \( \frac{1}{4} \) in. round india-rubber are placed round the bottom before the staves are put on. The rubber rings are placed in grooves which are made with a steel tool.

The side planks, or staves, are kept in their places by the pressure of the hoops alone, which are generally placed from 15 in. to 18 in. apart, with an extra hoop at the bottom. The hoops are placed, on very large deep tanks, only six or seven inches apart at the bottom, where the pressure is greatest, the distance apart gradually increasing to 18 in. or 20 in. at the top. The extra hoop is placed as close to the bottom hoop as the turn-buckles will permit.

One or more solution vats may be required according to the size of the plant.

**Leaching or Percolating Vats.**—These are made of many different shapes, sizes, and kinds of material. At first, small square tanks of wood were used, but the difficulty of keeping these tight led to the adoption of circular vats, which are also stronger.

In Australia, the favourite material is wood, but in South Africa and New Zealand, steel vats are at present preferred to wooden ones, in all the recently erected cyanide plants. In America, both steel and iron vats are preferred to wooden ones.

In Victoria and New South Wales, leaching vats made of ordinary corrugated iron with wooden bottoms have been used with very satisfactory results. Mr. W. Eddowes, who used them in Victoria, informed the author that they were light, strong, cheap and durable. For vats of large capacity, corrugated iron of No. 16 gauge is employed. The cost is said to be half that of steel vats of the same capacity. When the value of corrugated iron becomes better known, it will doubtless be more largely employed, more especially in outlying mining camps where transport costs are heavy.

The construction of the circular wooden leaching vats is in every respect the same as that of the solution vats already described, differing only in being provided with discharge-doors. The first vats in use at the cyanide works of the Waihi Gold and Silver Mining Company were 22\( \frac{1}{2} \) ft. in diameter, and 4 ft.
The appliances and plant for cyanide extraction. 65

deep. The sides were built of 5 in. kauri planks, 3 in. thick, bound together with five hoops of round iron, three of which were 7⁄8 in. in diameter, and two 1 in., having three turn-buckles on each hoop. Five inches is taken off the depth by the false bottom, filter-frame, and cloth. At the Company's new mill, there are ten concrete tanks in two rows, each tank being 50 ft. by 40 ft. and 4 ft. deep.

The foundations or supports of wooden vats, 20 ft. in diameter, with 7 ft. staves, to hold a charge of 70 tons of tailings, designed by the author for the Moanataiari Gold Mining Company, Thames, consisted of mudsills, 15 in. by 6 in., laid flat, on which rest the sole pieces, 10 in. by 8 in., supporting the props or uprights, 8 in. by 8 in., on which are laid the bearers of the same dimensions as the sole pieces. There are five rows of props—two rows of three, three rows of five—twenty-one in number altogether, spaced with 4 ft. centres both ways. The bottom of the vat rests on the bearers, which are cut so as to allow a space 4 in. wide between their ends and the side planks. This space is left to detect and repair any leaks that may take place round the sides of the vat.

Where the solid rock is exposed to build upon, the mudsills can be dispensed with. (For detailed drawings, see Plate I.)

The same support can be used for iron or steel vats, but it is advisable to place 9 in. by 3 in. planks across the bearers, so as to more evenly distribute the weight of the vat, the bottom of which rests on the planks.

At the Simmer and Jack Cyanide Works, Johannesburg, the leaching tanks are constructed of pine. They are 42 ft. in diameter, and 14 ft. deep; bound together with fifteen hoops of round iron. They rest on piers of solid masonry. The staves and bottoms are made of 9 in. by 3 in. material.

At the "Main Reef" Works there are six leaching vats, each 26 ft. inside diameter, with 8 ft. staves, and holding 135 tons of tailings.

The staves are 4 1⁄2 in. wide and 3 in. thick, and planed to the level by machines, and afterwards hand-dressed on the abutting edges. They are checked 2 in. to fit on the bottom, with a chime 6 in. below the check.

The bottom of the vat is made of 9 in. by 3 in. deals, planed by machine and grooved 3⁄4 in. by 1⁄2 in. by a saw, and is also hand-dressed on the edges. Clear-pine tongues, 1 in. by 3⁄8 in., fill the grooves. The joists across the tunnel, below the vat, consist of 9 in. by 3 in. deals, bolted together in pairs, and laid 2 ft. 3 in. apart from centre to centre.

These joists are first laid in position, then the bottom of the vat
is laid down, cramped up, and the circle struck out. The bottom
is now sawn to the circle, and when bevelled all round is ready
for the staves, which are driven up as tightly as possible.
Six hoops of round iron are used to keep the staves in their
places; the top pair 1 in. in diameter, the middle pair 1\(\frac{1}{2}\) in., and
the lowest pair 1\(\frac{3}{8}\) in. in diameter, with screwed ends.
Each hoop is made in three sections, rolled to the required
curve, and connected by cast-iron turn-buckles.

![Diagram of Turn-Buckle]

**Fig. 3.—Showing Construction of Turn-Buckle.**
Scale: \(\frac{1}{4}\) in. = 1 ft.

The screwed ends pass through the turn-buckles, and while
each hoop is being drawn up it is hammered with a heavy sledge-
hammer.

Two carpenters, practised at the work, can dress the material
for a vat, 28 ft. in diameter and 8 ft. deep, in about a week, and
erect it in about four days.

With large vats, constructed of brick and cement, in an exca-
vation in the ground, there is no means of ascertaining what leak-
age is going on; and, in a process in which gold solutions are
being dealt with, an exceedingly small leak, in the course of the
year, would represent a considerable loss. For this reason their
construction cannot be recommended where material is procurable
for the construction of wooden or steel vats.

At the Langlaagte Estate Company’s Cyanide Works, S.A., the
tanks are round and constructed of brick, faced with hydraulic
cement. Their size is 40 ft. in diameter and 10 ft. deep. At the
Crown Reef Works the tanks are also of brick, 40 ft. square and
10 ft. deep. At the Waihi-Silverton Works, N.Z., the tanks are
constructed of \(\frac{3}{8}\) in. steel, being 16 ft. in diameter and 4 ft. deep.
At the Moanataiari mine, the steel vats are 20 ft. in diameter and
7 ft. deep. At the Cripple Creek Gold Exploration Company’s
Works, in Colorado, the vats are made of iron, being 20 ft. in
diameter.

For the direct treatment of dry-crushed ore, the leaching vats
are seldom over 4 ft. deep, on account of the difficulty of percola-
tion with a greater depth of ore; but, with tailings comparatively
free from slimes the depth varies from 8 ft. to 14 ft.
The leaching vats, on account of the enormous weight they hold, must be built on strong firm foundations, so as to prevent settling, and the leakage which would be sure to follow. In South Africa they are often built on piers of stone; and in New Zealand and Australia, where timber is plentiful, on massive frames of wood. Whatever the foundations, there should always be free access to the bottom of the tanks, so as to be able to detect and repair leaks.

Steel and iron vats are now coming into general use. They possess many advantages over wooden ones. They are generally coated with a composition consisting of a mixture of coal-tar, pitch, and kauri gum. (See Plate II.)

The filter-frame in steel vats is supported on a ring of iron riveted to the side about 3 in. from the bottom. The filter webbing is laid on the frame and kept in its place by means of a ring of angle-iron, which is constructed in four, six or eight pieces, or lengths, so as to be easily handled. The ordinary method of grouting the cloth between the ring of iron and the side of the vat with a small rope is the best.

In large steel vats, the ring for the reception of the filter-cloth is omitted, and in this case the filter-frame is constructed so as to leave an annular space an inch wide all round the vat to permit the filter-cloth to be grouted between the frame and the vat.

**Solution Pipes.**—Each tank is provided with a separate drain-pipe, 1½ in. or 2 in. in diameter, with two stop-cocks near each other, one over the strong solution launder or pipe, the other over the weak solution launder, leading to their respective zinc extractors. When filtration is assisted by a vacuum, a third stop-cock is provided for the air-pipe.

The sizes of the pipes for charging the vats with the solutions are as follows:

<table>
<thead>
<tr>
<th>Vats, 20–24 ft. in diameter,</th>
<th>2½ in.</th>
</tr>
</thead>
<tbody>
<tr>
<td>24–32 ft.</td>
<td>3 in.</td>
</tr>
<tr>
<td>32–40 ft.</td>
<td>4 in.</td>
</tr>
</tbody>
</table>

Two or three lines of pipe with two mains, one for the weak and one for the strong; and with three mains, one for the weak, medium and strong respectively, running parallel with the line of leaching vats, afford the simplest, most economical, and effective method of collecting the solutions as they percolate from the vats. This system enables the solutions from each vat to be tested separately and readily, and by this means any mishaps can at once be detected.

Instead of having stop-cocks on the end of the drain-pipe, from each vat, a short length of rubber hose is sometimes fixed on the
end; and by moving the hose the solution can be drained into the strong or weak launder as required.

The solution pipes from the leaching vats are led to the extractor house or room where the solutions are run into the extractors. It is a good practice to fix a short length of rubber hose on the end of each pipe so as to be able to conduct the solution to either the strong, medium or weak extractor, as required.

From the end of the extractor boxes the barren solution is led into a sump. From the sumps the solutions are either pumped direct into the leaching vats, or into the solution vats, whence they can be led into the leaching vats as required. The arrangement of the pipe lines is shown in Plate III.

**Filter-Frames.**—The old filter-beds of gravel and sand have been entirely superseded by light wooden frames, over which are placed filter-cloths or webs, consisting of either extra strong Hessian, loose canvas, cocoa-matting, or burlap. For the filtration of slimes, or dry-crushed ores, which always contain a large percentage of very fine sands, a webbing of strong Hessian or canvas is used; and for tailings or concentrates a webbing of cocoa-matting or burlap.

In some plants a duck-cloth filter is laid over cocoa-matting. The duck-cloth for a 100 ton vat costs about £2, 10s. ready made.

At the Waihi Cyanide Works the filter-frames, designed by the general manager, Mr. H. P. Barry, consist of narrow laths placed parallel, and about an inch apart. On these laths are nailed, transversely, narrow moulding-like laths, also about an inch apart. An open frame-work or grating is thus obtained, having openings an inch square.

At the Main Reef Cyanide Works, Johannesburg, the filter frame is made of 3 in. by 1 in. slats, placed on edge, 6 in. apart, their ends being kept 1 in. from the sides of the vat. Strips of wood, 1 in. square, are nailed on the top of the slats 1 in. apart to form a support for the cocoa-matting.

The filter-frames for large vats are constructed in sections. The sections, when fitted together, form a circular frame about an inch less in diameter than that of the vat. This leaves an annular space between the frame and the vat, which permits the filter-cloth to be firmly grouted in its place by means of a small rope passing round the circumference of the vat.

The author has used such filter-frames for over five years, and finds that they possess many advantages over the old gravel-filters.

At plants where intermediate filling is adopted, the intermediate vats are provided with automatic distributors.
water, and often a want of fall for the sludge, the "bottom discharge" is largely practised, the residues being shovelled through a hole in the bottom of the vat into a truck immediately below. At the Barret Company's Works the tailings are shovelled into a launder below the vat, and a stream of water carries them away.

At the Crown, Woodstock and Talisman Cyanide Works in New Zealand, where the ore contains a percentage of coarse gold, the residues are slowly sluiced over extensive amalgamated copper plates placed immediately below the discharge holes.

**Fig. 4.—Butters' Bottom-Discharge Door. Scale: 1 in. = 1 ft.**

At the Langlaagte Company's Cyanide Works, near Johannesburg, the residues are discharged from the large brick leaching tanks by means of steam travelling cranes, which lower the bodies of the empty trucks into the tanks, where they are filled by Kaffir labour. When filled the trucks are raised and placed on their carriages, to be wheeled away to the dump.

**Discharge Doors.**—When side-discharge by sluicing is used, one or two outlets are generally provided for each vat; but in the case of bottom-discharge there are two, four, six, or eight discharge openings to each vat, according to its size.

At the Witwatersrand Goldfields the bottom-discharge is
employed for discharging the round wooden leaching vats. When filling a deep tank with tailings, a length of wrought-iron pipe or cylinder, three or four feet long, is placed over each discharge-hole, and then the tailings are dumped in. The pipe raises the outlet within a few feet of the surface, and thus facilitates the discharge.

On these fields, Butters' bottom-discharge doors are largely used. Fig. 4 shows their construction.

The arrangement is very simple and effective. On the bottom side of the tank, a cast-iron ring, A, is bolted to the cast-iron cylinder, B, inside the tank. Inside the cylinder is a projecting lug, C, upon which rests the hanger, D, which forms part of the screw, E. The cast-iron cover, F, when placed in position, is fastened by the butterfly-nut, G, and by screwing this firmly the whole arrangement becomes watertight. The faces of the ring

![Diagram](image)

**Fig. 5.—Irvine's Bottom-Discharge Door.**

A—Recess for Packing. Scale: \(\frac{1}{4}\) in. = 4 in.

and cover should be planed perfectly even, so as to make a good joint. The joint is also made tight by a luting of clay.

Fig. 5 shows another bottom-discharge door, designed by Mr. W. F. Irvine. It is simple in construction and not likely to get out of order, but would be more convenient in shallow than deep vats.

Plate IV. represents the working drawing of an improved bottom-discharge door, simple in construction, very efficient, and easy to handle. It was originally designed by Mr. G. R. Walker, improved by Mr. A. Price, and perfected and used in its present form by the author in the Moanataiari and other cyanide works. The Koppel patent tank door recently placed in the market differs from the above only in that the pressure to close the door is applied by a screw at the side instead of at the centre.

The door is swung on a loose hinge, and faced with a wide ring
of rubber-insertion. It is closed by lifting, or swinging the door with the left hand over the discharge hole, and then turning an iron button with the right hand so as to carry the weight of the door, which is now approximately in position. The loose dog is then inserted in the lugs and the door screwed up tightly with both hands. The iron-button turns easily on a small stud-bolt, and serves to keep the door in position preparatory to the final tightening.

To open the door the dog is unscrewed, and the button turned to one side, when the door falls open by its own weight, swinging away from the workman. The operation of opening or closing occupies two or three minutes.

The drawing shows a door designed for a steel vat, but it could easily be adapted for a wooden vat by lengthening the cylindrical part, which projects into the vat the necessary length to make it level with the filter-frame, less, of course, the thickness of the coir matting or filter-cloth.

Plate XII. in Chapter XI. shows the bottom discharge-door designed by Mr. Roche for the Waihi Company's new cyanide plant at Waikino.

Fig. 6 shows a modification of a side-discharge door for wooden vats, designed by Mr. W. R. Feldtmann, of Johannesburg.

Another method of side-discharge, designed by Mr. Irvine, is in use at the Crown Reef Company's fine cyanide works, where the large square brick and cement vats are provided with doors which permit of the ingress of the discharging trucks.* The door frames

* Feldtmann, Notes on Gold Extraction, 1894, p. 3.
are bolted to the cement walls, and the plate-iron doors are drawn tight against these by means of an ingenious arrangement of sliding lugs, bolts, and nuts.

The side-discharge sludge doors used in New Zealand and Australia are of the simplest construction and yet perfectly efficient. They consist of a cast-iron frame with two projecting lugs, one on each side, and a projecting bar running along the top side. The lug on the right side is placed with the notch upwards, that on the left with the notch in the reverse position.

The opening is closed by a cast-iron door, which is kept in position by the pressure of a screw acting through a loose iron dog, the ends of which fit into the lugs so as to obtain the necessary leverage.

The door is suspended in front of the opening, preparatory to fixing up, by a hook of bent round iron, which is supported on the projecting bar on the frame. It is rendered watertight by a facing of rubber insertion, fixed on with tar, or by a luting of clay.

These doors seldom give any trouble. They are easily opened or closed by a few turns of the screw. The different parts are shown to scale on Plate V.

Steel leaching tanks with bottom-discharge, designed by the author for the Moanataiari Cyanide Plant, are shown on Plate II.

**Sumps.**—There are at least two of these in every cyanide plant, to receive the cyanide solutions after passing through the zinc-extractor, one for the strong solution and one for the weak. In plants dealing with acid ores or tailings there is often an additional tank besides these for storing the alkaline wash-solutions.

The size of the storage sumps depends on the size of the plant. In most plants they are the same size as the leaching vats. They are constructed either of steel, concrete, bricks faced with cement, or wood. The latter is the favourite material. The construction of wooden sumps is the same as that of the storage tanks or vats.

In many plants the sumps are placed below the level of the lower or extractor floor of the cyanide building, and in such cases they are decked over with planks, having a man-hole for repairs or cleaning. The depth of the solution is indicated by the tell-tale.

In some tailings plants the working cyanide solution is made up in strong solution sumps, thus saving the construction of a storage vat, but requiring an extra sump. This method could not be used with advantage for the treatment of dry-crushed ores in which it is necessary to apply the strong solution slowly from below, so as to prevent the formation of lumps and channels in the pulp such as would occur were the solution turned on to the dry ore from above.
Blaisdell Vat Excavator.—This machine has been designed for discharging the sands from the leaching vat through a central discharge door. The sands fall on to a belt-conveyor, by means of which they are elevated and discharged into any desired vat in a series.

The machine consists of a number of rotating steel discs, attached to arms which are rotated from a central spindle by means of an electric motor. The discs are similar to those in an agricultural disc cultivator. They are set at an angle with the arm, and when the arm revolves they push the sand towards the centre of the vat, where it falls through the discharge opening. To prevent the sand heaping up towards the centre there are two short projections carrying arms set at right angles to the long ones.

This machine requires about seven horse-power to operate it. It is claimed to have a capacity of 100 tons per working hour in a 30-feet vat.

The Blaisdell excavator, like the Butters distributor, is carried on a track that runs the whole length of a series of vats.

This machine, as well as a centrifugal distributor designed by the same inventor, are in use at the tailings plant of Chas. Butters & Co., Ltd., Virginia City, Nevada.

Zinc Extractors.—There are at least two of these in every plant, one for the strong solutions and one for the weak. In works treating sands and slimes it is usual to provide extractors for the strong, medium, and weak solutions respectively. The number of extractors will depend on the capacity of the plant.

Extractors are commonly constructed of wood, but in America boxes made of sheet-iron and steel, either painted or enamelled on the inside, are used in many cyanide plants. They are generally made in the form of long rectangular boxes. Circular vats have been used in New Zealand, South Africa, and elsewhere for some years, but they have not come into common use.

The extractors made of wood are divided into a number of compartments, generally from five to ten, by means of partitions and baffle-boards fixed in such a way that the solution finds its way upwards through the zinc-turnings in each compartment.

The first and last compartments are filled with sand or tow to act as filters, the upper one to remove any clayey matter in the solutions, the lower one to prevent the escape of slimes.

The remaining divisions are sometimes provided with removable shallow trays of wood, with convenient handles, often of bent round iron, to enable them to be easily lifted up when necessary. The trays support, at the bottom, a wire screen of ¾ in. mesh, which permits the free circulation of the solutions, and enables
the gold and zinc slimes, as they form, to fall through into the bottom of the compartment.

Instead of projecting handles, the trays are often provided with an iron plate on two sides, with two holes in it to take the ends of hooks for lifting the trays and their contained zinc.

The extractors are cleaned-up from plug-holes on the side, one to each compartment. To facilitate the "clean-up" the bottoms of the compartments slope to the side, and also to the lower side-corner, where the discharge-hole is situated.

A launder of wood or iron is fixed on the discharge-side, immediately under the plug-holes, so as to receive the slimes when they are washed out of the boxes. The top of the extractor, as well as the side launder, is protected with a lid or close-fitting grating of wood or iron, provided with locks, to prevent the contents of the extractor being tampered with.

To facilitate the withdrawal of the wooden plugs from the discharge-holes, short lengths of rubber hose are fixed in the holes. The rubber is yielding, and, while rendering the holes watertight, enables the plugs to be withdrawn without the force which is generally necessary when they are driven into the bar-holes.

The extractor should be constructed of well-seasoned pine, with sides of 1½ in. or 2 in. boards, and the divisions of 1 in. boards, well dressed. The size will depend upon the capacity of the plant, and will vary from 12 ft. to 20 ft. in length, from 2 ft. to 3 ft. in depth, and from 15 in. to 45 in. in width. The extractors mostly used are shown on Plates VI. and VII.

It is now well known that the bulk of the precipitation takes place in the first two or three boxes, and in cyanide plants handling large volumes of cyanide solution daily, the tendency has been of late to reduce the number of divisions in the extractor to six or eight, and to increase their dimensions. In this case the extractor is simply a long box or trough divided by baffle-boards into compartments, without side plugs or side launder. At the periodical "clean-up" each compartment is cleaned separately; or sometimes two or more, often all the compartments, are made to communicate with each other by plugs at the bottom of baffle-boards so that the slimes can be washed into the lower one of the set thus communicating, and thence be baled out into a tub or run into a bucket. (See Plate VII.)

The construction of the extractor permits an endless variety of design according to the individual fancy of the engineer, always bearing in mind that the object to be attained is to compel the solutions to pass upward through the zinc at a uniform and slow rate.

At the Waihi Company's large mill at Waikino, the solutions
are collected in a rectangular tank divided into three compartments, each forming a coir-filter or clarifying box, for the strong, medium, and weak solutions respectively.

Each compartment is provided with a false-bottom, 9 in. deep, covered with cocoanut matting, which is kept in place by a light wooden frame.

From the clarifying boxes or compartments, the solutions run into the different extractor boxes, which are made 23 ft. long by 3 ft. wide by 2 ft. 6 in. deep. The extractors have flat bottoms and are divided into nine compartments, namely, seven for precipitation and two for settling, one at each end, which is filled with tow.

The trays are 6 in. deep and covered with \( \frac{1}{3} \) in. mesh screens. The slimes are washed into the end compartment through plugged holes in the baffle-boards.

The solutions leave each box by a 3 in. pipe connecting with the main leading to the sumps. The cement floor is constructed so as to convey all drainage from the extractors to a small well.

The circular vats in use at the Bau mine in Upper Sarawak in Borneo are 8 ft. in diameter and 4 ft. 6 in. deep. They are provided with a false-bottom, under which the solution is led by a vertical pipe in the centre of the vat. On the top of the false-bottom there is a sieve on which the zinc shavings are laid. The capacity of these extractor vats is about 10 tons of solution per hour. The extractions are said to be perfectly satisfactory. The vats require little or no attention, the fresh shavings being added to the top as required.

The sheet-iron and steel extractors used in America are made in the form of long rectangular boxes, divided, like the wooden boxes, into a series of wide and narrow compartments. They are enamelled with porcelain, or painted with a paint which is not affected by cyanide or alkaline solutions. Extreme care has to be exercised in the construction of these boxes to prevent the zinc coming in contact with the iron. A zinc-iron couple would waste zinc, and cause gold to be deposited on the iron.

Extractors made of sheet-steel painted or enamelled, or of cast-iron enamelled, are now supplied in San Francisco and London, made in separate sections divided into the usual wide and narrow compartments. The sections are placed in a series on descending shallow steps. They are made either rectangular or circular.

**The Roasting Furnace.**—In Australia and New Zealand in small cyanide plants this consists of a shallow cast-iron pan or plate, built over a small furnace. A funnel-shaped hood of light wrought-iron is placed over the furnace to carry off the zinc oxide and other fumes.

The hood is suspended by a chain or steel-wire rope passing
over a pulley overhead, and is balanced by a weight on the end of the chain. The upper or narrow end of the hood telescopes for a few feet into the pipe leading to the condensing flue, and by means of the balance weight the hood can be lowered or raised over the roasting pan or plate as required.

The first part of the condensing flue is a nearly horizontal length of iron pipe, on which a stream of cold water is sometimes allowed to play, so as to condense the mercury vapours and zinc fumes.

In large cyanide works, a built-in muffle-like roasting furnace constructed of fire-brick is commonly used. It possesses the advantages that a better regulation of the heat can be effected, and that with care the loss through dusting is less than with open pans.

A roasting furnace capable of taking 50 lbs. of precipitate at a time is shown in Plate VIII. figs. 1 and 2.

Fig. 3, Plate VIII., shows a special furnace for roasting slimes. It is a combined reverberatory and muffle furnace. There is a fire-box at one end, a cast-iron hearth in the middle, on which the material lies, and an exit-flue at the other end. The fire-bridge is built in two parts, with a passage between them leading to flues below the cast-iron plate, so that part of the heat is deflected and passes under the plate, while the remainder passes over it with a plentiful supply of air, thus effecting the oxidation of nearly all the zinc.

The Filter-Press or Vat.—In large works, where hundreds of pounds of precipitates have to be handled in the course of a week, a filter-vat with a strong close canvas webbing is found very convenient for washing and drying. The false bottom of the vat is connected with the air-pump. In plants where the slimes are acid treated, a small filter-press is used for washing and drying.

Tell-Tales.—These should be provided for the dissolving and solution tanks, and the storage sumps. They should be marked in feet and inches, so that the workmen can draw off a given depth of solution without any calculation. The cubic content of an inch of depth of each vat or tank in pounds should be marked on a board, so that when so many pounds of solution are required, their equivalent in inches of depth can be ascertained at a glance, thus saving delay and error from needless calculations.

Solution Pump.—This is generally a small centrifugal pump, with a 2 in., 3 in. or 4 in. discharge-pipe. It is driven off the shaft which conveys power to the air-pump.

The Air-Lift for Sands, Slimes, Solutions, etc.—On sloping ground the milling plant and cyanide works can generally be so arranged that the sands and slimes find their way to the leaching vats by gravitation. In cases where natural slopes are
not available for mill sites, the battery sands have to be lifted to the leaching vats by artificial means. In South Africa, bucket-wheels; in America, bucket-wheels and sand-pumps; and in Western Australia, bucket-wheels sand-pumps and air-lifts, are commonly employed to lift the tailings.

The air-lift was first introduced and worked on a large scale, some four years ago, by Mr. J. W. Archibald, manager of the Mount Malcolm Proprietary Gold Mines, Limited, at Murrin-Murrin, in Western Australia. Since then it has come into favour in Kalgoorlie, and seems likely in time to supersede both tailings wheels and sand-pumps in Western Australia.

So far the air-lift has not shown a high efficiency for the power employed; but it possesses several special advantages which seem to recommend it to mine owners, namely: cheapness of erection, continuous operation, and little wear and tear, having no complicated parts to get out of order. The Associated Mine has two in operation; Kalgoorlie, six; Boulder No. 1, four; Westralia Mount Morgans, two; Lake View Consols, one; besides many others.

Judging from the success which has attended the use of air-lifts in Western Australia, it is certain that this form of elevator is deserving of more general application, and for that reason I append the following lucid description of their construction, for which I am indebted to the courtesy of Mr. Max von Bernewitz of Kalgoorlie.

He says: "In erecting an air-lift the first work is that of sinking a well-hole the same depth, or better still, 2 ft. more than the height the pulp has to be lifted. The size of the hole depends on the size and number of lifts to be erected: for example, if one 6 in. lift is to be put up, a well-hole not less than 3 ft. in diameter should be sunk. The reason why a hole is sunk is that the lift has always in it a column of water and pulp, equal to height to be lifted, whereby the air meets with the necessary resistance required to start the column in motion.

"Next, a large pipe, say 2 ft. 6 in. in diameter, with a tapered bottom and of the same depth as the well-hole, is lowered into the latter. The delivery pipe or lift is then fitted together for the correct length, and lowered into the large pipe. Finally, the air-pipe is fitted together and run down at the side of the lift-pipe. At the bottom of the air-pipe is a bend which looks up into the lift about 2 ins. It is desirable to have flanged joints on all the pipes, in order that they may be quickly and easily disconnected in case of an accident.

"At the top of the lift-pipe, a bend may be fitted on, but the practice now is to bolt a launder directly on to a flange at the top
of the pipe. By this means the air, as it reaches the top, can escape at once."

It has also been found that the larger the air- and lift-pipes, the better is efficiency attained; also, that air-lifts should be in duplicate in case of mishap.

If the air is taken from a receiver at, say, five atmospheres pressure, a pressure several times greater than actually required by a lift, and expanded before doing its work, it is found that there is a loss of power. But if the air is taken direct from the compressor to the air-lift, a far greater efficiency is obtained, as the rising and falling of the level of the pulp acts as a perfect governor to the compressor. The practice on most of the mines is to take air from a receiver at, say, five atmospheres pressure, and conduct it to a lift through a reducing valve; but this is unsatisfactory, and a separate compressor giving large quantities of air at a low pressure is the more economical method.

Again, when a compressor supplies air for many purposes the working of the lift is greatly interfered with. Take, for example, the mill at the Kalgoorlie Mine. Here a compressor running at a high speed supplies air to actuate two solution pumps; also air to agitate slimes; to force the agitated slimes into filter-presses; to dry the presses and work three air-lifts, of which one lifts pulp 20 feet, one sands 30 feet, and one clear water, 15 feet high. All the above require air at different pressures, and interfere with the working of the air-lifts. It is certain that a small compressor for the lifts alone would be more satisfactory.

On the W. A. goldfields, where fuel, water, and wage costs are high, the working of air-lifts is fairly expensive, but in countries where compressors can be driven by water-power with little or no attendance, the cost should not be high.

In the Kalgoorlie Mill a little trouble was experienced with a 20 ft. lift, which elevates red hot roasted ore previously mixed with dense salt water. This lift elevates 700 tons of pulp per diem with an air pressure of 0·7 atmosphere. The pulp when raised is almost at boiling point, and a hard deposit quickly sets around the pipes, and gradually chokes them up at the bottom. This is due to the deposition of lime and magnesia from the ore.

In this mill a proportion of six parts of water to one part of ore was found necessary. The above lift-pipe was 8 in. in diameter, and air-pipe 1½ in.

In the Hainault Mill, a 25 ft. lift has worked for over twelve months with no stoppages, whereas the sand-pump was always hung up for repairs.

At the Associated and Westralia Mount Morgan’s mills, lifts are in operation elevating thick slimes from settlers to the agitators.
At the latter mill there is a lift which raises pulp from 30 head of stamps, crushing 120 tons per diem. Its dimensions are:

Height of lift-pipe above well, about 20 feet.
Depth, below top of well 22
Total length of lift-pipe 42
Diameter of lift-pipe 6 inches.
Diameter of air-pipe 1½

Air-pressure 0·8 atmosphere.

At Kalgoorlie, air-lifts are employed to circulate cyanide solution from the bottom to the top of leaching vats, and to raise solutions from sumps. At the Boulder No. 1 Mill, two air-lifts are for lifting mercury in connection with the Reicken process, having the following dimensions:

Height of lift-pipe, about 12 feet.
Depth of well 4
Diameter of pipe forming well 3 inches.
Diameter of lift-pipe 1 inch.
Diameter of air-pipe ½

Air-pressure, 3 to 5 atmospheres.

SIZE OF PLANT.

The size of the plant will depend on the intended capacity per month and the condition of the material to be treated. A plant capable of treating 2000 tons of freely-percolating tailings per month would probably not be able to treat more than half that quantity of dry-crushed ore or slimy tailings.

With dry-crushed ores the depth of the charge seldom exceeds 4 ft., while with sharp tailings the depth sometimes reaches 10 ft. or 12 ft. Cyanide plants for the treatment of dry-crushed ore are therefore provided with a large number of shallow vats; and those for tailings with a smaller number of deep vats.

The following gives the number of leaching vats required for the treatment of different quantities of dry-crushed ore and tailings.

**FOR DRY ORE OR SLIMY TAILINGS.—**Size of vat: 22½ ft. diameter, 4 ft. deep; charge 30 tons.

<table>
<thead>
<tr>
<th>Number of Vats</th>
<th>Monthly Capacity</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>350 tons</td>
</tr>
<tr>
<td>4</td>
<td>700 tons</td>
</tr>
<tr>
<td>6</td>
<td>1050 tons</td>
</tr>
<tr>
<td>8</td>
<td>1400 tons</td>
</tr>
<tr>
<td>10</td>
<td>1750 tons</td>
</tr>
<tr>
<td>12</td>
<td>2100 tons</td>
</tr>
</tbody>
</table>
THE APPLIANCES AND PLANT FOR CYANIDE EXTRACTION. 81

For Freely-Percolating Tailings.—Depth of vat, 10 ft.; charge 100 tons.

<table>
<thead>
<tr>
<th>Vats</th>
<th>Capacity</th>
<th>Per Month</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td></td>
<td>1200 tons</td>
</tr>
<tr>
<td>4</td>
<td></td>
<td>2400</td>
</tr>
<tr>
<td>6</td>
<td></td>
<td>3600</td>
</tr>
<tr>
<td>8</td>
<td></td>
<td>4800</td>
</tr>
</tbody>
</table>

In small plants it is advisable to have a spare leaching vat in case of a temporary breakdown in one of the vats in use.

COST OF PLANT.

The cost of the cyanide plant depends largely on the locality, the material used in the construction, and, to a certain extent, on the condition of the material to be treated. For example, a plant to treat 2000 tons of dry-crushed ore per month will cost more than a plant to treat 2000 tons of tailings.

The approximate cost of plants of different capacities in Australia and New Zealand, when wood is the material used in the construction of the vats, foundations, and buildings, is given below:

For Dry-crushed Ore—By Percolation.

<table>
<thead>
<tr>
<th>Monthly Capacity</th>
<th>Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>350 tons</td>
<td>£1200</td>
</tr>
<tr>
<td>700</td>
<td>1500</td>
</tr>
<tr>
<td>1050</td>
<td>1850</td>
</tr>
<tr>
<td>1400</td>
<td>2000</td>
</tr>
<tr>
<td>1700</td>
<td>2400</td>
</tr>
<tr>
<td>2000</td>
<td>2900</td>
</tr>
</tbody>
</table>

For Tailings—By Percolation.

<table>
<thead>
<tr>
<th>Monthly Capacity</th>
<th>Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>1200 tons</td>
<td>£2000</td>
</tr>
<tr>
<td>2400</td>
<td>2800</td>
</tr>
<tr>
<td>3600</td>
<td>3500</td>
</tr>
<tr>
<td>4800</td>
<td>4000</td>
</tr>
</tbody>
</table>

The above estimates include the cost of the laboratory, assay, and melting furnaces, and all the appliances for a successful cyanide plant, but does not include the cost of a bucket tailings wheel or sand-pump for elevating sands, etc.

The cost of steel or iron vats is about the same as that of wood. The steel leaching vats at the Waihi-Silverton, 16 ft. diameter
and 4 ft. deep, with central discharge, cost £56 each, and the wood foundations about £10 each.

At Johannesburg the cost of a cyanide plant is about 25s. per ton of tailings to be treated per month. Thus a plant to treat 300 tons of tailings per month would cost about £4000, not including the cost of tailings wheel or assay and smelting offices. For larger plants the cost is smaller in proportion. The Rand cyanide plants, it should be noted, are not roofed over as they are in many other countries.

In America steal leaching vats cost about $3\frac{1}{2}$ cents per lb. free on board cars. Thus a 100 ton vat weighing 9400 lbs. would cost about £67. For red wood vats the cost is as follows:

<table>
<thead>
<tr>
<th>100 tons</th>
<th>200</th>
<th>500</th>
</tr>
</thead>
<tbody>
<tr>
<td>£42 F.O.B.</td>
<td>80</td>
<td>120</td>
</tr>
</tbody>
</table>

The American red wood vats are stated to give every satisfaction. They are generally coated with P. and B. Paraffin paint, of which 40 gals., costing £8, is sufficient to coat a 500 ton vat twice (that is the inside only), two coats on the sides and three on the bottom.

Where classifying, sliming by pans or tube-mills, treatment of slimes by agitation, filter-pressing, or decanting is undertaken, the cost of construction may be increased from three to ten fold according to the capacity of the plant, character of ore, cost of labour, materials, transport, etc.
CHAPTER VI.

THE ACTUAL EXTRACTION BY CYANIDE.

Synopsis of the Process.—No hard and fast rules can be followed in the working of the cyanide process, as modifications have to be introduced to meet the special requirements of the different classes of ore; but the essential features of the process, whether for the treatment of ores or tailings, are practically the same in all cyanide plants.

The first operation is the filling of the vats. In the direct treatment of dry-crushed ore, the pulverized material is filled in to a depth of three or more feet, according to its fineness. With a uniform granular product, five or even six feet of ore may be filled into the vat, and, in the case of freely-percolating tailings, the vats may be safely charged to a depth of from six to ten feet.

With tailings it is the practice to fill the vat within 20 inches of the top, because the strong solution or preliminary wash-water when applied causes the charge to shrink several inches.

In the case of acid ores or tailings, a preliminary treatment with a caustic alkali, or at least a water-wash, is necessary in order to remove, or neutralize, the mineral salts and acids which decompose the cyanide solutions.

Oxidizing agents such as barium peroxide, potassium ferricyanide, sodium peroxide, and cyanogen bromide, have been tried in America, South Africa, and New Zealand. The results obtained on a working scale with the first and last of these are very satisfactory.

The strong solution, containing generally from 0.2 per cent. to 0.5 per cent. of potassium cyanide, is then applied. The proper strength of solution must be determined experimentally, and with most ores it will be found advisable to use a larger bulk of solution rather than to unduly increase the strength. Silver ores require a larger bulk of solution than gold ores in order to obtain a fair extraction.
The cyanide solution will drain off weak at first, due partly to dilution with the displaced water and partly to the dissolution of the gold and silver, and contact with cyanicides.

The operator will also discover that there is a loss in cyanide every time a solution comes in contact with the ore. For this reason the values ought to be dissolved in the shortest time possible, so as to reduce the consumption of cyanide.

In the case of a dry-crushed ore the solution is allowed to permeate the mass from below, but in the case of tailings it is allowed to flow in over the surface, and soak downwards. The filling of the strong solution generally takes from two to four hours. With tailings it is allowed to stand in contact for twelve hours, after which percolation is begun. With dry-crushed ores slow artificial percolation is commenced at once.

The leaching with the strong solution takes from twenty-four to forty-eight or more hours, according to the character of the bullion; but in the case of ores containing a proportion of fairly coarse gold, it is customary to make the strong solution up to the working strength and pass it through the charge till an adequate extraction is obtained.

The weak cyanide solution, often called the first wash, is then applied. It is pumped from the strong sump, and generally varies in strength from 0·1 per cent. to 0·25 per cent. of potassium cyanide. It is allowed to percolate as rapidly as possible, the filtration being facilitated, when necessary, by means of an artificial vacuum, which may be created by a steam exhaust, or an air-pump connected with a vacuum-cylinder.

The first weak cyanide wash is succeeded by two or three washes of solution from the weak sump, containing from 0·02 per cent. to 0·1 per cent. of K\textsuperscript{1}Cy.

A final washing of clean water is then applied, which serves to displace the preceding weak cyanide wash. By this means the quantity of cyanide solution in circulation remains much about the same.

In the treatment of pyritic ores it is sometimes advantageous to apply a weak cyanide solution before the strong solution. By this means a larger extraction and a saving of cyanide is effected.

The quantity of the strong solution used is about one-third of the weight of the ore. The cyanide and water-washes are each about one-sixth of the weight of the ore.

During the treatment of slimy tailings it is found advantageous to sometimes turn the material over by hand-labour, and thereby effect a more complete washing out of the cyanide solutions from the slimes, which always have a tendency to entangle and retain
them or even to transfer it from one vat to another and subject to a second leaching.

Discrepancies between the theoretical and actual extractions are often a source of much annoyance and perplexity to the metallurgist, but in most cases they will be found to be due either to imperfect sampling or incorrect tonnage. At most mines a ton weighs considerably over 2240 lbs.

At the Witwatersrand Goldfields, in South Africa, the tailings are sometimes subjected to a double treatment—i.e., after the first cyanide treatment the residues are charged into other leaching vats, and again treated with cyanide. A higher extraction will result, but it is doubtful if the difference would cover the cost of the extra handling and cyanide.

The average extraction on the Rand where only high-priced white labour was available, amounts to about 72 per cent., and with double treatment this is raised to 85 per cent. With tailings of fair value the extra recovery is, however, said to leave a margin of profit.

The sequence in which the different operations are applied may be tabulated as follows:

1. Filling the leaching vats.
2. The preliminary treatment with water or alkaline washes if necessary.
3. Leaching with the strong solution containing from 0.3 per cent. to 0.6 per cent. of KCy.
4. First wash, with cyanide from strong sump, containing from 0.1 per cent. to 0.26 per cent. KCy.
5. Second wash, with cyanide from weak sump, containing from 0.02 per cent. to 0.1 per cent. KCy.
6. Third wash, same as second.
7. Fourth wash, same as last.
8. Fifth wash, with clean water.

The gold and silver in the ore are dissolved by the strong solution, and removed or carried out by the first and second washes. The cyanide solutions are allowed to flow through the zinc extractors, the strong solutions through the strong box, and the weak solutions through the weak box. The first two washes, which generally carry most of the dissolved bullion, are conducted through the strong extractor.

FIRST STAGE.

Filling the Vats with Dry-Crushed Ore.—In the case of dry-crushed ores, the charging of the vats is a simple operation, the only disadvantage being the clouds of dust, which seem to be
inseparable from the handling of dry-pulverized ores. At Waihi 90-stamp mill the vats were generally charged by trucks running on to a traveller, provided with hand-traversing gear so as to enable the sand to be tipped in different parts of the vat. In order to prevent packing, the sand was discharged from the trucks on to the platform of a small traveller fixed below the main traveller. By this means the pulverized material was dispersed in a gentle shower over the whole vat.

At some plants the dry pulp is conveyed from the mill to the vats by screw-conveyors or by belt-conveyors, both very efficient methods of transport.

At the Kapai-Vermont Cyanide Works, Kuaotunu, N.Z., the vats were filled directly from the dust-bins, which was situated overhead in an elevated position. By means of a chute, provided with a universal canvas joint, the material was evenly spread over the vats.

When the vat was charged the surface of the ore was made smooth by passing a wide, shallow wooden hoe or rake over it.

This method of filling possesses many advantages over hand-filling by trucks. It is cheaper, more expeditious, and healthier for the workmen, as it raises less dust.

**Filling the Vats with Tailings.**—One of the most serious disadvantages of wet-crushing with the stamp-mill is the production of fine slimes. All ores, even the most silicious, form a proportion of slimes when wet-stamped; and when clayey or earthy matter, country-rock, iron or manganese oxides are associated with the ore, the proportion of slimes is often excessive.

In many places the slimes are very valuable, sometimes even more so than the sands. In leaching processes they prove very refractory, as they seriously interfere with the percolation and washing, thereby causing the extraction to be both costly and imperfect. When the slimes are irregularly distributed through the sands the cyanide solutions form channels through them, and imperfect leaching is the natural result.

At the Witwatersrand Goldfields, where wet-crushing, followed by copper-plate amalgamation, is at present universal, there are two methods in use for dealing with the tailings before treatment with cyanide. The pulp from the copper-plates is lifted by a bucket wheel, run through a lauder, and then classified by *spitzkasten* or *spitzluten* into two products, namely:—

(a.) Sands and slimes, 80 per cent.
(b.) Concentrates containing some sands, 20 per cent.

The concentrates are collected in a storage tank, and kept under water to prevent oxidation, while the sands and slimes pass on to the cyanide vats for treatment, either by what is known as
the "Direct Filling" process or by the "Intermediate Filling" process.

At the extensive works of the Langlaagte Estate Gold Mining Company, the tailings, after leaving the plates, are concentrated, and then run into three large settling dams, each capable of holding 7000 tons. The sands settle in the dam, while the slimes are carried off by the overflow, and allowed to run away. The tailings, now free from slimes, are hauled from the dams in trucks by means of two endless steel-wire ropes, and run up to an overhead tram-line, from which they are dumped into the leaching vats ready for treatment.

**Intermediate Filling.** — At the Robinson, Princess, and many other mines at Johannesburg, the tailings, after being subjected to concentration, are run into intermediate settling-tanks, which are circular in shape, and sufficiently large and numerous for the requirements of the mill. At the Robinson plant, treating 330 tons per day, there are six circular wooden vats, each 24 ft. in diameter, and 11 ft. in depth. This gives a settling surface of 32 square feet per stamp.

From these intermediate settling vats the tailings are distributed to the leaching vats. When the slope of the ground permits it, the settling vats are placed above the leaching vats, the tailings being discharged from holes in the bottom; but when they are below the level of the leaching vats, the tailings are hauled up in trucks actuated by endless wire ropes.

To ensure a fairly even distribution of the sand and slimes in the leaching vats, a simple and ingenious automatic distributor, invented by Messrs. Butters & Mein, is now in use in all parts of the world. It consists of a central casting, with a vertical spindle revolving in a foot-step, and carrying a conical hopper at the top, from which radiate twelve or sixteen iron pipes with bent ends. These pipes vary in size from $1\frac{1}{4}$ in. to $2\frac{1}{2}$ in. in diameter. To their discharge ends are attached flattened nozzles to assist in spreading the tailings over a wide area. A coarse screen is placed over the central hopper, or bowl, so as to prevent stones or pieces of wood, or other débris, from choking the pipes. The distributor is fixed on an iron column in the centre of the vat, and the reaction of the pulp escaping from the bent pipes causes it to slowly revolve in a manner similar to that of an ordinary garden sprinkler. (Fig. 4, Plate VIII., and Plate IX.)

Instead of erecting a distributor in each vat it is now the custom to provide one overhead distributor for each row of vats, suspended by a rod from a trolley running on rails above the centre line of the vats.

A movable launder or pipe feeds the pulp into the hopper, as
near the centre as possible so as to obtain a more even distribution of the sands. The size varies according to the capacity and diameter of the vats.

The usual slope of the arms is 1 in 12; and the diameter of the pipes varies from 1.5 to 2.5 in. The pipe-arms are generally made the same diameter throughout, but at the Horseshoe mill, * South Dakota, the long arms are 3.5 in., medium arms 2.5 to 3 in., and the short arms 2 in. in diameter.

A sketch of the suspended distributor in use at this mill is shown in fig. 7.

The suspended distributor figured by Fulton is similar to those which have been in use in New Zealand for some years. In this

![Diagram of suspended distributor](image)

**FIG. 7.—** Butters and Mein's Sand-Distributor. (After C. H. Fulton.)

class the whole of the weight of the machine comes upon the footstep, and in order to reduce the friction it is customary to provide a ball and roller bearing. The machine should be made as light as possible consistent with strength. It is balanced by fixing the pipes unsymmetrically around the central basin.

In many mills the distributor is placed on a pillar fixed in the centre of the vat, being hoisted up and moved from vat to vat as required.

To ensure success, the collecting vat must be filled with clean water before admitting the pulp, otherwise the slimes would settle with the sand until the overflow of water began. While the machine is running, there must be a continual overflow from

the vat to carry off the fine slimes. The discharge or overflow from the settling vat takes place at all points of the circumference, being received into an annular ring, surrounding the top of the vat, which conveys it to the slime-pit. The overflow of slimes must be continuous until the vat is full of sand; therefore, when the battery stops, a regular quantity of water should still be supplied to the vat.

In some of the mills in South Dakota dry-filling is said to give a more leachable distribution of the sands and slimes than wet-filling.

The vats are provided with filters, and when full of tailings the water is allowed to drain off for fifteen or twenty-four hours, while six hours before discharging, holes are dug down to the discharge doors to let more water flow out. When drained, the tailings are discharged into the leaching vat below, through "bottom-discharge" doors, or into trucks, which are then run into the tank house, where their contents are emptied into the leaching vats. At the same time the required amount of lime (quicklime ground in a ball-mill) is added in such a way that it gets distributed through the whole mass.

The advantages claimed for intermediate filling are:

(a.) By means of the distributor, nearly all the sands are collected in the intermediate tanks, the bulk of the slimes escaping. (The quantity of slimes depends on nature of ore, size of screen, and size of battery-box.)

(b.) When discharging through the bottom, the sands during the operation get thoroughly mixed, thus being in the best condition for treatment.

(c.) Oxidation of pyrites is very slight, so that little cyanide will be consumed.

(d.) A higher extraction can be obtained owing to the presence of the fine sands, from which 85 per cent. of the gold contents can be extracted by cyanide.

(e.) The cost of treatment is very small.

The principal disadvantage is the tendency of the distributor to form deposits of slime, through irregular distribution, at places on, or near, the bottom of the tank, thereby causing some trouble in draining off the water.

**Direct Filling.**—This method is in use at the works of the City and Suburban, Crown Reef, and other companies at the Witwatersrand, and consists in conducting the pulp, as it leaves the copper-plates, into a classifier or spitalute. In this the pulp is divided into two streams: one, the overflow, carrying slimes and fine sands; the other, carrying the coarse sands, together with some fine sand and slimes, which are conveyed by an india-rubber
hose to the leaching vat, where they are distributed by moving the hose over the whole area of the vat. The excess of water is carried off by adjustable gates fitted inside the vat, taking with it some fine sand and slimes.

The advantages claimed for direct filling are:—

1. That pyritic tailings are exposed to the minimum of oxidation.
2. A second handling of the tailings is avoided.
3. A rough preliminary classification is effected, thus separating the fine slimes.

The principal disadvantages of this method are:—

1. The packing of the tailings, which prevents a complete draining of the contained water.
2. The unequal distribution of the sands and slimes, which militates against a perfect extraction by favouring the formation of channels in the mass during the subsequent leaching.

The method of classifying the pulp by a spitzlute, and direct filling, has been in operation at the Great Mercury Cyanide Works, Kuotunu, N.Z., since 1892. When the leaching vat is charged, the water is drawn off through the filter-bed, and after the tailings have drained they are turned over by hand so as to loosen them, and at the same time thoroughly mix any slimes present with the sands.

SECOND STAGE.

The Preliminary Water or Alkaline Wash.—This treatment is only necessary in the case of ores or tailings containing the decomposition products of pyrites, or other base sulphides.

The drying of a pyritic ore in a kiln for dry-crushing and direct treatment generally results in the formation of soluble sulphates, which are destructive to cyanide. At the works of the Woodstock Gold Mining Company, at Karangahake, N.Z., the consumption of cyanide from this cause was very heavy; but the preliminary treatment with a caustic alkali, according to the later report of the management, has effected a great saving of cyanide, accompanied by a higher extraction.

The products of the partial oxidation of iron pyrites in tailings are principally free sulphuric acid, soluble sulphates, and insoluble basic sulphates, all of which cause a large consumption of cyanide, and must, therefore, be removed or neutralized before cyanide treatment is commenced.

If the tailings are very acid, and a considerable proportion of the salts are found to be soluble in water, the general practice is
to apply a preliminary water-wash. In order to neutralize the remaining acid, a quantity of the solution of caustic soda, equal to about 1 lb. of the salt to every ton of ore, is allowed to stand in contact with the tailings for two hours, and then drained off into the alkali-ump.

The necessary quantity of caustic soda can be determined experimentally by the method recommended by Feldtmann. (See Chapter IV.)

At the Witwatersrand Goldfields powdered caustic-lime is generally used in place of soda. With very acid tailings as much as $2\frac{1}{2}$ lbs. to the ton is used. It is applied by adding the requisite amount to each truck-load before being dumped into the leaching vats.

The author generally used lime at the Government Experimental Works at the Thames, and found it preferable to caustic soda, as it is not attended with the production of ferrocyanide of zinc, which fouls the zinc in the extractor. On the other hand, it must be remembered that the addition of lime only improves the extraction when sulphides are not present in the material being treated.

When the cyanide contains sulphide, the extraction is certain to be slow and unsatisfactory. The sprinkling of lead acetate on the top of the tank will precipitate the sulphur of the sulphide as lead sulphide, thus leaving the cyanide free from sulphide.

In the case of very acid tailings, Feldtmann strongly condemns the practice of conducting the preliminary washing in the leaching vat, on account of the possibility of the acid acting on the residual cyanide in the vat, and thereby liberating sufficient hydrocyanic acid to dissolve an appreciable quantity of gold, which he says would be lost, as it is not precipitated by zinc.

He suggests that the washing should be effected in one vat, and the leaching in another; and considers that the extra costs of handling would be more than made up by the higher extraction.

It is doubtful if hydrocyanic acid has any action on gold; but it is probable that the ascending acid by uniting with any alkali present would form an alkaline cyanide which would at once react on any gold to which it had access.

THIRD STAGE.

Strong Solution Leaching.—With dry-crushed ore, the strong solution, about one-third the weight of the charge, and generally containing from 0·2 per cent. to 0·5 per cent. of available potassium cyanide, is allowed to pass into the vat from below. When the solution stands two inches above the surface of the ore
the stop-cock is shut, and the solution lying below the filter is then drawn off, and filtration commenced. The strong solution generally takes from twenty-four to thirty-six hours to percolate through.

In the case of tailings, the strong solution is added on top. After standing twelve hours, to allow the solution to permeate all the lumps of slimy material, the solution is slowly drained off, and passed through the zinc extractor. Small quantities are occasionally drawn off to assist diffusion of the solution through the mass.

It is a very common practice to apply a weak solution from the sump containing, say, 0.05 to 0.10 before applying the strong solution, the intention being to allow the acids and foreign matters destructive to cyanide to expend themselves on the weak solution, before applying the strong solution.

When the ore contains a large proportion of silver—say, from five to eight parts to every part of gold—it will be found necessary to adopt one of two courses in order to obtain a satisfactory extraction.

Either a much greater bulk of potassium cyanide solution must be used to leach the ore—say, a quantity equal in weight to that of the ore taken—or else a much stronger solution must be used.

The former, only, of these two courses would be applicable if the ore contained even a small proportion of copper oxide, carbonate, or sulphide, or of sulphide of antimony, since the solubility of these in all solutions of cyanide, but especially in the strong, would render a fair extraction impossible, besides causing a heavy loss of cyanogen.

The sulphide ores of silver are more slowly soluble in cyanide than gold, and for this reason the treatment of such is always more expensive than that of ordinary tailings or gold-bearing ores. On the other hand, the chloride of silver (kerate or horn silver) is more readily acted on than gold, the extraction generally exceeding 80 per cent.

In the practical treatment of ores and tailings by cyanide, one of the first anomalies to attract the notice of the metallurgist is the fact that the strong solution, while it 'loosens' the gold, so to speak, yet does not carry it away, this being effected by the first and second cyanide washes.

It is found that the first portions of the strong solution, draining from the charge, contain only from 0.02 per cent. to 0.1 per cent. of cyanide; but the remaining portions come off stronger and stronger after a lapse of eight or twelve hours, until, towards the end of the strong leaching, the maximum strength is reached,
after which the strength declines a little before the application of
the first cyanide wash.

The first portions of the solution during the strong cyanide
leaching, being weak in cyanide and low in gold, are, therefore,
passed through the weak zinc extractor, while the later portions,
together with the first and second cyanide washes, are conducted
to the strong extractor.

It will be found in practice that fresh solutions are more active
in dissolving gold than used solutions which in passing through
the zinc-extractors become charged with the inert compound zinc-
potassic-cyanide.

**Strong Sump-Solution Wash.**—After the strong solution
has completely drained off, the strong sump-solution is applied
from above, being run in on the top, or surface, of the ore. Its
strength varies from 0.1 per cent. to 0.25 per cent. of KCy, and
the quantity applied from one-third to one-quarter of the weight
of the ore.

This weak solution is sometimes allowed to slowly percolate
through the charge, but more often it is drawn off as rapidly
as possible, for the more rapidly the wash solutions are drawn
off the more effective is the washing. The filtration is assisted
by opening the stop-cock connecting the bottom of the leaching
vat with the vacuum-cylinder.

The percolation of the weak solution generally takes from twelve
to twenty hours, the time depending on the condition of the pulp
forming the charge.

**Weak Cyanide and Water Washes.**—The number and
strength of these will depend entirely on the character of the ore,
or tailings, being operated on. In some cases it is found necessary
to apply as many as three or four weak cyanide washes, from the
weak sump, and then a final water wash; in other cases the whole
treatment consists of the strong cyanide leaching followed by a
cyanide wash from the strong sump, a cyanide wash from the weak
sump, and one or two final water washes.

The quantity of each wash is in most cases one-half that of
strong cyanide solution.

The effect of the different washes should be carefully deter-
mined by assaying the residues after each washing, and also the
wash solutions as they drain from the vat. By this means the
necessary number of washings will soon be ascertained.

The assay sample should represent a fair average of the charge
in the vat, and is easily and reliably obtained by taking a large
number of cores, the full depth of the charge, by means of a tube
shaped something like a cheese sampler. The scores should be
dried and then sampled down for assay.
Briefly summarized, the aim of the different solutions which have been applied to the material in the leaching vat is as follows:—

(a.) Alkaline wash, to neutralize acidity so as to prevent waste of cyanide.

(b.) Strong solution, to effect solution of gold.

(c.) Weak solutions and water washes, to displace cyanide solution and to prevent cyanide from being thrown away with the residues.

The different operations to be undertaken in the dry-crushing and direct method of treatment may be summarized as follows:—

1. Dry crushing.
3. Copper-plate amalgamation of tailings.
4. Concentration of tailings from plates.
5. Treatment of concentrates.

With wet-crushing the different operations are:—

1. Wet-crushing.
2. Copper-plate amalgamation.
3. Concentration.
4. Treatment of concentrates, by cyanide or other means.
5. Treatment of tailings by cyanide.
6. Treatment of slimes by cyanide.

Remark.—When leaching slimy sands the successive solutions and washes should only be partially drained, just sufficient to allow assay samples to be taken. If very fine sands, or sands containing slimes, be thoroughly drained, they become almost impervious to subsequent solutions. Hence it is important to partially drain so as to leave a cushion or jacket of solution round each particle of sand, the said jacket preventing packing, and thereby permitting its easy replacement by the solution following.

The circulating of the solutions through the charge in the vat is sometimes necessary with obstinate extractions. It can be easiest effected by means of a centrifugal pump, or a small ejector arranged as an air-lift. The latter is in use in New Zealand, and is found to aerate the solutions effectively, besides circulating them.

CRUSHING WITH CYANIDE SOLUTION.

In the Ohinemuri Goldfields of New Zealand, the gold occurs in an extremely fine state, and hitherto dry-crushing and cyaniding have been used, the objection to wet-crushing being the difficulty experienced in treating the slimes which were always high grade.
THE ACTUAL EXTRACTION BY CYANIDE.

The successful experiments conducted at the Crown mines with wet-crushing last year, and the subsequent adoption of this process at the Company's mill, have shown that, with careful manipulation, the simultaneous treatment of the slimes can be carried on with very little extra cost.

The following description of wet-crushing with cyanide solution has a special interest.* The ore is delivered to the rock-breaker without drying. With the ore there is fed into the mortars a constant stream of a stock cyanide solution of about 1 per cent. strength, regulated by stop-cocks near the boxes. The rate of flow is kept between 35 cwt. and 2 tons per hour, being gauged at the supply-tanks. As the stamps, which weigh 1000 lbs., with 96 drops to the minute, crush about 2 tons per stamp per day, the amount of solution is consequently about equal to the ore crushed. Previous experiments showed that a larger proportion of solution, while it increased the output, caused the pulp as well as the gold to be washed through the screens in a state too coarse for cyanidation. From similar considerations the discharge is raised to a height of 8 or 9 inches, the screens being 40-mesh. From the front of the boxes the pulp flows to a wheel-and-bucket elevator, and thence to the vat-shed. The vats are of the size usual in dry-crushing on those goldfields, viz., 22½ ft. in diameter and about 4 ft. in depth, not including the space of 4 inches below the filter-cloth. The pulp flows into the centre of a wooden, two-armed distributor, driven by overhead gearing. Suspended to the arms are five sets of rectangular pieces of sacking, each set being composed of three or four pieces of varying lengths, so that they may be lifted as the vat fills. Their object is to keep the slimes on the surface of the sands from settling. These slimes are being continuously decanted off by an adjustable pipe into a receiving vat on the same level, provided with an agitator similar to that used in an ordinary settler. Thence they are pumped from time to time by a small centrifugal pump into the settling vats, also provided with light agitators. These vats are similar to the sand vats, but have the filter-cloth and frame removed. From these the clear solution is kept continually draining into a sump, whence it is pumped to the head of the precipitating boxes. After passing through the boxes it is pumped into the supply tanks, whence it feeds the stamps or is used for washes.

The sands, which contain a certain amount of slimes, are treated as ordinary tailings, percolating, with a vacuum of 20 inches, at the rate of about 5 to 8 tons per day. The extraction obtained is over 80 per cent., and with a few modifications in the treatment

will probably be better still. The strength of the strong solution is 5 per cent., and the solution draining off is run through the towers with that coming from the slimes. In this way the stock solution is kept up to about 1 per cent. in strength, with little variation. The slimes give no difficulty in the solution of gold, the 1 per cent. solution being apparently quite strong enough. Each slime vat is filled with slimes till the latter settle down to about the middle of the vat, the proportion of solid matter in this state to the solution being about 13 : 20. The agitator is then started, and a weak wash added, equal, if possible, to the amount of slimes. In half an hour these are sufficiently mixed, the agitator is stopped, and the clear solution is gradually decanted off by adjustable, fan-mouthed pipes. In this way, neglecting, for the sake of simplicity, the amount of solid matter in the slimes, the original amount of gold solution in the vat is reduced to a half. A second wash reduces it to a quarter, a third to an eighth, and so on. In this particular instance about 20 tons of slimes are given four washes, or from 15 to 20 tons of cyanide solution each, and one of water. The extraction of gold usually amounts to 90 per cent., a few exceptions being probably due to the imperfect decanting of the slimes from the sand vat, from which some of the coarser material might easily descend into the slime vat.

The method of working is as follows:—The ten-head employed in this process fills one vat with sand in three days. There being three sand vats, and the stamps being “hung up” during Sunday, seven days are available for treating the sand in each vat. For treating the slimes there are four vats. This allows the slimes from a day and a half’s crushing to be put into one vat, each of the four vats being filled in the six days. This gives five and a half days for washing, etc. The washes decant off, with the aid of lime, at the rate of from 15 to 20 tons per day. The extraction of gold in the precipitating boxes is so far very good. The chief advantages of this method over dry-crushing are: (1) Elimination of the expense of drying the ore; (2) increase in stamp duty; (3) economy in consumption of cyanide; (4) saving of labour in filling; (5) absence of dust. In the mill under review the stamp duty has been increased from one and a half tons to two tons per day per stamp; the consumption of cyanide decreased by about half. The “cyanides” formed in roasted ore are absent, while the strength of the solution used in the boxes is too weak to suffer much loss by agitation. Crude as are some of the appliances used in this process, the results justify the expectation that dry-crushing with cyanidation will soon be entirely displaced by wet-crushing with cyanide.
THE ACTUAL EXTRACTION BY CYANIDE.

In the past few years the New Zealand method of crushing with cyanide solution has been successfully installed at several important mines in South Dakota. The ores are highly silicious, and both oxidized and unoxidized ores are mined and treated. They may be classed as dry silicious ores.

Maillard Mines.—According to Gross* the battery solution is kept at a strength of from 1·2 to 1·3 lbs. of KCN and a protective alkalinity corresponding to from 0·8 to 1 lb. of NaOH per ton of solution. Two solutions only are used in the mill, the battery solution, assaying about 50 cents in gold per ton, and the barren solution with a strength of from 1·5 to 1·6 lbs. KCN and a protective equivalent of from 1·0 to 1·2 lbs. of NaOH per ton of solution.

The lime, which is fed with the ore into the battery, has an average composition of CaO, 92·5; MgO, 0·5; Al₂O₃ + Fe₂O₃, 2·0; insoluble, 1·5; and H₂O + CO₂, 2·5 per cent.

The stamp-duty per day was not high, averaging 2·66 tons during the latter 6 months of 1903, and 2·80 tons for the first 5 months of 1904. During May 1904 the stamp-duty was increased to 2·96 tons per day. The quantity of solution going to the battery was between 4 and 5 tons to 1 ton of ore, which, in connection with the hardness and compactness of the ore, accounts largely for the small stamp-duty.

The stamps deliver a product carrying about 60 per cent. of sands and 40 per cent. of slimes.

The separation of the sands and slimes is effected by the Merrill system, which consists in using a succession of inverted cones for throwing off the slimes, and finally using an upward stream of clear water for producing a clean sand. A clean sand is made rather than a clean slime, so as to obtain an easily leachable product. (Fig. 8.)

Sands-Treatment.—The sands are filled into the vats by a Butters’ distributor, provided with six arms, ball and roller bearings.

The sands-vats, of which there are 6, are 30 ft. in diameter by 6 ft. deep, having a lattice-filter frame; 8-oz. duck cloths are used on top of cocoa-matting; the 8-oz. duck has a life of from 8 to 10 months, and has been found to be more satisfactory and economical than the heavier grades. The sands-vats hold 140 tons, and are filled in about 60 hours by dry-filling, instead of starting to charge the vat already full of solution.

When full, battery solution is run on for a period of ten days on the average. Then follows barren sump solution for about

six more days. The vat is then drained, and 15 tons of wash water passed through. Each vat is treated with about 900 tons of battery solution and 450 tons of sump solution, in addition to about 700 tons of solution filtering through the solution during filling.

*Slimes Treatment.*—The slimes are treated in eight vats, including the two for loading, each 24 ft. in diameter and 12 ft. deep. All are connected to two No. 4 centrifugal pumps with 4 in. suction and discharge, which, running at 550 revolutions per minute, handle 50 tons of wet pulp per hour.

The stuffing-box of the centrifugal pumps is supplied with solution or water under pressure to avoid undue wearing of the shaft and the cutting out of the packing.

The loading-vats are provided with a partition through the
centre of the vat extending to within 30 in. of the bottom, thus allowing the slimes to settle sufficiently so that the clear solution may be decanted from one side of the partition while the vat is being filled on the other side. The time of filling one of the vats is 12 hours; the stream being then turned into the other loading-vat.

The slimes passing into the loading-vat have 12 tons of solution to each ton of dry slimes, and during a loading of 12 hours about 150 tons (or one-half) of the incoming solution are decanted sufficiently clear to render objects visible through a distance of 2 or 3 ft. This loading-vat, just filled, is decanted as closely as possible, the slimes being transferred by the centrifugal pump to vat No. 1, forming one-half of a charge, barren solution being added at the same time. The material in the second loading-vat, after decantation, is transferred to vat No. 2 as one-half of the charge with barren solution; and upon the decantation of these two vats, the combined contents, now constituting the full charge, are pumped to a third vat. Two more transfers and dilutions are then given with barren solution, and, finally, one with water. After each transfer and dilution, several hours of agitation are given by pumping out from the bottom and discharging into the top of the same vat. For the past 11 months the average time of agitation for each charge was 13.87 hours during solution dilution and 1.97 during wash-water treatment.

It will be noted that the two largest dilutions are obtained on the half charges when the contained solutions are the richest. A charge of 60 tons of dry slimes from a 24-hour run gets practically the following dilutions with barren solution:—

- First half charge, 30 tons, 1 dilution of 100 tons.
- Second half charge, 30 tons, 1 dilution of 100 tons.
- Full charge, 60 tons, 3 dilutions of 65 tons.

This treatment makes a total dilution of 395 tons, or a little more than 6.5 tons of barren solution to 1 ton of dry slimes, the actual figures for the past 11 months being 6.57 tons of barren solution and 0.93 ton of water per ton of dry slimes. Theoretical calculations on this amount of dilution, on the assumption that the total extraction has taken place before the first decantation, and taking the value of the barren solution at 10 cents per ton, the dissolved gold passing out with the slimes-tails, should assay from 12 to 20 cents per ton solution, starting with a head-solution of a value from $1 to $2. However, the extraction has not all taken place, but continues slowly throughout the entire treatment, and for a period of six months the solutions finally going out with the slimes-tails showed an average
value of 46·1 cents per ton, with an average value of $1.80 for the head-solution, and a value of 10·6 cents for the barren solution. During a period when precipitation was giving trouble, the value of barren solution averaging 26·7 cents, the value of the head-solution averaged $1.48, and that of the solutions passing out with the slimes averaged 49·8 cents per ton.

The decantations are brought down to a pulp containing from 55 to 60 per cent. of moisture (50 per cent. moisture being equivalent to equal weights of dry pulp and solution). After the decantation of the wash-water, the top layer of the thinner slimes is drawn off, and thrown back to the charge next following. In this way the slimes going to waste are drier, averaging, for the past 10 months, 47·0 per cent. of moisture.

The gold and silver are precipitated with zinc shavings in the ordinary zinc boxes. The gold-zinc slimes are treated by sulphuric process and smelted with the following flux:

- Dried gold slimes, ....... 10 parts
- Sodium bicarbonate, ...... 4 "
- Borax glass, ............. 1 "
- Sand containing 60 per cent. SiO₂, .... 1·5 "
- Fluorspar, ............... 0·2 "

The charge melts easily, and gives a clean liquid slag.

The consumption and costs are as follows, calculated from a period of eight months:

- Cyanide, ............... 1·25 lbs. per ton of ore treated.
- Lime, ................. 5·49 "
- Zinc, .................. 1·34 "
- Zinc, ............. 0·295 cyanide solution.
- Cost of milling and treating, .... 6s. 8d. per ton of ore.
- Cost of cyaniding, ........ 2s 9d. "

Recovery of bullion:

- Gold, .................. 75·52 per cent.
- Silver, ................. 44·30 "

Of the extractions 47·9 per cent. are obtained in the battery, 27·5 per cent. from the sand, and 25·1 per cent. from the slimes treatment.

COARSE CRUSHING AND CYANIDING.

In a few rare cases it is found that an adequate extraction of the gold can be obtained without reducing the ore to the customary state of fine subdivision required with most ores.
Coarse crushing and cyaniding has been in use at the Mercur mine, Utah, for some years, and is now in successful operation at the Bau and Bidi mines in Upper Sarawak, Borneo.

It is somewhat singular that the ore and its mineral contents at Mercur and Upper Sarawak are of a closely related origin; that is, the gold ores at each are manifestly genetically connected with solfataric action. Those at Mercur contain cinnabar, orpiment, and realgar; those at Bau and Bidi, antimony, native arsenic, and realgar; and at each a comparatively high extraction is obtained notwithstanding the presence of such active cyanicides.

When treating ores containing base sulphides soluble in the cyanide solution, it is advisable to test the solutions for alkaline sulphide and sulphocyanides, at frequent intervals. In most cases the addition of the desulphurizer lead acetate will be beneficial.

**Mercur Mine.**—The description of the ore and particulars of the cyanide treatment are as follows:—

"Silica in a form similar to silicious sinter, or geyserite, characterizes the ore. Cinnabar is most abundant in this rock, and forms beautiful incrustations in the cellular varieties. Wherever found in the district, cinnabar is considered a sure sign of gold. Orpiment and realgar occur in large quantities in some of the ore. There is about as much iron as is usually found in impure limestone and clay. Barite and gypsum occur more or less crystallized; also masses of limestone are found mineralized in rings, the outside assaying from 6 dollars to 8 dollars, and the centre a trace in gold. No free gold is visible in the ore, even with a microscope. One remarkable feature is the absence of silver. The average of the ore milled is kept close to 12 dollars per ton."*

At the Mercur mill the ore is delivered by the railroad to an ore-bin 40 ft. long, 20 ft. wide, and 20 ft. deep. It is crushed in a Dodge crusher, from which it passes to a set of Walls corrugated rolls, and is finally trammed to the cyanide vats, after being crushed to 1 in. mesh or less. The vats are 12 ft. 8 in. in diameter, and hold 15 tons. They are made of tank iron, with red-wood bottoms. The filter-cloth on the false bottom is burlap, and lasts from four to six weeks. From the tanks the solution is conveyed to a collecting tank, from which it is pumped by Blake single-acting pumps to the precipitating room. The zinc boxes are from 24 to 36 in. wide, 10 to 12 in. deep, and about 20 ft. long.

Fine crushing is found to be unnecessary, as the ore is very

* Mining and Scientific Press, 23rd May 1896.*
porous, and much of it disintegrates into mud when solutions are applied.

It is interesting to note one change made in Mercur practice. Formerly the strong solution was run through the ore continuously, the surface being kept always covered. Now a series of washes is run through, the solution each time being down below the surface. The extraction has been increased thereby, and much time saved on each vat.

The solution used is from 0·1 per cent. to 0·3 per cent. in strength. It was at one time the practice to estimate the strength of the solution by its action in the zinc boxes, and by its alkaline feel. At the present time more acute methods are practised. Still an instance has come within our notice of an operator determining the strength of his solutions wholly by their odour.

At the Mercur mill the practice was formerly to standardize solutions by adding cyanide to the lower end of the zinc boxes, "the judgment of the operator determining the time and amount."

The zinc slimes were dried in an old retort belonging to the amalgamating mill. The door is closed but not luted, and at about 160° C. the product ignites, producing fumes of a complex nature, causing salivation and a headache. The slimes are finally taken from the retort, and the burning completed on a sheet-iron table. This product is then shipped to a smelter for refining.

At present the residues from a 12 dollar ore assay about 1.75 dollars, giving an extraction of 85 per cent. The average ore value is about 6 dollars per ton. The cost of treatment is itemized as follows:—Mining, 35 c. per ton; railroad hauling and milling, 80 c.; cyaniding the ore, 1 dollar 35 c.;—total cost per ton, 2 dollars 50 cents. The consumption of cyanide is at present about ½ lb. per ton of ore.

**Sarawak Practice.**—At Bau and Bidi in North-West Sarawak, Borneo, the gold-bearing material is coarse crushed and treated in vats by percolation, and notwithstanding the occasional presence of arsenic and antimony, a satisfactory recovery of the values is obtained at a low cost per ton of material handled.

As a rule, exceptional or novel methods of cyanide treatment are not due to the caprice of the management, but are rather the result of a gradual adaptation of first principles to cope with certain peculiarities of physical or chemical condition presented by the ore to be treated. Such is the case in Upper Sarawak. The gold occurs in a condition that is unique, and profoundly interesting to the geologist from a genetic standpoint.

**Geological conditions.**—The gold deposits occur along a narrow strip of country about seven miles in length, the general trend
being N.E.—S.W. The country rocks are a limestone, shales, and sandstones, lying nearly horizontal, and said by Boehm and others to be Jurassic, a northerly continuation of the great Jurassic system of New Guinea. The limestone is the dominant and lowest member of the series. It is intruded by numerous dykes and sills of quartz-porphyry, and dislocated by faults.

Character of Ore at Bidi.—For the following information respecting Bidi ores the author is indebted to the lucid papers of T. C. Scrutton and W. Hamilton, read before the Mining and Metallurgical Institution of London on 21st December 1905.

The gold-bearing material may be divided into three classes, namely:—

(1) Silicious deposits.
(2) Shale deposits.
(3) Earthy (clay) deposits.

These deposits occur as hills or mounds, surmounting pipes passing down into the limestone. In all cases the material is won by open cuts and benches.

The clays (3), which yield the greater part of the gold, form a layer spread over the irregular surface of the limestone, ranging from 1 to 20 feet thick. They are brightly coloured with oxides of iron and manganese. In places they enclose masses of silicified shale and quartz, and in the pipes detached masses of limestone. The clays are probably residual clays, resulting from the underground corrosion of the shale and limestone by hydrothermal action.

The following is the analysis of a typical sample of the clay taken within 1000 yards of a neck in the limestone:—

W. Hamilton.*

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<td>Gold</td>
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<tr>
<td>Silver</td>
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98·9 10 dwt. per ton.
11·5 " "

From these results it would appear that both the arsenic and antimony occur in a native state. The absence of CO$_2$ would indicate that the lime occurs in a combined form probably with the SiO$_2$.

This clay yields an extraction of 78 per cent. of its gold contents when leached in 6-in. lumps with cyanide. Grading experiments show that 99.1 per cent. of the clay passed through a 100-mesh sieve.*

The silicified shales occur in situ, forming rounded hills surmounting a pipe or vent in the limestone. In these cases the shales lying near the pipe are highly silicified, often being a hard silicified breccia, consisting of fine particles of shale cemented with silica. The shale carries gold values ranging from 1 to 10 dwt. per ton. Its thickness seldom exceeds 30 feet.

The silicious deposits (1) form mounds consisting of irregular masses of quartz embedded in clays which pass downward into a lenticular cavity or pipe in the limestone.

At the surface the quartz is often vesicular, white or grey in colour, or tinged yellow with orpiment. Deeper down it is dark blue, and the holes are filled with native arsenic. It also contains realgar and stibnite, the latter occurring as small detached needles, but masses of stibnite of all sizes up to those weighing a ton are not uncommon.

With increasing depth the contents of the pipes become more and more mineralized with native arsenic and realgar, and are generally richer in gold, until at a depth of 30 feet the filling consists of semi-liquid clay of a dark blue, lemon-yellow, orange, or deep red colour, mixed with masses of dark blue or black silicious stone highly charged with arsenic and antimony. At the ends of the pipe-cavity this stone is associated with calcite intruding through fissures in the limestone.

The cavity is usually lenticular in form, and varies from 20 ft. to 200 ft. in length, and from 10 ft. to 60 ft. in width.

At a depth of about 40 ft. below the surface of the limestone the walls usually close in rapidly, the fissure being filled with calcite and blue silicious stone containing needles of antimony, as found in the ends of the cavity.

The evidence seems to leave little room for doubt that the pipes were the vents of solfataric action, almost similar to those existing at Ohacowai Hot Springs, north of Auckland. The vents and fissures have probably become filled with calcite since the cessation of hydrothermal activity.

As a general rule, the gold is so fine that it will not show in the

---

pan,* and is attached to the native arsenic rather than the antimony (stibnite).†

Cyanide Treatment.—At the Bidi mill and cyanide plant the rock is crushed to 1-in. or 2-in. cubes in Blake-Marsden rock-breakers. The clay undergoes no preliminary treatment whatever, except that when made very wet by rains and floods it is stacked in drying sheds in layers varying from 2 ft. to 4 ft. thick until the moisture has been reduced to about 20 per cent.

The cyanide plant contains 30 steel vats, 18 of which are 6 ft. deep and 27·5 feet in diameter, with a capacity of 100 tons. The other 12 vats are 3 ft. deep and 27·5 ft. in diameter, with a capacity of 50 tons. All the vats are made of 3-in. mild steel plates, riveted with ½-in. rivets with a 1½-in. pitch.

The shallow vats are charged with clay only, and the percolation is good. The charging takes about two hours and a half, and the discharging, by shovelling through four bottom doors into wagons on two lines of rail, two hours. The treatment occupies seven days.

The 50-ton vats after charging are upward-leached with strong cyanide solution, 0·08 per cent. KCy, which causes the charge of clay to sink down 8 or 9 inches. The solution is kept in contact with the charge for two hours, after which it is allowed to drain and run to the zinc boxes. Solution is now run on at the surface five times a day, which allows each vat to be shut up for one hour after each pumping, and to drain dry before the next solution is applied.

Each vat receives 6 tons of solution at a time, that is, 30 tons per day. For the first four days the solution used has a strength of 0·08 per cent, and for the next three days about 0·04 per cent. The alkalinity is regulated by the working of the zinc boxes.

The 100-ton vats are charged with both clay and crushed stone. A layer of clay 2 feet deep is placed on the bottom and levelled, over this is laid a layer of stone 1½ to 2 feet deep, and over this a layer of clay 2 feet deep. The treatment occupies about ten days and is as follows:—

Strong solution, 30 tons, 0·07 per cent. KCy, upward-leaching taking three hours.
Solution in contact with ore, four to six hours.
Solution allowed to drain takes from four to five hours.
Solution pumped up on top twice a day until 150 tons of strong solution have been used.
Each solution is shut up six hours and drains six hours.

† Cecil J. Brooks, Discussion, loc. cit., p. 7.
Sump solution, 180 tons, 0.05 per cent. KCy, five days, applied on top.

The solutions are kept at a definite standard of alkalinity, partly by addition of lime to each waggon of ore, and partly by addition of soda to the sumps. The average consumption of lime is 8.8 lbs. of lime per ton of ore, and the maximum amount of soda added 0.1416 per ton of solution.

Large quantities of lime are required to neutralize the organic matter in the clays. The lime also helps the solutions to percolate more freely.

Prolonged soaking has not been found to give good results, but on the other hand each new solution must be given time to permeate the whole of the charge before draining begins.

Slow draining of the solutions is a necessity, i.e. at the rate of five tons per hour. When this rate is exceeded, too large a proportion of moisture remains in the vat, doubtless due to the rapid closing of the interstices, thereby cutting off the supply of air.

The silicious shales are crushed through a rock-breaker set to 2-in. gauge, roasted in a shaft furnace, drenched with water to split the rock up and expose the cementing medium which carries the gold, and then cyanided. The extraction amounts to 70 per cent.

Extractor House.—The precipitation is effected by zinc shavings. There are 16 zinc boxes, containing in all 936 cubic feet of zinc, which allows 1 cubic foot of zinc per ton of solution in 24 hours, at the usual rate of flow.

The precipitation gives very little trouble, and when working satisfactorily 0.05 per cent. cyanide solution entering with 2 dwt. of gold per ton is reduced to a solution containing 3 grains per ton.

With so much arsenic and antimony present it is found necessary to regulate carefully the relative quantities of lime and soda used, so as to obtain the requisite alkalinity of solution. When too much soda is used large quantities of arsenic and antimony are dissolved from the ore in the vat, and become deposited on the zinc, forming a coating which in cases prevents any precipitation of the gold; but generally, owing to the large area of zinc, the efficiency of the precipitation is not greatly affected. Defective precipitation is remedied by placing several lumps of cyanide in the head of the boxes.

An excessive use of lime is always avoided, as the lime becomes deposited in the extractor boxes, where it mixes with the slimes, which are thus made excessively bulky and more costly to handle in the clean-up. Soda has a stimulating effect upon the zinc boxes, but is only used in cases of emergency, in order to keep arsenic and antimony out of the solutions.
The presence of much antimony in the solutions is always indicated by an offensive smell in the extractor house.

According to Scrutton,* the consumption of zinc amounts to about 0.58 lb. per ton of ore.

The Clean-up.—The arsenic that collects in the zinc boxes adds considerably to the labour and cost of the clean-up. Direct acid treatment is found inadmissible, owing to the formation of arseniuretted hydrogen, and the subsequent difficulty in melting the low grade bullion obtained. The process now in use may be summarized as follows:—

1. Rinsing but not rubbing the zinc to avoid formation of fines.
2. Filter-pressing all the slimes in a Johnson 19-in. gold press.
3. Roasting raw slimes thoroughly in large open pans to drive off arsenic and antimony and oxidize the zinc.
4. Crushing roasted slimes in set of hand rolls.
5. Acid treatment in vats; 350 lbs. at a time requiring 3½ jars \( \text{H}_{2}\text{SO}_{4} \). Time, 12 hours.
6. Filter pressing.
7. Washing.
8. Drying.
9. Fluxing—Slimes 100 parts, borax 50. Fusion in No. 120 crucibles, skimming and adding more slimes and flux through a long funnel until the charge contains about 30 lbs. of bullion, when contents are poured into conical mould. On cooling, a layer of speise about ½ inch thick forms on top.
10. Separating bullion from speise by melting bullion and speise in No. 30 pot and pouring into rectangular mould. After the bullion solidifies the speise is easily scraped off, leaving a clean surface.
11. Re-melting the bullion in the same pot, taking a dip sample for assay, and then pouring into mould.

The speise is smelted with thin sheet-iron and borax for about three hours, and the fused mass then poured into a conical mould. The speise collects at the bottom, and is easily separated from the arsenide of iron above. The speise is now re-melted, sampled, and poured. It shows a fineness of about gold 100, silver 300, and is shipped to London.

Costs.—The cost of cyaniding is given by Scrutton as 2s. 2d. per ton of ore treated, a most satisfactory result. The total cost of ore winning, crushing, cyaniding, laboratory, power, general

charges, prospecting, etc., amounts to 10s. 3·4d. per ton of ore treated.

*Cyanide Treatment at Bau Mine.*—The gold-bearing material is somewhat similar to that at Bidi, but is less mineralized. At the mill the ore is crushed to pass a half-inch mesh screen, and then elevated into a 400-ton ore-bin, into which the clay is dumped direct after passing through a 2½-inch grizzley.*

The proportion of clay to ore going into the bin is roughly 5 to 3, and about 10 lbs. to 14 lbs. of lime to the ton of ore are added to the vat charge.

The vats are 40 feet in diameter and 8 feet deep. They have a capacity of about 340 tons dry weight.

The practice is to use about 400 tons of 0·05 per cent. cyanide solution, followed by an equal quantity of wash-water. The extraction amounted to 75 per cent. of the gold contents of an ore averaging about 5 dwt. of gold per ton.

Zinc shavings are used to precipitate the gold from the cyanide solutions, but instead of the customary rectangular boxes, circular tubs 8 feet in diameter and 4 feet 6 inches deep have been in use for some years. The tubs are fitted with a false-bottom, under which the solution is led by a vertical pipe in the centre of the tub. On the top of the false-bottom there rests a sieve on which the zinc shavings are laid.

The capacity of these extractors is about 10 tons of 0·02 per cent. KCy solution per hour. The tubs require little or no attention, and there is no need to handle the zinc, the fresh shavings being added at the top as required. The solution flows upward and escapes through numerous holes round the top of the tub. The capacity of the mill is about 110,000 tons per annum. The total cost of all charges amounted to only 7s. per ton. Chinese and native labour is used in all the different operations of mining, crushing, and treatment of the ore. As at Bidi, the overburden is stripped, and the gold-bearing material filled direct into waggons from open workings.

CHAPTER VII.

THE PRODUCTION AND TREATMENT OF SLIMES.

With all methods of wet-crushing and pulverizing the formation of a certain proportion of slimes seems inevitable, and when the ore contains metallic oxides, clay, or other earthy matter, the production is often very large. In ores containing a certain proportion of very fine or 'float' gold, the slimes are generally a valuable product, and their successful treatment has engaged the attention of metallurgists for many years.

Since the introduction of the cyanide process, many attempts have been made to leach the slimes resulting from wet-crushing, on an economic scale, and the subject is one to which the author has devoted much thought and experiment for a number of years. The problem is principally a mechanical one, consisting in the difficulty of separating the solutions from the slimy mass, rapidly, effectively, and at such a cost as to permit the treatment of slimes of low value.

Up to the present a great many different devices have been tried, with varying degrees of success. Among these may be mentioned compression by hydraulic and other presses; agitation and centrifugal force; agitation and decantation; and agitation with filtration, aided by an artificial vacuum.

The dry-crushing of ores by means of the Californian stamp battery is always attended with the production of so large a proportion of slimes that only a shallow depth of the pulverized material can be leached in the direct method of cyanide treatment.

The stamp battery was invented for wet-crushing, for which it stands unrivalled for all classes of ore. For dry-crushing it is a most unscientific machine, on account of its inability to discharge the dust when reduced to the requisite fineness. Nevertheless, no better has yet been invented, except perhaps the Edison rolls.
Definition of Slimes.—When the cyanide process was first introduced, and for some years after, it was the practice of the millman to prevent the formation of slimes in the battery pulp by every contrivance that skill and experience could suggest. A large proportion of slimes in the sands was looked upon with something akin to apprehension by the most accomplished cyanide operators of that time.

What the metallurgist demanded was a leachable product. When the sands contained so much fine matter that percolation became difficult, they were said to be slimy, or to contain a large proportion of slimes. When percolation was found to be impossible, even by an artificial vacuum, the product was generally spoken of as being "all slimes."

It is well known that what is slimes for one class of material at the same degree of fineness is not slimes for another. Thus, while the degree of fineness is an important, perhaps the prime, factor in creating what is termed slimes, it is not the sole factor. "Slimes" is a relative rather than an absolute term, and for this reason it is impossible to formulate a strictly scientific definition of it.

It is well known that perfectly homogeneous pulverized matter of all kinds at a certain degree of fineness possesses the power to absorb and retain moisture in such a way as to resist displacement, except some external force is used to expel it.

The author prefers to define slimes as the product of fine grinding which is unleachable by percolation without the application of external force.

The quantity of dry slime in the moist pulp can be calculated by the formula suggested by Caldecott.*

Let \( s = \) specific gravity of dry slime.

\( a = \) specific gravity of pulp found by weighing equal volumes of water and pulp, and calculating ratio.

\( P = \) percentage by weight of dry slimes in pulp.

Then \( P = \frac{100s (a - 1)}{a (s - 1)} \)

Production of Slimes with Tube-Mills.—The commercial success of slimes treatment in Kalgoorlie has directed attention to the probable advantages to be derived from the finer crushing of all gold- and silver-bearing ores preparatory to cyanide treatment.

The production of a certain proportion of slimes, generally ranging from 15 per cent. to 35 per cent. according to the

* W. A. Caldecott, *Jour. Chem. and Met. Soc. of South Africa*, vol. i. No. 3.
character of the ore and mesh of screen, attends the pulverizing of ores by all classes of machines. The cost of this proportion of slimes will bear the same ratio to the total cost of pulverizing that the slimes bear to the quantity of original ore. Thus, if the crushing charges amount to 3s. per ton and the slimes produced amount to 25 per cent. of the ore crushed, it is evident that the cost of producing this proportion of slimes will be 9d. per ton.

The sliming of an additional 25 per cent. of the sands will cost from 2s. to 3s. per ton of slimes, according to the charges for wages, power, material, depreciation, etc. The problem is—Will the additional extraction from the slimes yield a profit? The solution is a simple arithmetical exercise, when the respective values carried by the slimes and sands, and the relative extractions (and costs) have been ascertained.

At Kalgoorlie, practically the whole of the ore milled is slimed, a practice which is found to be a necessity in order to obtain an adequate extraction of the values.

For all ores there will be a maximum paying point between the extremes of coarse milling and sliming. This point can only be determined by careful working tests; and in the majority of cases it will be advisable to work from the coarse to the fine, rather than from the fine to the coarse.

The quantity of slimes produced will depend primarily upon the physical and chemical character of the ore, being least with hard chalcedonic silicious ores, and greatest with clayey talcose ores.

The machines used for sliming at Kalgoorlie are the improved wheeler pan and the tube-mill, often called grit-mill or flint-mill.

The comparative efficiency of these machines, and much useful information respecting the method of running, maintenance, and cost of sliming, were elicited in the discussion of four papers* on fine grinding, read before the Institution of Mining and Metallurgy, London, in October 1904.

Mr. Walter Broadbridge † stated that numerous experiments made at the Ivanhoe mine proved that the cost of grinding with pans and mills is practically the same, but for very fine sliming the mills are the best. The advantage, however, was with the pans in first cost:—

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The following tests, also made at the Ivanhoe mine, are interesting, as showing the difference when mills and pans are variously equipped:

<table>
<thead>
<tr>
<th>Sands before Grinding</th>
<th>Sands after Grinding</th>
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<tbody>
<tr>
<td></td>
<td>16 ft. Mill fitted with old liners and small flints</td>
</tr>
<tr>
<td>On 40 Mesh</td>
<td>%</td>
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<tr>
<td>&quot; 60 &quot;</td>
<td>38.4</td>
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<td>&quot; 100 &quot;</td>
<td>41.6</td>
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<tr>
<td>&quot; 150 &quot;</td>
<td>15.4</td>
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<tr>
<td>Thro’ 150 &quot;</td>
<td>1.6</td>
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<td>2.8</td>
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<td>100.0*</td>
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**Same Tests with Pans.**

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<tr>
<th>Sands before Grinding</th>
<th>Sands after Grinding</th>
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<tr>
<td></td>
<td>Pans fitted with shoes half worn out</td>
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<tr>
<td>On 40 Mesh</td>
<td>%</td>
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<tr>
<td>&quot; 60 &quot;</td>
<td>38.4</td>
</tr>
<tr>
<td>&quot; 100 &quot;</td>
<td>41.6</td>
</tr>
<tr>
<td>&quot; 150 &quot;</td>
<td>15.4</td>
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<tr>
<td>Thro’ 150 &quot;</td>
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<td>2.8</td>
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<tr>
<td>100.0*</td>
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* [99.8]
Thus there is a difference in fine slimes of 13·59 per cent. with
the mill and 6·35 per cent. with the pan.

Mention was made of pans fitted with compensation weights—an entirely new feature with Wheeler pans. The Ivanhoe was
the first mine to use them. They are made of cast-iron in the
shape of a ring, and fit on the top of the muller to which the shoes
are fastened, and can be put in in five minutes. These weights
are only added when the shoes are half-worn, so that the horse-
power keeps constant. The following trials give very conclusive
results:

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<tbody>
<tr>
<td>Pan with Weights</td>
<td>Nil</td>
<td>3·3</td>
<td>11·7</td>
<td>11·7</td>
<td>76·3 = 100·0</td>
</tr>
<tr>
<td>Pan without ,,</td>
<td>Nil</td>
<td>3·5</td>
<td>48·5</td>
<td>10·2</td>
<td>37·8 = 100·0</td>
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<tbody>
<tr>
<td>Pan with Weights</td>
<td>1·1</td>
<td>1·7</td>
<td>24·2</td>
<td>18·5</td>
<td>55·5 = 100·0</td>
</tr>
<tr>
<td>Pan without ,,</td>
<td>0·5</td>
<td>16·7</td>
<td>50·2</td>
<td>8·9</td>
<td>23·7 = 100·0</td>
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The foregoing figures show that very good work is being done
at the Ivanhoe mine with Wheeler pans.

It would be interesting to know the relative efficiency of pans
and tube-mills when working on the harder silicious ores of the
Rand.

The cost of fine-grinding at El Oro and some Kalgoorlie mines
is as follows:

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<tr>
<td>Hannan’s Star, treating 1·6 per hour,</td>
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<td>Oroya Brownhill, ,, 1·77 ,, ,, 3 6 ,, ,,</td>
<td></td>
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<tr>
<td>Lake View Consols, ,, 1·87 ,, ,, 4 9 ,, ,,</td>
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<tr>
<td>El Oro, ,, 2·00 ,, ,, 3 4·5 ,, ,,</td>
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The following useful points were elicited during the discussion:
(1.) That the smaller tube-mills, say 13 ft. 7 in. by 3 ft. 8 in.,
are easier to manage than the large mills, and have a relatively
higher efficiency.
(2.) That with dry-crushing, small flints give the best results, and with wet-crushing, large flints.
(3.) That the mills must be run to their full capacity, or the flints will grind themselves, thereby increasing the cost of maintenance.
(4.) That hard iron liners are better than steel, and flint linings better than iron (S. Robinson, loc. cit., pp. 135, 136).
(5.) That the consumption of cyanide due to contact with the iron linings is very little (Grüssner, loc. cit., p. 88).
(6.) That the wear of flint balls amounted to about 1 lb. per ton of ore treated.
(7.) That iron liners lasted from six to nine months.
(8.) To obtain the highest efficiency, the tube-mill must be run at a regular maximum rate of speed, which must never be exceeded or dropped (W. H. Trewartha-James, loc. cit., p. 118).

According to the last authority, the thickness of the pulp and the rate of feed were points of great importance. If the pulp was too thin, the balls worked upon themselves and the liners. Even small variations, when the mill was properly adjusted to its work, became critical. The regulation of the thickness of the pulp was the crucial point in tube-mill practice. The linings in the Hannan's Star mills were all bolted through the shell, which was found to be an advantage.

It should be noted, he said, that at the Hannan's Star mill they did not attempt to get an immediate or direct product at one operation from the tube-mill. The ground pulp was elevated to the spitzkasten; and the underflow went back to the mill, passing and re-passing till fine enough, so that the mill, which was nominally dealing with sands from 78 tons per day, and which was actually sliming only 38 to 45 tons of sand per day, passed through it something like 268 tons during the 24 hours.

He was told that the tube-mill could be modified to produce a completely slimed product at one operation. So far as he could see, that could only be done by lengthening the tube. If lengthened too much, of course there would be excessive wear. If it were not long enough, one certainly would not get, as the ore passed through the mill, the effect one wished to produce. He was personally of opinion that the tube-mill industry was only in its infancy to-day.

Although they had had the tube-mill working certainly for over five years, he did not think that they knew very much about it yet, because, with the stress of actual requirements, they had no time to experiment. The Oroya Brownhill Co., and he thought the Ivanhoe, had adopted the smaller 13-ft. mill, and they claimed they got a greater efficiency per horse-power with those smaller mills.
THE PRODUCTION OF SLIMES.

Data at first sight appeared to warrant that opinion, but he thought it was a case of one of those comparisons which was made fairly enough in good faith, but which was not sufficiently accurate to be really true. He believed that the 16-ft. mill gave them more work units for their horse-power than the 13-ft. mill, and he believed that a larger mill would do still better work for them. He believed the tendency would be to extend the length of the tube so that the product should be as near as possible the product sought, and the amount of returns the smallest possible.

(9.) The pulp fed to the tube-mill should not contain more than 33 per cent. of water (S. Robinson, loc. cit., p. 134). At Hannan’s Star the pulp is thickened to the utmost possible (Trewartha-James, loc. cit., p. 134). A pulp of 50 per cent. thickness gave good results, but a thicker pulp (60 per cent.) was in successful use (Alfred James, loc. cit., p. 145).

Mr. M. Davidsen supplied the following useful information:—

For grinding hard quartz ores only natural flint pebbles should be used. The pebbles vary in size from $\frac{3}{4}$ inch to 3 inches or 4 inches in diameter.

In wet-crushing the large pebbles work quite as well as the small pebbles, and even better, inasmuch as the use of large pebbles allows the mill to be fed with coarser material, say passing a No. 6 sieve.

The best work is done when the mill is half filled with pebbles. If the interior volume of a tube-mill is $N$ cub. ft., the weight $W$ of pebbles which should be loaded in the mill in pounds is—

$$W = 44N.$$  

The number of revolutions which give the best results is obtained as follows:—

Let $D =$ interior diameter of tube in inches

$$V = \text{revolutions per minute}$$

Then

$$V = \frac{200}{\sqrt{D}}.$$  

The power required is obtained from the expression—

$$P = 0.15N$$

where $N =$ interior volume of tube in cub. ft.

The theory of the tube-mill has been made the subject of elaborate experiment by Fischer * and White,† whose results are

* Herman Fischer, Zeit. des Vereines deutsch. Ingenieure, 26th Mar. 1904; and Engineering and Mining Journal, 14th Nov. 1904, p. 791.
not much different from those of Davidsen. And more recently an instructive and useful series of working experiments has been carried out at El Oro mill, Mexico, and described in a paper entitled "Fine Grinding by Tube-Mills," by G. Caetani and E. Burt.*

The following particulars concerning tube-mill practice at Kalgoorlie will be of interest (Grüssner, loc. cit., p. 89).

OROYA BROWNHILL, G. M. Co., W.A.

Five mills, 3 ft. 7 in. by 13 ft. 2 in.; hard iron liners.
One mill for each 10 stamps of 1050 lbs. each. Capacity, 31,500 tons through 25 mesh in 15 months, or 37.5 tons battery pulp per 24 hours per mill went through the spitzluten into the mill to grind the sands to 220 mesh.

Power consumption 18 h.p. per mill = 1s. 4·9d. per ton crude ore.
In 15 months, 2 sets of plates = 2·56d. per ton ore, and 16 tons flints = 1·32d. per ton ore, have been worn out. Total for power and wear = 1s. 8·78d. per ton crude ore.

LAKE VIEW CONSOLS, W.A.

Three mills, 3 ft. 11 in. x 16 ft. 5 in.
Stamps and mesh same as Oroya Brownhill. Capacity of 1 mill, 67 tons battery pulp per 24 hours, grinding to 220 mesh.

Power consumption, 30 h.p. per mill.

One set of liners wore out in 145 days, corresponding to 10,530 tons crude ore. Wear of flint stones, 848 lbs. per 1000 tons of crude ore.

THE IVANHOE GOLD CORPORATION, LTD., W.A.

Sands fed into the mill: coarser than 40 mesh = 38·4%; 40-60 = 41·6%; 60-100 = 15·4%; 100-150 = 1·8%; through 150 = 2·8%. Finished product: coarser than 40 mesh = 0·3%; 40-60 = 10·14%; 60-100 = 39·63%; 100-150 = 9·63%; through 150 = 40·3%. Capacity of 3 ft. 7 in. x 13 ft. 2 in. mill, 38 tons per 24 hours; power = 17 h.p.

HORSE-POWER REQUIRED.

Oroya Brownhill, 13 feet tube-mills require 18 H.P
Lake View Consols, 16 " " 30 ",
New Goch mine, Rand, 26 " " 66 ",
El Oro mine, Mexico, 25 " " 80 ",

THE TREATMENT OF SLIMES.

TREATMENT OF SLIMES.

The different processes of slime treatment at present in use may be classified as follows:—

1. Agitation and Decantation, as practised in Africa, America, and New Zealand.
2. Filter-press method, as practised in Western Australia and New Zealand.

AGITATION AND DECANTATION.

South African Practice.—The main features of the decantation process as practised in South Africa are as follows:—

1. Spitzbute separation of sands and slimes.
2. Spitzkasten concentration of slimes.
3. Collection and settlement of slimes in collecting vats provided with decanting syphon.
4. Agitation of slimes with cyanide solution.
5. Settlement of slimes and decantation of solution for precipitation.

At the Rand mines, the agitation is generally effected by a centrifugal pump, but in some cases revolving stirrers are used.

In 1896 Mr. C. Butters, then manager of the Rand Central Ore Reduction Company, erected a large slime plant at the Robinson mine, at the cost of nearly £60,000. The salient features of the treatment are the classification of the slimes into three products. The first and second products, which consisted of fine sands, were treated by ordinary leaching in vats; while the fine slimes were leached in vats with revolving agitators, by a process patented in New Zealand by the author early in 1893. The agitators were about 10 ft. in diameter, and provided with a filter-frame and webbing. During agitation the solution was drawn off by means of an air-pump connected with a vacuum-cylinder. A fuller description of this process is given further on.

On the Rand, the treatment of slimes has naturally been a very serious problem, to the solution of which much time and money have been devoted. The following method of treating slimes, known as the “natural settlement” process, has been in use at the Crown Reef since August 1896, with satisfactory results. It was first devised by Mr. J. R. Williams, and afterwards adopted by the Robinson Slime Works.* To the water carrying the

* Chem. and Met. Soc. of S. Africa, July 1897.
slimes from the separator plant is added milk of lime, the slimes being thus precipitated in a flocculent form. The supply of lime is regulated by an automatic feeder, as too much lime is as bad as too little, since it interferes with the efficient precipitation of the gold.

The slimes are settled in three large pointed boxes, two of them 20 ft. x 20 ft. and 10 ft. deep, and the third 40 ft. x 40 ft. and 10 ft. deep. The settled slimes are drawn off at the bottom and pumped into the first two of eight treatment vats, about 90 per cent. of the water contained in them having been separated.

The vats are each 32 ft. in diameter and 10 feet deep, having a conical bottom. More water having been separated from the slimes by allowing them to settle, they are sluiced into a pump by a jet of cyanide solution and transferred to a second series of two vats filled with solution, and the strength of which is brought up to 0·1 per cent. of KCy. A centrifugal pump is used for circulating the pulp. After settlement, and prior to cyanide treatment, the pulp consists of about 1 part dry slimes to 1 part water.*

About 80 per cent. of the gold is dissolved in the passage through the pumps, but agitation is continued from one to two hours by withdrawing the solution at the bottom, and discharging it in oblique jets at the top and through the sides.

The slimes are then allowed to settle, and the clear solution is drawn off through side-cocks (which have been replaced by syphon pipes at the Robinson Works), and passed to the precipitation boxes.

The residual slimes are then pumped successively into the third and fourth series of two vats, where they are further agitated with extremely dilute solutions of cyanide and allowed to settle. These solutions do not pass to the precipitation boxes, but are transferred to the preceding series of vats.

The treatment having been completed, the residue slimes, containing, say, 50 per cent. moisture, are now sent to the residue slimes dam, either by sluicing out direct with water or by passing through the same centrifugal sludge pump used throughout the operations, which can be connected up to deliver into the discharge pipe or launder going to the slimes dam. In treating 2½ dwt. to 3 dwt. current slimes, an extraction of, say, 75 per cent. is obtained; the total amount of solution used would amount to 5 or 6 tons per ton of dry slimes.†

The "strong" solution from the second series of vats is run

† Clement Dixon and Torrente, loc. cit., p. 775.
into two settling tanks, 15 ft. in diameter and 5 ft. deep, where it is allowed to clarify. Electrical precipitation is used. From 6643 tons treated, an actual extraction of 60½ per cent. was obtained, at a cost of 3s. 9d. per ton.

The concentration of the solution, as regards its gold contents when decanted, was first successfully introduced by Mr. J. R. Williams at the Crown Reef plant, in order to reduce the amount of solution passing through the precipitating boxes. Naturally, when a given amount of gold in solution is obtained from the slimes per twenty-four hours, if this amount of gold can be concentrated into a small volume of solution, a small precipitating capacity is required. For these reasons Mr. Williams introduced the system of double washing; that is to say, the first solution that is applied to the slimes after decantation is not clarified and passed through the precipitation boxes, but is run into what is called an "intermediate tank," and then pumped up for use again upon a fresh charge of slimes. After this second decantation it is run through the clarifying apparatus to the boxes for precipitation. The settled slimes are then re-pulped with an equal volume of solution from the precipitated sump; that is, solution which has passed through the precipitation boxes. Theoretically, the enrichment of the unpulpetitated solution could be carried on until it was equal to the value of the slimes, but in practice it has been found that the first wash which has not been precipitated, and the second wash which has been precipitated (two washes altogether), make the simplest and most perfect system.

Kalgoorlie Practice.—According to Robert Allan, M.A., B.Sc., the raw slime treatment at the Oroya Brownhill gold-mines is as follows:

The raw sludge from the pulp condensing plant is agitated with ordinary cyanide and bromo-cyanide [the Diehl process is used] in ten covered-in agitation vats (provided with the usual mechanical stirrers), from 20 to 25 ft. in diameter and from 7 ft. 6 ins. to 8 ft. deep. They hold from 90 to 110 tons of pulp, equivalent to from 41 to 50 of dry slime. Each agitator takes about four hours to fill.

The solution, which forms about half of the bulk of the sludge, is plant water containing about 8 lbs. per ton or .04 per cent. of cyanide; it is made up by the addition of more cyanide to 0.1 per cent. strength.

The contents of each vat are first agitated for three hours, then bromo-cyanide in amount proportional to the value of the pulp—about 1 lb. to each ounce of gold—is added, and the agitation continued for another twelve hours, when solution is effected. Towards the end of the agitation—about two hours before its
completion—lime in amount from 2 to 5 lbs. per ton is added. During the agitation about \( \frac{3}{2} \) to 1 lb. of ordinary cyanide per ton is consumed, whilst the whole of the bromo-cyanide is destroyed. After agitation the raw or roasted pulp is discharged into a sunken agitator.

**New Zealand Practice.**—The method of slimes treatment adopted by the Waihi Company at their new mill at Waikino in 1901 is based on the best South African practice, differing only in the separation of the gold-containing solutions, which are separated from the slimes by filter-presses instead of repeated decantation as at the Rand.

A detailed description of the methods of slime treatment at the Waihi Company’s mills of 320 stamps, furnished by the company’s metallurgist, Mr. E. G. Banks, will be found in Chapter XI.

**Author’s Experiments.**—Early in 1893 the author treated a large parcel of ore from the Monowai mine, in the Thames district. The ore consisted of hard bluish and grey-coloured splintery quartz, containing a considerable proportion of sulphides of iron, copper, lead, and zinc.

The ore was dried, dry-crushed, sampled and assayed, showing a value of £5, 5s. per ton. The crushing was effected in a stamp battery, which produced a large quantity of the finest slimes. These slimes rendered it impossible to effect the leaching by percolation, even with the aid of a vacuum. When mixed with water, the slimes, when only 4 in. thick, settled on the filter-cloth, forming an impervious bed, through which it was found impossible to draw off the solution.

The pulp was then subjected to agitation, by which the dissolution of bullion was effected in six or seven hours. The separation of the solution from the pulp, however, was a long and tedious operation, and extended over eight days. It was effected, but not very satisfactorily, by agitating the ore, allowing the slimes to settle, and then drawing the clear solution off by a syphon. The weak solution and wash-waters were added in succession, and the same operation performed after each.

In order to ascertain the degree of fineness to which the ore was reduced when crushed, a number of experiments were made with a 60-mesh, 40-mesh, and an ordinary battery-punched screen, and it was found that the results were in each case as follows:

**With 60-Mesh**—

<p>| | | | | |</p>
<table>
<thead>
<tr>
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</thead>
<tbody>
<tr>
<td></td>
<td>A. 18%</td>
<td>B. 5%</td>
<td>C. 100%</td>
<td></td>
</tr>
<tr>
<td></td>
<td>remained on</td>
<td></td>
<td>passed through a</td>
<td></td>
</tr>
<tr>
<td></td>
<td>. . . .</td>
<td>. . . .</td>
<td>. . . .</td>
<td>8100 sieve.</td>
</tr>
<tr>
<td></td>
<td>. . . .</td>
<td>. . . .</td>
<td>. . . .</td>
<td>3600 ”</td>
</tr>
<tr>
<td></td>
<td>. . . .</td>
<td>. . . .</td>
<td>. . . .</td>
<td>1600 ”</td>
</tr>
</tbody>
</table>
THE TREATMENT OF SLIMES.

WITH 40-MESH—

A. 22% remained on . . . 8100 sieve.
B. 12% " " " 3600 "
C. 3% " " " 1600 "
D. 100% passed through a . . . 900 "

WITH PUNCHED-SCREEN, EQUAL TO 30-MESH—

A. 26% remained on . . . 8100 sieve.
B. 18% " " " 3600 "
C. 13% " " " 1600 "
D. 4% " " " 900 "
E. 100% passed through a . . . 625 "

Taking the values of the different products separately, it was found that the finest in all cases gave the highest values. This also received confirmation from the circumstance that the fine dust, which had collected on an elevated platform during crushing, assayed higher than the dry material in the bin.

The relative values were as follows:—

From dust-bin, . . . £5 5 0 per ton.
" platform, . . . 6 13 6 "

The ratio of gold to silver in the pulp from the dust-bin was nearly 1 to 9, and in the dust from platform 1 to 12, thus showing an increase of the silver contents.

Vacuum Suction Process.—In 1893 the author obtained, with two of his assistants, a patent for a combined agitation and leaching process, which may be described as follows:—

The appliances used in the operation consist of a shallow circular vat, a vacuum-cylinder, and an air-pump. The vat is provided with four revolving arms, to which soft rubber brushes are attached. The bottom of the vat is provided with a false-bottom, consisting of a wooden grating covered with wool-packing or other webbing. The operation is conducted as follows:—The leaching solution, made up to the required strength, is first conducted into the vat. The revolving arms are then set in motion, and the dry pulp or fine slimes introduced. The agitation is continued for six hours, or until the extraction is complete. A stop-cock, in a pipe connecting the false-bottom of the vat and the vacuum-cylinder, is then opened, and the air-pump started. The effect is immediate. The clear solution at once begins to drain over into the vacuum-cylinder, the brushes on the revolving arms preventing the slimes from settling and choking up the filter-cloth. When the slimes have been drained down to a thick paste, the first wash is added, the pump again started, and the slimes drained
as before. The subsequent washes are applied in the same manner, and when the washing is completed, a plug or door is opened, and the leached slimes are sluiced out. The whole operation of leaching and washing takes from eighteen to twenty-four hours.

This process was adopted by the author for the treatment of several tons of ore from the Monowai mine, at Waiomo, with complete success. This ore contained a large proportion of clay and iron oxides, and, when dry-pulverized, formed a pulp which defied all the ordinary methods of percolation. Trial tests were also made with parcels of very fine slimes, and in all cases the results were most satisfactory.

The Moore vacuum suction process now in use in America and New Zealand is an ingenious adaptation and improvement of the vacuum suction principle used by the author in 1893.

Agitation with Compressed Air.—The author continued his experiments for the treatment of slimes in another direction, and in 1895, with Mr. G. W. Horn, obtained a patent for an improved leaching process for the treatment of slimes and other fine matters by cyanide or other solvents.

The necessary apparatus consists of a solution vat, an air-compressor, and a leaching vat provided with a filter-frame and webbing, both at the top and bottom. The vat may also be provided with an air-tight cover, so as to permit the creation of an artificial vacuum to facilitate the filtration.

In practice, the leaching vat, which may be constructed of wood or any suitable material, and of any convenient size, is charged with cyanide solution. The slimes are then introduced and well mixed with the solution, which is then allowed to penetrate and stand in contact with them for two hours. At the end of that time the top filter-frame and webbing is put on, and additional cyanide, or other solution, is allowed to penetrate and permeate the slimes from below, thus displacing the solution already in the vat, and causing it to pass through the top filter-web as a clear liquid, which is conducted away by a launder.

In order to prevent the solution selecting channels through the slimes, or other matter in the vat, during the upward filtration, the material is agitated by the discharge of compressed air through it from distributors placed in the sides and bottom of the vat.

The gold in the slimes, being necessarily very fine, is dissolved very quickly, and the percolation can generally be started two hours after charging the leaching vat. When a portion of the gold is coarser than usual, or when it exists in the form of amalgam, it is found that the agitation caused by the discharge of compressed air during the leaching greatly accelerates the dissolution.
The novel features of the process consist of upward filtration, and agitation by means of discharges of compressed air through the mass.

The results of a number of small tests, on the finest slimes obtainable, have proved most satisfactory, and more extensive trials will soon be undertaken. One air-compressor would be sufficient for a number of leaching vats; and the author is confident that the process is capable of very wide and useful application.

Agitation with compressed air has now become an important feature in the treatment of slimes and concentrates throughout Australia, New Zealand, and America. It possesses many advantages over mechanical agitation, and, besides, causes a better aeration of the pulp.

Acting on a suggestion thrown out by Mr. C. Wichmann, the author obtained from Mr. E. G. Banks, metallurgist to the Waihi Gold Mining Company, the results of a number of experiments, showing the relative values of the different degrees of fineness of the pulverized ore. These results are very instructive.

### Experiments showing degree of Fineness and relative Values to which “Martha” Ore was reduced by stamping through 30-mesh Screens.

<table>
<thead>
<tr>
<th></th>
<th>Value per Ton</th>
</tr>
</thead>
<tbody>
<tr>
<td>A. 0·36% remained on 30-mesh (900 holes)</td>
<td>£4 2 6</td>
</tr>
<tr>
<td>B. 2·16%</td>
<td>3 18 10</td>
</tr>
<tr>
<td>C. 9·29%</td>
<td>3 11 2</td>
</tr>
<tr>
<td>D. 25·72%</td>
<td>4 9 0</td>
</tr>
<tr>
<td>E. 74·28% passed</td>
<td>5 7 1</td>
</tr>
</tbody>
</table>

### Similar Experiments with 40-mesh Screens.

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<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>A. 0·3% remained on 40-mesh (1600 holes)</td>
<td>3 12 7</td>
</tr>
<tr>
<td>B. 7·8%</td>
<td>3 7 11</td>
</tr>
<tr>
<td>C. 14·7%</td>
<td>3 11 4</td>
</tr>
<tr>
<td>D. 85·3% passed</td>
<td>3 19 2</td>
</tr>
</tbody>
</table>

### Experiments showing value of Dust rising from Stamp—Dry-crushing.

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>A. Dust obtained from floor of mill to 10 ft. high</td>
<td>£6 16 4</td>
</tr>
<tr>
<td>B.</td>
<td>7 2 4</td>
</tr>
<tr>
<td>C.</td>
<td>7 3 1</td>
</tr>
<tr>
<td>D.</td>
<td>7 19 11</td>
</tr>
<tr>
<td>E.</td>
<td>8 13 8</td>
</tr>
<tr>
<td>Average value of ore from which this dust came</td>
<td>4 1 4</td>
</tr>
</tbody>
</table>
These experiments proved conclusively that when using 30 and 40-mesh battery screens the highest values were carried by the finest sands and slimes.

**FILTER-PRESS METHODS.**

This process has been revived in New Zealand during the past few years, but it is in Western Australia that it has received its widest and most successful application. The scarcity of water and the clayey character of the oxidized ores of that tropical country presented a difficult problem to the metallurgist and chemist. Compared with the decantation process, a higher average extraction is being obtained, with a lower consumption of water; but the assay value of the residues is higher than in South Africa, where it has been reduced as low as 9½ grains of gold per ton. A consideration of the practice shows that the higher extractions at Kalgoorlie are due to the higher grade of the slimes treated.

1. For some years the dissolution of the gold by the cyanide was effected in the filter-press, the solution being forced through the cakes of pulp. This practice was used mostly for the treatment of slimes from oxidized ore. It can only be employed with economic advantage for slimes containing gold that is rapidly dissolved. Artificial aeration of the slimes between the application of the different charges of cyanide solution was a special feature of this method of filter-press practice.

2. At the present time the filter-press is commonly used as a filtering machine only. The thickened pulp, containing about 1 part of dry slime to 1 part of water, is treated with cyanide solution in agitators in which the agitation is effected either by mechanical stirrers, compressed air, or a centrifugal pump, or by two or more of these together.

The slime pulp is generally thickened by first adding lime to it and then passing it through a series of spitzkasten. At some mills the pulp from the spitzkasten is further thickened by settlement for twenty-four hours in a settlement tank, the clear solution being removed by decantation with a floating syphon.

The pulp can also be thickened by vacuum suction in large agitator vats provided with a vacuum partition on the bottom, or around the sides, or on both sides and bottom. The vacuum partition is connected with an air-chamber into which the solution is drawn. By this method of thickening the author has been able to reduce the moisture in slimes to 30 per cent. of the residual pulp. The thickened pulp is then sluiced into a second agitator vat with a jet of cyanide solution passing through a small nozzle.
The Treatment of Slimes.

When the gold is dissolved the pulp is (a) either passed into a monteju or pressure tank, from which it is forced by air at a pressure of 50 to 90 lbs. to the square inch into filter-presses for the removal of the cyanide solution and washing; or (b) the monteju is dispensed with, and the pulp pumped to the filter-press with a plunger pump, either direct from the agitator or from a collecting vat. The second method of press-filling is now in common use in Western Australia. It effects a considerable economy in filling, since in the older method it was necessary to use one monteju of high-pressure air at each filling of the press, besides additional air for agitation.

The obvious objection to the method of thickening by spitzkasten followed by cyanide treatment is the accumulation of solution which takes place. In mills crushing with cyanide solution this objection is not so serious, as a large volume of weak solution is required for the mortars.

(3.) At one of the Waihi mills and elsewhere the spitzkasten thickened pulp is first passed through the press to expel the water, after which the cakes are agitated with cyanide solution, pressed a second time and washed.

Kalgoorlie Filter-Press Practice.—Details of Process.—The following details of the process as practised in Western Australia are extracted from an instructive paper recently published by Mr. Donald Clark, B.E., Director of the Bairnsdale Mining School in Victoria.*

The ore at Kalgoorlie, as a rule, is dry-crushed, Krupp's ball mills being mainly used; after crushing, it is wholly roasted. Several types of furnaces are on the field, including the shaft furnace (similar to those at Mount Morgan), Brown's, Ropp, Holthoff-Wethy, and Edwards' mechanical furnace. It is noteworthy that the last is most valued, on account of its lowness in cost and its perfection in roasting.

The analysis of typical ore from the Lake View Consols mine is appended:

<table>
<thead>
<tr>
<th></th>
<th>Before roasting</th>
<th>After roasting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Insoluble</td>
<td>62.80</td>
<td>66.50</td>
</tr>
<tr>
<td>FeS</td>
<td>8.27</td>
<td>Fe₂O₃</td>
</tr>
<tr>
<td>CaCO₃</td>
<td>13.89</td>
<td>CaO</td>
</tr>
<tr>
<td>MgCO₃</td>
<td>7.37</td>
<td>MgO</td>
</tr>
<tr>
<td>SO₂</td>
<td></td>
<td>6.58</td>
</tr>
<tr>
<td>S (undecomposed)</td>
<td>11</td>
<td></td>
</tr>
</tbody>
</table>

These results show that the ore is not a complex one, since it

* Clark, The Australian Mining Standard, 5th Dec. 1901.
consists of the usual gangue and a comparatively small percentage of pyrites. The presence of telluride of gold gave rise to the greatest difficulties in the treatment of this ore.

The method finding most favour is to pass the whole of the roasted ore into a hydraulic separator, the fine gold and slimes flowing over the top, while the coarse gold, the partly roasted sulphides, and the coarse sands are drawn away at the bottom. The slimes are led into spitzkasten, where 50 per cent. of the water flows off at the top clear, and the mud, containing 50 per cent. of dried slimes, is run into agitators, where it is agitated with cyanide solutions and afterwards filter-pressed. The coarse sands from the hydraulic separator are run over copper plates, then over percussion tables, where the coarser concentrates are extracted. The sand which escapes from these is run into a vat, and there treated in the usual way with cyanide solutions. The concentrates from the table are ground to slimes in an amalgamating pan; the slimes are sent to the agitators, thence to the filter-presses.

The Dehne filter-press is the favourite press in use and has given great satisfaction. The Martin press has lately been placed in the market and is worked as follows:—

The slimes are forced in through the slime valve and passage, and thence find their way through the ports or side openings of the open frames, which they fill up, the liquor escaping by the drain cock.

When the water ceases to run, these plates are full of fairly caked slime. The cocks are then closed. Solutions for cyaniding or washing, as the case may be, are then forced in through the solution channels. These find their way into the solution plate, and are forced from the corrugated plate through the perforated one, then through the slime cake, through the filter-cloth, the perforated plate, and down the corrugated surface of air-plate, whence they are led away to the zinc boxes. It should be stated that any imprisoned air is first got rid of by opening the air escape cocks, and displacing the air by the solution. When the liquid starts to run through the air-cock, the solution plate is closed.

When cyaniding in presses, the escaping water is clear enough to be led back to the separators; the cyanide solutions are forced through at a pressure of 90 lbs. per square inch until the gold is extracted. It takes 90 minutes to fill, 90 minutes to extract the gold, and 90 minutes to empty three such presses. In order to displace any wash-water or solution, a current of compressed air is turned on to the solution passage for a few minutes, and the press is ready for opening. The cakes or slabs of slime are usually 3 ft. square, and a 50-cake press 2 ins. thick contains 80 cubic
feet, or about 3 tons. Messrs Martin & Co. are now making presses 42 in. square for 3-inch cakes, the 50-cake press holding nearly 6 tons. When the presses are used in conjunction with the agitators, as many as from 10 to 12 charges may be put through in twenty-four hours, or, in other words, as much as 60 tons per day. According to Mr. J. Moss, the cost for filter-breaking without any grinding power is 10s. 10½d. per ton, the average residues being 1·95 dwt. per ton. The total cost of treatment from the mine to the tailings dump, allowing for first cost in labour and supplies and second cost in repairs and renewals, is put down at 35s. 9·4d. per ton.

The Dehne machines can, if necessary, be constructed of gun-metal or wood, with all the channels and valves lined with material suitable for withstanding the corrosive effects of any particular material.

Details of the Dichtl and Riecken slime processes, both of which are in operation at Kalgoorlie, will be found in Chapter XIII.

THE MOORE FILTRATION PROCESS.

It was first installed at the Golden Gate mill, Mercur, and discarded. It is now in operation at Lundberg, Dorr & Wilson mill, South Dakota, and at least in one mill each in Colorado, California, and New Zealand, in a variously modified form.

For the details of the Moore process, as practised in South Dakota, the author is indebted to a paper by C. H. Fulton.*

"At the Lundberg, Dorr & Wilson mill, the Moore slimes-process is used on the slimes. In this process there are three rectangular tanks, 15 ft. long, 7 ft. wide, and 5·5 ft. deep; the first has a double-hopper bottom, the sides inclined 45° in order to aid in the circulation of the pulp, and to discharge more readily some heavy slimes which may fail to adhere to the filter-frames and fall to the bottom of the vat. There is a set of 34 frames, each 4·5 ft. by 6 ft. in area, made of 2-in. material. The filter is made of 18-oz. duck. Both sides of the frames are effective as filters, and the total filtering-area is therefore 1836 sq. ft. The interior of the frames is connected with a pump and a compressor, which furnish suction or pressure respectively. The suction is equivalent to 18 in. of mercury. The set of frames is suspended from a hydraulic crane which transfers it from one vat to the other.

"The method of treatment is as follows:—The slimes, after agitation by a centrifugal pump in an 8-ft. sheet-iron cone,

are passed to the first tank, the frames are immersed in the slimes and the suction is started. A coating of slimes deposits on the filters, and the clear solution is discharged by the pump. At the expiration of from 40 to 65 min. the layer of slimes on the filters has accumulated to the thickness of an inch, which is equivalent to a load of 4 tons, the set of frames is then lifted out with its adherent load of slimes (suction meanwhile being continued) and immersed in the next tank filled with barren solution, which is sucked through the slimes for 40 min. The frames are then transferred to the next tank, which is filled with water. In like manner the wash-water is sucked through the slimes for 40 min., and the frames are then lifted out and carried to a position above the discharge-hopper. The suction is changed to pressure, which causes the slimes to peel off into cars placed below the hopper. A little scraping has to be done in order to clean the frames.

"There being no data yet available concerning the working results of the Moore process, a comparison with the decantation-process cannot be made. The Moore process, however, discharges drier slimes, those at the Lundberg, Dorr & Wilson mill containing from 34 to 36 per cent. moisture, as compared with 50 per cent. of moisture, which is the usual proportion of the slimes from the decantation-process.

"At the mills where the upper layer of slimes is pumped off, as described for the Hidden Fortune mill, the slimes are discharged with from 46 to 47 per cent. of moisture. At some of the mills of the district, where the tailings must be confined within narrow limits, and hence cannot flow freely to waste, considerable trouble is experienced from the high content of moisture in the slimes-tailings. The slimes-tailings from the Moore process are much more easily held in check."

Sun-drying Slime Process.—At the Try Fluke mine at Kuaoetunu in New Zealand, the slimes were dried in the sun, broken up, mixed with sands in the proportion of one part of slimes to two parts of sand, and then treated with cyanide with satisfactory results.

This slime process is in use in India, and in the United States at the Dexter plant, at Tuscarora in Nevada, where 1 part of dry slime is mixed with 1 part of sand and leached with a 0·20 per cent. solution of cyanide for three to five days, the solution running in at the top as fast as it runs out at the bottom until an adequate extraction has been obtained. It was tried for the treatment of slimes collected in dams at the Rand, but without success.*

CHAPTER VIII.

THE CYANIDE TREATMENT OF CONCENTRATES.

Pyritic concentrates may be successfully treated by the following methods:

Raw Concentrates.
(a) By percolation.
(b) By agitation.

Roasted Concentrates.
(a) By percolation.
(b) By agitation.

CYANIDING OF RAW CONCENTRATES.

By Percolation.—The chief obstacle to successful treatment by this process is the tendency of the raw concentrates to pack so tightly as to prevent percolation of the cyanide solutions, even when aided with an artificial vacuum under the filter-cloth.

When treating low-grade pyritic concentrates at the Moanataiari mine in 1899, the author found that the best results were obtained when the concentrates were mixed with sand in the proportion of 5 parts of sand to 1 part of pyrites. At the Rand, where large quantities of concentrates have been treated by percolation, a smaller proportion of sand to pyrites, about 3 to 1, is found to give a freely leachable charge. Obviously, the proportion of sand to pyrites will depend on the fineness of the pyrites.

The prime factors to ensure success are time, sufficient aeration, and patience. Aeration is effected by turning over the charge in the vat every second or third day. Time and patience are required to continue the application of cyanide solutions and frequent turning over of the charge until a satisfactory extraction has been obtained.
The method of treatment does not differ materially from that followed in the treatment of sands, except in the frequent application of strong cyanide solution. The succession of operations is as follows, but it may be modified with advantage to suit local peculiarities of material:

1. Water-wash.
2. Alkaline-wash.
3. Weak cyanide solution, say 0.10 KCy. (This solution is generally destroyed.)
4. Strong cyanide solution, say 0.25 KCy. This solution is returned to vat six or ten times, with short aeration between application of solutions.
5. Charge turned over in vat.
6. Strong cyanide solution (0.25 KCy) applied for two days as before.
7. Charge turned over in vat.
8. Strong cyanide solution as in (6) if necessary.
9. Two washes of weak solution, 0.05 KCy.
10. Final water-wash, or two if required.

In all cases the strong solution must be applied until the extraction is satisfactory. From time to time the solution should be tested for alkaline sulphides.

Mr. C. M. P. Wright gives the following useful details of the cyaniding of concentrates by percolation at the Choukpazat gold-mines: *—

"The concentrates consist of 30 to 40 per cent. sulphides and 60 to 70 per cent. coarse sands: of the sulphurets, other than iron pyrites, 5 per cent. consists of franklinite, galena, chalcopyrite, and a very little altaite. Of these minerals the franklinite is by far the richest, assay from 7 oz. per ton and upwards; the galena holds practically no gold, and the chalcopyrite and iron pyrites vary from about 18 dwt. per ton to 2 oz. per ton, depending on the general value of the ore.

"Our mode of treatment is simple. After an alkaline or plain water-wash follows a weak solution wash (catch-sump strength 0.10 per cent. to 0.12 per cent. KCN), and then nine washes of 0.3 per cent. strength: the contents of vat are carefully turned over; and 0.3 per cent. solutions follow until for two successive days the effluent comes down to 0.26 per cent., when the treatment is considered complete. Two more solutions, catch strength of strong solution sump, usually 0.25 per cent., and followed by two weak solution 0.07 per cent. washes, and by a final water-wash, or two if required.

* Wright, Inst. Min. and Met., London, 20th Nov. 1902
"Every solution and wash that passes through the percolation vat passes into the same sump through the same zinc box. A careful check is kept of cyanide used, of solutions taken from weak solution sump, and of strong solution thrown on to tailings vat so that the level of the special sump is kept constant and at same time the somewhat loaded strong solution is exchanged for a cleaner weak solution.

"The check upon consumption of cyanide and zinc is complete and automatic. Usual tonnage charged monthly, 17 to 20 tons; duration of treatment, 24 days; extraction, 84 per cent. Formerly the zinc in zinc box was made up mid-monthly; this was found to be not necessary and wasteful of zinc. This zinc box is now carefully packed at the commencement of the treatment and left absolutely untouched till the clean-up: it is an eight compartment 1,000-ton per month box, and each compartment is filled."

Mr. Wright states that owing to the presence of copper in the concentrates, all the zinc became coated with that metal immediately after being put into use. The cost per ton of concentrates treated was 12s. 10d., and the extraction 84 per cent.

Roasting previous to Cyaniding.—With pyritic concentrates and even high grade pyritic ores, a higher percentage of the gold contents can always be obtained by roasting previous to cyaniding, and in the Cripple Creek camp of Colorado and Kalgoorlie in Western Australia this process is largely adopted for the treatment of sulpho-telluride ores.

By roasting, the cyanide solutions are kept freer from soluble salts than when treating raw ore; a higher percentage of gold can be extracted at a smaller cost; the time of treatment is shorter; and it is found that in the case of clayey ores, roasting causes dehydration, thereby rendering them porous, and making filtration comparatively easy.

The author found that Moanataiari concentrates, which yielded only 30 per cent. of their values in the raw state, yielded, when roasted, 90 per cent., with a smaller consumption of cyanide.

In order to ensure success it is essential that the sulphides should be subjected to a "dead" roast.

A quick and satisfactory test to determine if the ore is dead roasted, and amenable to cyanide treatment, is described by Wallace Macgregor as follows * :

"Take samples of the roasted ore at discharge end of furnace, cool, take from 100 to 250 grams, place in a beaker with 200 cc. of water; stir this by shaking for about a minute, then filter into

a beaker or flask, and to the filtrate in the beaker add a small quantity of cyanide solution made up to the same strength as that used in the regular work of the plant. It is best to add the cyanide solution slowly and carefully, noting the result. If no cloudiness at all appears the ore is dead roasted, or at least well fitted for treatment by cyanide solution, and the consumption of cyanide will be normal.

"If a brown coloration takes place there are still some soluble salts of iron left in the ore, which will cause a somewhat higher consumption of cyanide, and may lead to a precipitation of compounds of ferro-cyanide in the zinc boxes. If, on addition of cyanide to the filtrate, a blue coloration, soon becoming a greenish blue precipitate, is formed, then the ore is very badly roasted, and one may look for a high consumption of cyanide, and the circulating solution will be made foul."

As to the economy of roasting ores, that must be determined for each individual case. There are oxidized ores and tailings from stamp mills which pay a fair profit by direct treatment with cyanide. These ores and tailings may be so low-grade that roasting would be out of the question, although if used it would increase the extraction of the gold on subsequent treatment with cyanide by probably 10 per cent. or more. There are cases of heavy sulphur-rettled ores and concentrates where roasting interferes with the extraction of the gold, and causes a very heavy consumption of cyanide.

The advisability of roasting ores previous to cyaniding should be carefully determined in each case, both by laboratory experiments and small plant tests, where 1 to 10 tons of dead roasted ore could be worked.

The main features of the cyanide process used for the treatment of the sulpho-telluride ores consist in drying the ore; dry-crushing in stamps, Griffin or other mills; roasting the pulverized ore in furnaces; amalgamation in pans; cyaniding sands, if any, in vats by percolation and slimes in agitators; filter-press separation of solutions from slimes, and final washing of slimes.

A detailed description of the plant and process used at some of the leading Kalgoorlie mines will be found in Chapter IX.

In the Ohinemuri Goldfields of New Zealand, the gold occurs in an extremely fine state, and hitherto dry-crushing and cyaniding have been used, the objection to wet-crushing being the difficulty experienced in treating the slimes which were always high grade.
CHAPTER IX.

LEACHING BY AGITATION.

The first attempt to introduce the cyanide process on a working scale, for the recovery of gold and silver from their ores, was made by the Cassel Gold Extracting Company, of Glasgow, at the New Zealand Crown mines, Karangahake, in 1889. The operations were under the supervision of Mr. J. McConnell. In the first plant, agitation formed a prominent feature, but in later years leaching by percolation became the more general and favourite method of treatment.

The new and extensive cyanide works of the Crown Mines Company, besides a percolation plant of twenty-four tanks, contained an agitation plant consisting of sixteen wooden tubs, or barrels, fitted with revolving paddles. The agitators were seldom used, preference being given to percolation.

At the concentration plant at the Sylvia mine, Thames Goldfield, an agitation plant was erected by Dr. A. Scheidel, in 1891, consisting of three agitators, 6 ft. in diameter and 6 ft. deep; three vacuum-filters, together with the necessary solution tanks, zinc extractors, and other appliances for cyanide treatment.

The ore was heavily charged with iron pyrites, and occasionally a small proportion of zinc-blende, argentiferous galena, and copper pyrites. It was wet-crushed in a 10-stamp Californian battery, classified in pyramidal boxes, and subsequently concentrated in jiggers, slime-tables, and buddles.

The concentrates of four grades were afterwards subjected to cyanide treatment by agitation. The extraction from the best slimes is said to have amounted to 96·45 per cent. of the gold, and 94·59 per cent. of the silver. The average extraction from all classes of concentrates amounted to 82·67 per cent. of the assay value.*

The strength of the cyanide solutions varied from 0·5 per cent. to 1 per cent., and the time of agitation from six to twenty-four

* The Cyanide Process, by Dr. A. Scheidel, p. 79.

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hours. About 300 tons of concentrates were treated, and yielded over £10,000 value of bullion, but the cost per ton is not given.

The extraction is said to have been satisfactory until an excess of copper ore appeared in the concentrates, which rendered them unsuitable for cyanide treatment.

The cyanide plant at the Thames School of Mines, designed and erected by the author, is provided with an agitator which serves a double purpose, being used also as a dissolving vat when ores are being treated by percolation.

The agitator is similar to those used at the Crown and Sylvia works, consisting of an upright tub or barrel provided with a central revolving spindle set in a foot-step at the bottom. At the bottom end of the spindle is fixed a screw, consisting of four paddles or blades. The foot-step, being in the agitator, is subject to great wear and tear, and this forms a most objectionable feature, as it must be continually renewed. This difficulty is easily got over by fixing a hollow cone in the centre of the agitator and placing the spindle in this, the motion being applied from below, as in most grinding and amalgamating pans. An agitator of this kind was erected in 1894 by Dr. Scheidel at the Utica mine, Calaveras County, in California.*

At the Victoria mill of the Waihi Company the treatment of the concentrates is effected by agitation with compressed air in tall vertical cylinders. This method is described at the end of the chapter.

The Actual Extraction by Agitation.—The practice of the author was to charge the agitator with the cyanide solution made up to the required strength, using from 40 per cent. to 60 per cent. of the weight of the ore. The agitator was then set going, about fifty revolutions per minute, and the ore or tailings gradually fed in, until the charge was complete. The agitation was continued until a satisfactory extraction had been effected, which generally took from six to ten hours. Samples for assay were obtained from the agitator by means of a small tin at the end of a stout string. When the extraction was considered adequate, the agitator being still in motion, the stop-cock was opened and the pulp allowed to discharge into a percolation vat, where the solution was drawn off and the residues washed, aided by an artificial vacuum. The cyanide solutions were then passed through the zinc precipitation boxes.

Leaching by agitation possesses many advantages over percolation for the treatment of pyritic concentrates or rich tailings. The dissolution of the gold is much more expeditious, taking hours where percolation requires days; and with suitable material the extraction is always high.

* The Cyanide Process, by Dr. A. Scheidel, 1894, p. 89.
LEACHING BY AGITATION.

On the other hand, agitation requires motive-power, and from the nature of the process the charges must be small, in no cases exceeding a few tons. There is a prevalent belief that agitation causes an excessive consumption of cyanide by decomposition by atmospheric carbonic acid gas; but the author thinks this source of loss much exaggerated, and is certainly much less than it was in the early attempts at cyanide treatment when agitation was prolonged from thirty-six to forty-eight hours continuously.

The author has found, by many trials, that from six to eight hours' agitation is sufficient to effect the dissolution of the gold in the most refractory ores when reduced to a sufficient degree of fineness.

Numerous experiments during the progress of agitation have shown that the greater portion of the gold was dissolved during the first hour.

The rate of the extraction at the different hour-periods, during the treatment of the Monowai sulphide ore, is given below, and will be found instructive:

<table>
<thead>
<tr>
<th>Time of Agitation</th>
<th>Gold Extraction %</th>
</tr>
</thead>
<tbody>
<tr>
<td>After 1 hour</td>
<td>85·4</td>
</tr>
<tr>
<td>,, 2 ,,</td>
<td>88·2</td>
</tr>
<tr>
<td>,, 3 ,,</td>
<td>90·6</td>
</tr>
<tr>
<td>,, 4 ,,</td>
<td>92·3</td>
</tr>
<tr>
<td>,, 5 ,,</td>
<td>92·3</td>
</tr>
<tr>
<td>,, 6 ,,</td>
<td>92·3</td>
</tr>
</tbody>
</table>

From the above it will be seen that the maximum extraction was obtained in four hours. This was a complex sulphide ore, containing sulphides of copper, zinc, iron, and lead. An analysis of the more mineralized portion by Mr. F. B. Allen, M.A., B.Sc., Director of the Western Australia School of Mines, gave the following results:

| Insoluble gangue | .      .      .      .      .      90·15 |
|------------------|--------|--------|--------|--------|--------|
| Copper pyrites   | .      .      .      .      .      3·78  |
| Iron pyrites     | .      .      .      .      .      4·40  |
| Galena           | .      .      .      .      .      0·25  |
| Zinc-blende      | .      .      .      .      .      0·26  |
| Alumina          | .      .      .      .      .      0·13  |
| Water and loss   | .      .      .      .      .      1·03  |

\[100·00\]
The bullion contents of this ore were, gold 1 oz. 5 dwt., and silver 14 oz. per ton. A 0.6 per cent. solution of cyanide was used for leaching by agitation, and the consumption amounted to 0.45 per cent., with an extraction of 92 per cent. A very large amount of copper was dissolved, and, becoming deposited on the zinc, caused much trouble in the precipitation of the bullion. The actual extraction was below 70 per cent.

The following interesting and instructive experiments by agitation with 0.25 per cent. and 0.33 per cent. cyanide solutions on “Martha” ore were kindly supplied by Mr. E. G. Banks, the metallurgist for the Waihi Gold Mining Company at Waihi:

Experiments on “Martha” ore, showing rate at which the precious metals were extracted by a 0.25 per cent. solution of KCy, and amount of KCy consumed.

<table>
<thead>
<tr>
<th></th>
<th>Gold.</th>
<th>Silver.</th>
<th>Saved per ct.</th>
<th>KCy used.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ORE BEFORE TREATMENT,</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1., ,, after 2 hours’ agitation,</td>
<td>0 16 8</td>
<td>3 4 13</td>
<td>10 20</td>
<td>40</td>
</tr>
<tr>
<td>2., ,, 4</td>
<td>0 8 4</td>
<td>2 13 22</td>
<td>50 0</td>
<td>16 5</td>
</tr>
<tr>
<td>3., ,, 6</td>
<td>0 6 14</td>
<td>2 7 10</td>
<td>59 7</td>
<td>26 5</td>
</tr>
<tr>
<td>4., ,, 8</td>
<td>0 4 14</td>
<td>2 4 40</td>
<td>71 4</td>
<td>31 2</td>
</tr>
<tr>
<td>5., ,, 10</td>
<td>0 3 12</td>
<td>1 19 20</td>
<td>78 6</td>
<td>33 3</td>
</tr>
<tr>
<td>6., ,, 12</td>
<td>0 3 12</td>
<td>2 3 1</td>
<td>78 6</td>
<td>33 9</td>
</tr>
<tr>
<td>7., ,, 14</td>
<td>0 3 6</td>
<td>2 2 11</td>
<td>80 1</td>
<td>34 2</td>
</tr>
<tr>
<td>8., ,, 16</td>
<td>0 3 6</td>
<td>2 2 11</td>
<td>80 1</td>
<td>34 2</td>
</tr>
<tr>
<td>9., ,, 18</td>
<td>0 3 10</td>
<td>2 3 3</td>
<td>79 1</td>
<td>33 2</td>
</tr>
<tr>
<td>10., ,, 20</td>
<td>0 3 1</td>
<td>2 4 8</td>
<td>81 4</td>
<td>31 3</td>
</tr>
<tr>
<td>11., ,, 22</td>
<td>0 2 15</td>
<td>2 2 8</td>
<td>88 9</td>
<td>34 4</td>
</tr>
<tr>
<td>12., ,, 24</td>
<td>0 2 11</td>
<td>2 2 11</td>
<td>84 9</td>
<td>34 2</td>
</tr>
</tbody>
</table>

**Fineness of Ore Treated,** 1.5% remained on a 40-mesh screen.

14.5% ,, 60 ,, 26% ,, 80 ,,

Agitation of Raw Concentrates with Compressed Air.—The method of treating the raw sulphurets by the Waihi Company is as follows:—The sulphurets are conveyed from the vanners to the tube-mills in tubs containing a few inches of water to prevent oxidation by contact with the air.

The tube-mills are 18 feet long by 3 ft. 6 in. in diameter and charged with 3½ tons of flint pebbles. Their capacity is about 0.5 ton of concentrates per hour, running at a speed of twenty-eight revolutions per hour.

The slumped concentrates are passed through a series of spitzluten and pyramidal settling-boxes, the coarser products being returned automatically to the tube-mill.

The slumped pyritic pulp is treated in ten vertical cylinders,
fitted with conical bottoms and connected in a series. The cylinders are made of sheet-steel, being 15 feet deep and 6 feet in diameter. Agitation is effected by a jet of compressed air in each cylinder.

For the following particulars of the actual treatment the author is indebted to an interesting article by Mr F. N. Rhodes.*

Cyanide solutions carrying from 12.0% to 3.0% KCy are used in the first three cylinders, 0.6% KCy in the middle five tanks, falling to 0.4 KCy in the last two.

The treatment lasts from three to five days, the pulp being passed successively from one tank to the other, that is, from the strong solution tanks to the weak.

The compressed air pipe is connected to a flexible rubber tube at the top, and by this means can be lowered or raised as required. When the concentrates show a tendency to settle, the tube is raised, but ordinarily it rests on the bottom.

After the dissolution of an adequate percentage of the gold, the pulp from the weak solution cylinders is forced into a filter-press. The residues range in value from 20s. to 28s. per ton, and are stacked on a dump for subsequent treatment.

The extraction is given at 95% with a consumption of 20 pounds of sodium cyanide of 129 to 130 per cent. KCy equivalent. The original value of the concentrates is about £40 per ton, and it is noteworthy that a considerable proportion of the value lies in silver sulphide.

There is little doubt that a saving of cyanide and a quicker extraction could be obtained by the use of a desulphurizing agent such as lead acetate.

CHAPTER X.

ZINC PRECIPITATION AND TREATMENT OF GOLD SLIMES.

The zinc for bullion precipitation is used in thread-like turnings, as this form gives the most surface for the least weight. It should be free from arsenic or antimony, although a little lead is an advantage, as it causes more rapid precipitation by forming a voltaic pair with the zinc.

As a general rule, one cubic foot of zinc turnings will precipitate the gold from two tons of solution in twenty-four hours.

Zinc on which a film of bullion has been precipitated is more active than pure zinc, and it is therefore advisable to replace the zinc dissolved in the upper compartments by moving the zinc forward from the lower compartments, and adding fresh zinc to the latter.

In practice, the cyanide solution is allowed to slowly drain through the zinc in the precipitating boxes. The rate of flow is soon determined by actual experience. It is generally found that 85 per cent. to 95 per cent. of the bullion will be precipitated in the first three boxes.

The solution, after leaving the boxes, should not contain more than six or eight grains of gold to the ton.

Zinc, in the form of zinc-dust and zinc-fume, is in use for precipitation, particularly the former, but neither seems likely to supersede zinc turnings, which possess the important advantage that they afford a continuous method of precipitation, whereas dust and fume have to be applied in charges in solutions collected in separate vats. Furthermore, the zinc extractor-boxes, when once in good working order, require very little attention except at the periodical "clean-up."

The principle of precipitation of gold by metallic zinc is based on the fact that cyanide has a stronger affinity for zinc than for gold, as shown by the following equation:

$$2AuKCy_2 + Zn = ZnK_2Cy_4 + 2Au.$$
ZINC PRECIPITATION AND TREATMENT OF GOLD SLIMES

By the above reaction it will be seen that 1 oz. of zinc should precipitate 6 oz. of gold, but in practice it is found that from 4 oz. to 12 oz. of zinc are required to precipitate 1 oz. of gold. The reactions which take place in the zinc precipitating boxes are at times most varied and perplexing, especially during the treatment of pyritic tailings or acid mineralised ores.

Part of the excessive consumption of zinc is no doubt due to decomposition by free cyanide, as may be ascertained by testing the solution for available cyanide before entering and after leaving the zinc precipitating boxes; but the consumption and consequent loss of cyanide by this cause is much less than generally supposed, and in all cases insufficient to account for the great waste of zinc.

Zinc is soluble in an aqueous solution of potassium cyanide without evolution of hydrogen. In the extractors, the cyanide comes in contact with so extensive a surface of zinc that a large quantity of that metal must pass into solution, but curiously enough the consumption of free cyanide in the extractor does not correspond with the consumption of zinc, and we can only conclude that a process of regeneration takes place in the extractor. It is quite certain that the fouling of cyanide solutions with that troublesome inert alkaline substance, zinc-potassium-cyanide, takes place in the passage of the solutions through the extractors.

It might naturally be expected that zinc would accumulate in the cyanide solutions to a detrimental extent, but this is found in practice not to be the case. The zinc does not accumulate to any extent, a result in all probability due to the action of the sulphides contained in the ore and cyanide which cause its precipitation as a sulphide of zinc.

The precipitation of the gold, doubtless from electro-chemical causes, is always more rapid and complete from moderately strong than from very weak cyanide solutions, but under all circumstances the solutions must be distinctly alkaline to ensure a satisfactory precipitation.

It has been suggested by some chemists that this is due to nascent hydrogen, liberated by the action of the free KCy on the zinc, taking the place of the gold, according to the following equations:

\[ 4\text{KCy} + \text{Zn} + 2\text{H}_2\text{O} = \text{ZnK}_2\text{Cy}_4 + 2\text{KHO} + \text{H}_2 \]

and

\[ 2\text{AuKCy}_2 + \text{H}_2 = 2\text{KCy} + 2\text{HCy} + \text{Au}_2. \]

The liberated hydrocyanic acid is capable of combining with any free alkali present, and thus there would be no loss of the
cyanogen combined with the gold. This reaction is shown by the following equation:

\[ \text{HCy} + \text{NaHO} = \text{NaCy} + \text{H}_2\text{O}. \]

Hydrogen gas is always evolved when gold is precipitated, and the gentle action of the gas bubbles, as they rise to the surface in the zinc boxes, is an indication of satisfactory precipitation.

During the treatment of pyritic tailings at Kuaotunu, the unsatisfactory precipitation of the gold was for some time a source of much trouble to the chemists in charge of the cyanide works, but this difficulty was overcome by making up the strength of the solution before entering the extractor to something like the original working strength.

In practice, this was effected without any extra trouble by simply placing a barrel containing a strong solution of cyanide at the head of the extractor, and allowing a steady drip into the cyanide solution, in the top compartment, which was filled with a filter of sand and gravel. By testing the cyanide solution a few times, the rate of drip to bring it up to the required strength was easily determined.

The author used this method with success in the treatment of cupriferous ores from the Monowai mine in the Hauraki Goldfields, in 1895.

It was found that the dissolved copper was precipitated much more rapidly from a weak solution of cyanide than from a strong one. In order to overcome this, the solutions were made up to the original working strength. This method has now been superseded by the lead acetate pickling process.

In the treatment of slimes by decantation and in filter-press processes, but especially in the former, there are formed large volumes of excessively weak cyanide solutions containing gold. It was a matter of early cyanide experience that zinc precipitation from such weak solutions was very imperfect, in fact far inferior to electrical precipitation.

The discovery was soon made that the lead-couple used for the precipitation of gold from solutions containing copper was also very effective for the precipitation of gold from extremely dilute cyanide solutions, and the practice in South Africa and elsewhere in slimes plants is to pickle the zinc, before use, in a trough containing a 10 per cent. solution of lead acetate, until all the zinc is covered with a black coating. The precipitation effected by this lead-couple is almost perfect, only a trace of gold, as a rule, finding its way into the sump.

At the Camp Bird mines, as described in the following page, a
ZINC PRECIPITATION AND TREATMENT OF GOLD SLIMES. 141

tin-mercury couple is used with very satisfactory results for solutions containing copper, and is stated to give better results than the lead-couple.

It is the practice at some plants to allow a dilute solution of lead acetate to drip slowly into the head of the zinc extractors, but the practice is not to be commended, on account of the difficulty of regulating the rate of flow. Besides, by this method an excessive amount of lead is certain to find its way into the bullion.

It is now the practice when the cyanide contains sulphides to sprinkle lead acetate on the top of the tank. By this means the alkaline sulphide is decomposed, and the cyanide freed from sulphide by the precipitation of lead sulphide.

When copper is present in the solution, it soon covers the zinc with a bright metallic coating, which begins in the lower boxes, and gradually encroaches on the upper ones. When the zinc is coated with copper, the precipitation of the gold is very slow and imperfect. By increasing the strength of the solution to near the working strength, before it enters the boxes, the copper may be largely kept in solution.

When the ore or tailings contain copper, the supplies of fresh zinc should only be added when the strong solution is passing through the extractor. By attending to this much of the copper can be kept in solution, with a correspondingly satisfactory precipitation of the gold.

It should, however, not be forgotten that when there is much copper present in the ore, it becomes imperative to allow the precipitation of the copper with the gold with the object of keeping the copper-contents of the solutions constant. In such a case to keep the copper in the solutions would soon result in their becoming overcharged with copper and thus useless for gold extraction purposes.

If the zinc turnings be placed in a solution of lead acetate, of, say, 10 per cent. strength, they will become covered with a porous coating of lead. This lead-coated zinc, by its electro-chemical energy, will effect the perfect precipitation of the gold, and leave the copper, even from the weakest solutions, unprecipitated. The resulting bullion obtained by this means is, however, always highly charged with lead.

At the Camp Bird mills, Ouray in Colorado, the waste solutions are very weak, of poor value, and contain some copper. The lead acetate method was tried without much success and abandoned in favour of a mercury-couple which is obtained by immersing the zinc in a weak solution of mercuric cyanide until it is coated with mercury. The mercury coated zinc is stated to give a very fair
precipitation, while the mercury is recovered and does not pass into the bullion as is the case with lead.

Experience has shown that ores containing much copper are not adapted for cyanide treatment, firstly, on account of the undue consumption of the cyanide; and, secondly, on account of the difficulty of precipitating the gold in the presence of the base metal; moreover, by continued use, the stock and sump solutions become charged with copper, and thus rendered useless for all practical purposes, such as washing, or forming the basis of working solutions.

Occasionally an inert gritty, greyish-white, porous precipitate of zinc cyanide forms on the zinc in the precipitating boxes. The reactions which lead to its formation have not yet been satisfactorily explained, but, whatever they may be, its presence is always accompanied by loss of cyanide and imperfect precipitation of the gold. This precipitate is seldom seen excepting in the treatment of pyritic ores and tailings. It can generally be prevented by a careful preliminary washing, and treatment with lime instead of caustic soda.

On the other hand, when a too free use of lime is made to reduce acidity, an incrustation of lime will form on the zinc and thus prevent satisfactory precipitation.

In some cases there may be, in the presence of organic compounds, an excessive and injurious evolution of hydrogen. During the treatment of decomposing pyritic tailings at the Great Mercury Cyanide Works, Kuaotunu, the evolution of hydrogen gas was so vigorous that it lifted the zinc out of the precipitation boxes, forming a thick froth. On this occasion the precipitation of the bullion was very imperfect and unsatisfactory, and suggested polarization.

When a scum forms on the surface of the solution in the precipitation boxes, both it and its cause should be removed without delay. In the case of accumulated tailings it will generally be found to be caused by the presence of decomposing organic matter, and the application of an oxidizing agent often exerts a beneficial effect.

In practice the zinc shavings are first placed in the weak solution box, and afterwards transferred to the medium, and thence to the strong. In the weak and medium solution precipitating boxes, the gold becomes plated on the zinc, and less zinc is destroyed than in the strong. The solution from the extractor boxes, containing only traces of gold, is returned to the solution tanks, where, if found necessary, its strength is made up by the addition of cyanide.

The precipitation of the gold from very weak solutions, especi-
ally in the presence of copper, has always been a difficulty with zinc, but of late years success has, to a great extent, been obtained by pickling the zinc, before use, in a solution of acetate of lead, and adding fresh cyanide at the head of the box, as was done by the author in 1895.

This method of precipitation has been in use in the Lydenburg district for over five years, and is now largely in use at the Rand, to assist the precipitation of the gold contained in the extremely weak cyanide solutions, which necessarily accumulate in large volumes in the treatment of slimes.

The following interesting particulars of the operations at the latter place have been given by Mr. T. L. Carter*:

A very important point is the preparation of the zinc. After it has been cut on the lathe it is taken to a trough which contains a solution of acetate of lead of about 10 per cent. strength. The zinc is thoroughly washed and stirred in the solution until it becomes of a dark hue. If it is not thoroughly stirred only the outside of the mass of zinc will become coated, the inside remaining quite bright. An empirical way to ascertain whether or not the solution is strong enough is to look at the zinc, and see if it is sufficiently and thoroughly coated. After thus preparing the zinc it should be placed in the box, and covered with the auriferous solution as quickly as possible, since leaving it in the air seems to affect it adversely.

The next important point is the addition of free cyanide at the head of the box. Twenty pounds of KCy are dissolved in an iron tank, holding about 75 gallons water. This 2½ per cent. solution is allowed to run freely into the auriferous solution entering the box for a period of about four hours, raising the strength of the solution passing through from 0.007 per cent. to 0.025 per cent. When the 20 lbs. of KCy are finished, another 10 lbs. are dissolved as before, and freely run into the box, taking a period of six hours, and raising the solution about 0.007 per cent. higher. By no means should the addition of the cyanide as described be neglected, for it is found absolutely necessary at the commencement to let this free cyanide run into the box. Twelve or fourteen hours after starting a slow drip is allowed to fall into the solution as it enters the box, bringing up the strength of the solution going through the box from 0.007 per cent. to 0.008 per cent., and this is dropped in through the run of the box. The precipitation is almost perfect. On account of the lead present in the slimes, the smelting gave much trouble at first, but, after many experiments, the following flux was found to give the best results:

Borax, ... 60 per cent.
Nitre, ... 19 "
Sand, ... 11.5 "
Soda, ... 7 "

The precipitation of gold by zinc, results in the formation of a double cyanide of zinc and potassium, and the continual use of the same solutions would lead to the belief that the working solutions would in time become charged with zinc salts. In actual practice it is found that this is not the case to any great extent.

Feldtmann states that, under favourable conditions, the zinc-potassic cyanide is of itself capable of dissolving gold from its ores. He considers that the small quantities of alkaline sulphides present in commercial cyanide, or formed by the action of cyanide on metallic sulphides, serve to precipitate a portion of the zinc as the insoluble sulphide.

In cases where ores contain considerable proportions of metallic sulphides, soluble in cyanide solution, sufficient alkaline sulphide may be formed to precipitate a portion of the dissolved gold.

To prevent any loss in this direction, Mr. J. S. McArthur suggested the addition of a solution of some soluble lead or other metallic salt which is known to form an insoluble sulphide in alkaline solutions, and this practice is now general when the cyanide contains sulphides. In this case, however, it would be advisable to avoid an excess of the lead, or other salt, so as to prevent possible complications in the extractor. The exact amount of salt required can be determined in the laboratory.

The Clean-up.—The periodical clean-up takes place once or twice a month. The first operation is to pass a current of clean water through the zinc boxes, so as to remove the cyanide solution, which is injurious to the workmen, often causing their arms to become covered with painful red boils.

The trays holding the zinc are then moved up and down in their compartments so as to allow the fine gold precipitates, and fine particles of zinc, to fall through the sieve and settle in the bottom of the box. The contents of the trays are then placed in a large trough, provided with an easily removable false bottom of finely perforated iron. The zinc is gently teased out and rubbed in this trough, which is partly filled with clean water, and in this manner as much as possible of the adhering gold is removed. After all the gold has settled, as a slimy mass, the water is syphoned off. The gold slimes remaining in the extractor are sluiced through plugholes into the side launder, or into the bottom compartment and collected in a trough. The fine
slimes or precipitates are rapidly settled by the addition of a little powdered alum to the solution.

In large cyanide works the precipitates are dried in a small filter-press or vacuum-filter. The discoloured zinc shavings are now returned to the precipitation boxes, fresh zinc being put in the lower compartments. The gold still remaining on the zinc is recovered at the next clean-up.

The gold slimes are treated in one of three ways, namely:

1. By smelting with suitable fluxes.
2. By sulphuric acid method.
3. By lead smelting method.

The smelting method is the oldest and still has many adherents. It is, however, laborious, slow, and attended with loss of bullion. It is gradually being superseded by the sulphuric acid method, which in its turn seems likely to be superseded by the lead smelting process.

1. **Smelting Process**: — *Roasting the Precipitates.* — The dry precipitates are roasted at a low heat, with free access of air. The object of the roasting is to oxidize the zinc in the slimes, and thus cause it to combine with the fluxes in the subsequent smelting, and thereby leave the bullion as fine as possible.

In Australia and New Zealand cyanide works the roasting furnace often consists of a large flat cast-iron plate, with raised edges. It is built over a small grate or furnace, and a hood of light sheet-iron is placed over the roasting place, so as to carry off the zinc fumes.

The roasting should be conducted at a moderate heat, i.e., it must never rise above a dull red, and the precipitates must be stirred continuously so as to expose fresh surfaces to the action of the atmospheric oxygen. During the early part of the roasting, dense white vapours of zinc oxide are given off, but as the operation advances they are observed to diminish and finally to cease entirely when the reaction is complete. Time, from one to two hours.

Mr. Feldtmann has found that the oxidation of the zinc is facilitated by the addition of a little nitre, say from 3 per cent. to 10 per cent. He suggests that it should be applied to the slimes as a strong solution before the drying, so as to get thoroughly mixed with the whole mass. The nitre not only helps to oxidize the zinc, but is also said to assist the subsequent fluxing by uniting with the zinc oxide, and forming a zincate of potash, which is not so readily reduced by the plumbago of the crucible as the oxide. At many works the nitre is added to the dried slimes in a powdered form; of course less nitre must be
used than is necessary to oxidize all the base metal present, as any free nitre remaining would rapidly destroy the plumbago crucible during the smelting process. Besides rendering the bullion fine, the nitre roasting gives a cleaner slag and greatly hastens the fusion.

When stirring and removing the roasted precipitates, care must be exercised to avoid a loss of bullion in the form of dust.

*Smelting the Oxidized Precipitates.*—The roasted precipitates are now placed on a large, shallow iron tray, mixed with the necessary fluxes, and fused in plumbago crucibles. The following fusing mixtures have always given satisfactory results:

<table>
<thead>
<tr>
<th>Clean Precipitates</th>
<th>Much Zinc and</th>
<th>Very Sandy.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Little Zinc</td>
<td>Little Sand</td>
<td></td>
</tr>
<tr>
<td>Precipitates,</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Bicarbonate of Soda,</td>
<td>6</td>
<td>20</td>
</tr>
<tr>
<td>Borax</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td>Sand</td>
<td>3</td>
<td>15</td>
</tr>
<tr>
<td>Fluor-spar</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

The chief essentials in a slag are fluidity, small bulk, and non-corrosion. The first is conferred by the “fused” borax and the last by the sand.

It is desirable that the fluxes used should be free from moisture, so as to avoid loss due to the escape of steam through the charge.

*Smelting Acid-Treated Slimes.*—Acid-treated gold slimes are necessarily of a basic character, hence an acid flux must be used so as to produce a neutral or non-corrosive slag. Manganese dioxide is now generally added for carrying base metals into the slag. Messrs. E. H. Johnson and W. A. Caldecott have shown that in a state of fusion it is even a more active oxidizer than nitre although it contains much less available oxygen.*

The following is the basis of flux used by these authors, the proportions being varied, within the limits, as varying conditions required:

Slimes, . . . . . . 100 parts.
Fused borax, . . . . 20 to 35 °°
Manganese dioxide, . 20 °° 40 °°
Sand, . . . . . . 15 °° 40 °°

The addition of soda to an already basic material was considered unnecessary. When sulphates are present, a little fluor-spar is found to assist the fusion: a few preliminary trial fusions should be made to ascertain the best proportions of the fluxes.

The authors quoted above state that a matte or base-looking bullion indicates too little manganese dioxide, whilst too much manganese dioxide yields an infusible slag or one containing much silver. It has long been known that manganese dioxide in fusion carries off silver, and for this reason in smelting slimes containing commercial values in silver, it must be used with care, or not at all.

Plumbago crucibles with removable clay-liners are generally used for the fusion of these charges. Crosse uses silicate of soda as a source of silica and clay-lined crucibles instead of removable clay-liners.

In works where large quantities of slimes have to be smelted, Nos. 50, 60, or 70 plumbago crucibles will be required.

*The Actual Fusion.*—The crucible, previously annealed, is placed on a flat brick resting on the bars of the furnace. A priming of borax is then placed in the crucible, and over this a charge of precipitates; fresh additions of precipitates are made as the charge fuses and subsides. When the crucible is two-thirds full, the slag is skimmed off and fresh portions of precipitates added until it is three-fourths full of molten bullion.

The crucible is now removed from the furnace, and the contents poured into ingot moulds which have been previously *well heated* and carefully oiled with the best olive oil. All excess of oil should be wiped out of the mould before pouring the metal.

The melting furnace may be constructed to hold two or three crucibles at the same time. It should be built of the best materials, as the heat required to melt the *slime mixture* is higher than that for ordinary smelting.

At the Langlaagte Cyanide Works the slimes, mixed with the fluxes, are charged into No. 50 plumbago crucibles, and melted in a reverberatory hearth furnace which holds 22 crucibles at the same time. The time required for melting varies from one and a half to three hours, according to the character of the materials and the temperature of the furnace.

The slag resulting from the smelting of slimes always contains a small proportion of gold. It is, therefore, generally pulverized in a single stamper, or in a small mill, and afterwards amalgamated with mercury. In some cases it is sent to the smelting works for treatment.

The ingots of bullion, obtained from the first smelting, are re-melted with borax; and, since gold forms but a very imperfect alloy with zinc, this second melting should be conducted at as low a temperature as possible so as to obtain an approximately uniform bar of bullion.

The zinc slimes generally contain from 30 per cent. to 65 per
cent. of bullion, the fineness of which, after melting, generally varies from 600 to 900.

The clips for assaying should be taken from different parts of the bar so as to obtain a representative sample for valuation, but the dip sample is always the most reliable.

McBride gives the cost of smelting gold slimes at 2d. per ounce of fine gold. At a clean-up giving 718 ounces of bullion, 817 fine in gold, the costs were as follows:

<table>
<thead>
<tr>
<th>Item</th>
<th>Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Borax,</td>
<td>£1 5 0</td>
</tr>
<tr>
<td>Soda,</td>
<td>0 6 7</td>
</tr>
<tr>
<td>Fluor-spar</td>
<td>0 5 6</td>
</tr>
<tr>
<td>Nitre,</td>
<td>0 2 6</td>
</tr>
<tr>
<td>Pots,</td>
<td>1 1 9</td>
</tr>
<tr>
<td>Coal,</td>
<td>1 16 0</td>
</tr>
<tr>
<td>Labour,</td>
<td>1 2 8</td>
</tr>
</tbody>
</table>

\[ \text{Total Cost: £5 19 8 = 2d. per oz. fine gold.} \]

2. **Refining by Sulphuric Acid.**—This method is commonly used in cyanide works in America and South Africa, but has not yet been adopted to any extent in Australia or New Zealand. The acid treatment of the precipitates is a simple enough operation, and was occasionally used by the author for the refining of small parcels. The necessary apparatus consists of shallow wooden tubs, or vats.

The operation is conducted as follows:—Clean water is passed through the zinc extractor, for some time, to remove all traces of cyanide. The precipitates are then removed from the boxes and placed in the first vat, with a sufficient quantity of dilute sulphuric acid. The acid should not be too strong, nor yet too weak; a mixture consisting of ten parts of water and one part of strong acid answers well.

The quantity of dilute acid will depend on the proportion of zinc present in the precipitates. With 50 per cent. of zinc, about six parts of the acid mixture to one of the precipitates will be required; and with very zincy precipitates, from ten to twelve parts.

The mass in the tub should be stirred occasionally, and then allowed to settle. Heat is generated, and large quantities of hydrogen gas liberated by the action of the acid on the zinc.

When the undissolved precipitates have been allowed to settle, the clear liquid should be removed by decantation into a second tub, and thence finally, after an interval, into a third tub. By this means any fine particles of bullion which have escaped in the
first decantation will be secured in the second tub; and that which has escaped during the second decantation will be found as a fine sediment in the third tub.

The bullion should now be washed in the tubes with clean hot water, to remove all base soluble sulphates and any free acid remaining. Then remove the bullion slimes and dry on the vacuum-filter. When dry, subject to an oxidizing roasting on a shallow iron pan, for an hour or so, to oxidize any base sulphates present.

Next, mix with 5 or 10 per cent. of borax glass, according to the amount of zinc oxide still present, and fuse in a plumbago crucible in which a priming of borax has already been placed. As the charge fuses and subsides, fresh portions of bullion should be added, until the crucible is three parts full of melted metal. To permit of this being done the slag can be skimmed off from time to time.

If manganese dioxide is used in the flux it will be necessary to protect the crucible from corrosion by using a clay-liner as described in the preceding pages under the heading devoted to the “Smelting of acid-treated Slimes.”

The bullion is generally from 850 to 900 fine, but with a little extra trouble can be worked up to 950.

With suitable appliances, this process possesses many advantages over the smelting process. It occupies less time, produces finer bullion, and, properly conducted, costs less.

When large quantities of precipitates have to be dealt with, the method of settling the slimes and decanting is too slow and expensive. In this case, the separation of the slimes from the acid solution, as well as the subsequent washings, must be effected in a vacuum-filter, or a filter-press, as practised in America and South Africa.

The filter used for the purpose is a wooden box, two or three feet square. It is provided with a filter webbing, or cloth, of fine canvas or twill duck, resting on a grating of wood, and fixed with slips of wood, so as to be easily detached for washing.

The false bottom, below the webbing, must be 15 in. or 20 in. deep, and provided with a solution gauge, the upper limb of which should be 2 in. below the air-exhaust pipe connected with the vacuum boiler.

Care must be taken to draw off the acid solution and washings by a plug-hole before they rise to the level of the air-exhaust pipe, which is placed immediately before the filter-frame.

When the acid solution is diluted to half its strength before filtering, the webbing lasts for several operations. A Johnston filter-press does the washing well and expeditiously.
Acid Treatment in South Africa.—The following description is an abstract of a paper read by E. H. Johnson before the Chemical and Metallurgical Society of South Africa. It describes the process employed at the Princess Works, where the slimes are submitted to acid treatment before smelting, and will be especially interesting for its figures of costs and metallurgical results:—

At the Princess Works the slimes from the zinc boxes are separated from the solution drawn off with them by the aid of a vacuum-filter. A water wash is passed through until the slimes are free from cyanide. The gross weight of the slimes, including moisture, is then taken, by weighing the buckets of moist slime during transference to a large sheet-iron tray placed alongside the acid tank, to determine the amount of sulphuric acid necessary to destroy the zinc.

Having found the approximate weight of slimes to be treated, sufficient water is run into the acid vat to form, on the addition of the acid, a 10 per cent. solution. One pound of acid for every pound of moist slime gives good results. This would be equivalent to about 1\frac{1}{2} lb. of acid to the pound of slimes, dry. The acid is then added, and the vat closed down tightly.

The stirring apparatus is kept continually moving during the time of feeding in the slimes, which are fed in gradually in the same condition in which they were taken from the filter-vat. It is beneficial to keep up a continual stirring for at least half an hour after the action has apparently ceased.

After all the slimes are in the acid, a jet of water is turned into the hopper to wash down any adherent slimes, and everything that has been used in the cleaning-up of the boxes, etc., is well washed in the same jet during removal. The vat is then filled with water and allowed to settle.

Working with dilute acid, and not heating, a perfect settlement takes place within an hour. When heating with a steam jet, settlement was much more difficult.

The washing is done by syphoning off the clear liquor, and filling the vat repeatedly with water, until the solution is neutral to litmus paper—usually four or five washings. It is well stirred at each refilling by means of a long wooden paddle, a rotary motion being given to the water. This causes the slimes to collect in the middle of the vat, and reduces the risk of loss during syphoning—the syphon being let down at the side. A sample of the washings taken continually during syphoning off showed, on careful assay of a large sample, 13 gr. of gold per ton of solution.

The drying of the resultant gold slime is conducted on an open drying hearth in large cast-iron enameled dishes. The cakes are subsequently broken up and transferred to small sheet-iron trays
in thin layers, and subjected to an increased heat. When cool
the slimes are ground, fluxed, and transferred to the crucible. It
fuses quietly and with but little fume, and normally yields 50 to
60 per cent. of the weight of slime as bullion.

The average fineness of the bullion obtained was, according to
the Work's assays, 821.9 and 819.6, according to London returns.
As a deduction of 2 milliemes is made on 800 bullion, this leaves
an actual difference with London of 0.3 millieme.

The slag, after panning out the prills (of which there is very
little), assays 23 oz. per ton, and one ton of slag has been accumu-
lated in two years for an output of 11,627 oz. of fine gold,
which is equivalent to a little under 0.2 per cent. of the total
gold produced.

The cost of reduction, including acid, is 6·7d. per fine ounce,
made up as follows, taking an actual "clean-up" as a basis:—
Dry weight of zinc-gold slimes, 504 lbs.; dry weight after acid
treatment, 100 lbs.; 672 lbs. of acid at 4½d., £12, 12s. 10d.; 66 lbs.
of borax at 37s. 6d. per cwt., £1, 2s. 1d.; 9 lbs. carbonate of soda
at 2½d. per lb., 1s. 10d.; 9 lbs. fluor-spar at 4d. per lb., 3s.; 5 bags
coke at 8s. 6d. per bag, £2, 2s. 6d.; 1 No. 60 crucible, £1, 7s. 6d.;
total, £17, 9s. 9d.; yield, 620 oz. fine gold, or 6·7d. per fine oz.

From information gathered from reliable sources Mr. P. S.
Tavener states that the cost of sulphuric acid refinement of slimes
in South Africa is as follows *:—

a. Sulphuric acid treatment from 4d. to 8d. per oz. of fine
gold.

b. Smelting cost exclusive of labour and furnace wear and
tear 4d. to 6d.

Approximately this gives a total cost of 1s. per oz. of fine gold
produced.

Acid Treatment in America.—I am indebted to Mr. Godfrey
Doveton, one of my former assistants, for the following par-
ticulars of the acid treatment at the Camp Bird Mills, Ouray,
in Colorado:—

"The gold slimes are refined by sulphuric acid in a wooden vat
12 ft. in diameter and 2 ft. deep, coated with paraffin paint.
They are not subjected to heat, sufficient being generated on the
addition of the acid to the moist slimes. About twelve hours are
occupied in dissolving the zinc, and some forty-eight hours in wash-
ing out the soluble sulphates. The washing is much expedited by
the use of hot water, at least for the first six washings. About
fifteen wash waters are usually required. After the first two
washes, having a high specific gravity, are removed, the time

allowed for settling is one hour. Usually it takes from eight to twelve hours to remove the first two washes, as the slimes do not settle so readily in the heavy acid liquid.

"All the washings are stored in a large settling-tank, which is cleaned up at long intervals. The settling-tank, after a year’s run, yielded bullion worth £20.

"The first two or three wash waters on assay are found to yield from 10 to 20 cents per ton, but subsequent washes contain from 5 cents to a trace,—only apparent on assay of large evaporations.

"The gold slimes are partially dried by filter-press, and are then transferred to a hearth calcining furnace and calcined. The hearth of furnace has an area of 36 square feet. The furnace-doors are securely padlocked, and the furnace serves the double purpose of a safe and calciner.

"From the furnace the slimes are removed as required, and charged into graphite crucibles, and smelted, having been previously mixed with 50 per cent. of a flux composed of the following parts:

\[
\begin{align*}
\text{Soda,} & \quad \ldots \quad 2 \text{ parts.} \\
\text{Sand,} & \quad \ldots \quad 1.5 \text{ } \\
\text{Borax powder,} & \quad \ldots \quad 4 \text{ } \\
\text{Sulphur,} & \quad \ldots \quad \text{as required.}
\end{align*}
\]

Thus 200 lbs. of slimes require 100 lbs. of flux. The charge is smelted with frequent skimming in from ten to sixteen hours for each crucible.

"The slimes contain a considerable quantity of copper, but comparatively little zinc, consequently little or no nitre is used; but we use instead a certain quantity of flowers of sulphur, which readily converts the copper to a matte or regulus, leaving the bullion comparatively fine. The regulus, which by the way is usually from 15 to 25 per cent. of the weight of the bullion, was found to contain, on assay—

\[
\begin{align*}
\text{Gold,} & \quad \ldots \quad 23.4 \text{ oz. per ton,} \\
\text{Silver,} & \quad \ldots \quad 408.0 \text{ } \\
\text{Copper,} & \quad \ldots \quad 60\% \\
\text{Zinc,} & \quad \ldots \quad 3\% \\
\end{align*}
\]

"The slag resulting from the bullion melting is ruby red in colour, and contains—

\[
\begin{align*}
\text{Gold,} & \quad \ldots \quad 15 \text{ oz. } \{ \text{Assayed without removing} \\
\text{Silver,} & \quad \ldots \quad 265 \text{ } \{ \text{shots or prills of bullion.} \\
\text{Copper,} & \quad \ldots \quad 6 \text{ to } 8\% \\
\text{Zinc,} & \quad \ldots \quad 12 \text{ } 16\%
\end{align*}
\]
"The bullion averages about 800 to 850 fine, the impurity almost entirely copper with a little zinc.

"A shipment of 3000 lbs. of slag and regulus mixed was lately sent to the smelter and netted us £120."

3. Tavener Lead-Smelting Method.—This process was first made use of by Mr. P. S. Tavener in August 1899. Since August 1901 it has been in continuous operation at the Bonanza Mines, Limited, Johannesburg, and already the process has been adopted by many leading mines at Johannesburg. The adoption of the lead-smelting of gold slimes marks a notable advance in cyanide practice. The old smelting process is laborious, and always likely to entail serious losses of gold where large quantities of slimes have to be handled, while the sulphuric acid method is cumbersome, slow, and costly.

The obvious advantages of lead-smelting compared with the sulphuric acid process may be briefly summarized as follows:—

a. Saving of cost per oz. of fine gold produced.
b. No by-products.
c. Less liability to loss in handling slimes.
d. More gold actually produced from a given weight of slimes.

The essence of every metallurgical process is its cost, and judged by this principle, lead-smelting possesses a marked advantage. In South Africa, where the sulphuric acid method has, perhaps, its greatest application, the average cost, according to Tavener, is not less than one shilling per oz. of fine gold produced. The lead-smelting process costs threepence per oz. of fine gold, including all charges, which means a saving of ninepence per oz. compared with the sulphuric acid method.

Thus the lead method at threepence per oz., in a mine producing 2500 oz. fine gold per month from cyanide works, would effect a saving of costs amounting to £93, 15s. per month, equal to about £1000 a year.

The Tavener lead-smelting process is cheap, rapid, and efficient compared with other processes, and with some modifications in practice is likely to be universally adopted.

The following working details of lead-smelting as practised at the Bonanza mine are extracted from a paper read by Mr. Tavener before the Chemical and Metallurgical Society of South Africa, October 1902.*

Mr. Tavener says that he can best describe lead-smelting by comparing it to a scorification assay conducted on a large scale, for the reason that the zinc slimes are melted and the gold

recovered in lead bullion. The lead bullion is then cupelled, or to use a better term, refined.

"The 'clean up,'" he continues, "is conducted in the ordinary way, with the exception that all the precipitate is at once pumped from the 'clean-up' tub into the filter-press. The fine zinc which remains at the bottom of the 'clean-up' tub is heaped up on one side and allowed to drain for about half an hour, and is then ready for the smelting room. The filter-press is cleaned out and the cakes taken in their moist condition to the furnace, and there both slimes and fine zinc are put in trays into a drying oven, and sufficient time is allowed to warm through, fifteen minutes for each tray being sufficient. Care is taken to keep the fine zinc separate from the filter-press slime, and on no account should they be allowed to get mixed.

"In charging the furnace, the slime is first dealt with. After warming in the drying oven, it is at once rubbed through a sieve, four holes to the linear inch, and then roughly weighed for fluxing, the necessary fluxes having been previously mixed.

"The slime is mixed with the fluxes and passed through a sieve to ensure thorough mixing. It is then shovelled direct into the furnace. When all the filter-press slime is fluxed and charged, the fine zinc is dealt with in the same way, and put into the furnace on the top of the slime in order to prevent loss by dusting, and also to have the greater portion of litharge present on the top of the charge. The fluxes used are residue assay slag and commercial litharge. The former costs nothing, for unless used in this manner it would be thrown away."

Discussing the question of fluxes, he says:—"I have found that the following, with little variation, will give satisfactory fusion and clean slag:—Slag, 25 to 30 per cent., made up of 10 per cent. assay slag; the balance, equal quantities of old slag and scalings from the pots of previous smelt. In the event of the lead-smelting method being adopted, I should like here to point out the advantage of storing the slag now being obtained from crucible smelting, since in the lead process its gold contents are converted into bullion free of cost. If it were not treated, clean slag would have to be used. During the last year I have been able to deal with several tons of this material assaying over fifty ounces fine per ton, left behind on the mine after the Boer occupation. The same applies to clay liners and anything else carrying gold, now termed by-products. With a lead-smelting furnace the word by-product is forgotten, since none is obtained."

The quantity of litharge to be used will depend on (a) value of slimes, (b) weight of charge in furnace, and (c) the percentage of gold required in the resultant lead bullion.
Mr. Tavener gives the following proportions of fluxes for gold slimes:

- Gold slimes, . . . 100 parts by weight.
- Litharge, . . . 60 "
- Assay slag, . . . 10 to 15 "
- Slag previously used, . . . 10 to 15 "
- Sand (SiO₂), . . . 5 to 10 "
- Sawdust, . . . 1% of weight of litharge.

For fine zinc he uses the following proportions:

- Fine zinc, . . . 100 parts by weight.
- Litharge, . . . 150 "
- Slag, . . . 20 "

The products of different mines will necessarily vary, and the proportions to give a clean, well-oxidized slag can easily be determined by experiment on a small scale with assay crucible tests. Mr. Tavener, however, mentions that considerably less assay, or other slag, will effect a good fusion in a reverberating furnace than in a crucible, and he states that it would be safe to use 30 per cent. less slag when smelting in the reverberatory than was found necessary in the crucible trial.

The quantity of litharge should be so proportioned that the lead bullion should not carry more than 8 per cent. of gold, or 10 per cent. at the maximum. It was found preferable to make a larger quantity of lead than to have it too rich in gold contents.

No reducer is used with the fine zinc, which is relied on to reduce sufficient lead, leaving an excess of litharge in the slag to ensure success. With the gold slimes charge, 1 per cent. of sawdust is added on the weight of litharge, but if a larger proportion of litharge has been used, then from 1 ½ to 2 per cent. of sawdust is necessary.

The furnace work is described as follows:

"When the entire charge of zinc and slime is in the furnace, it is banked up from the sides to the centre so as to avoid the possibility of particles remaining on the sides above the slag level as the charge reduces and settles down. A covering of litharge is spread over the surface, and on this again a light covering of easily fusible slag is spread. The furnace is charged the day previous to smelting, and one of the night-shift men lights a slow fire about 3 a.m., which serves to dry the charge. At 5 a.m. the damper is opened and the fire urged, and in half an hour the furnace is at a smelting heat. By 9 or 10 a.m. the charge is reduced, then sweepings from cyanide works, smelting room, or
any slag requiring resmelting, is added and is quickly absorbed in
the molten bath. When all this has been fed in and melted, and
the slag become fluid, it is well stirred with a rabble, and sawdust
is thrown in to reduce the excess of litharge in slag. This opera-
tion is repeated until the slag, which remains on the rabble when
withdrawn from the furnace, is judged by its appearance to be
clean. The slag is now run off into pots through the slag-door,
the level of which is 4 in. above the centre of the lead bath—a
bath of 12,000 oz. of lead bullion almost occupies this space.

"Before filling the furnace, the slag-door is built up about
12 in. by placing flat cast-iron plates, ½ in. thick, bedded in fire-
clay, one on top of another, and in front of these plates a bank of
fire-clay is also made. In order to run the slag off, all that is
necessary is to break away this bank, plate by plate, and so allow
the slag to flow over into the pot. When the pot is full it is
wheeled away and another is put in its place. The filled pot is
run outside, and after standing a minute or two, is tapped, and
the molten slag allowed to run out on the ground to cool; that
which remains on the sides and the bottom of pot is brought back
for further use. When no more slag will flow from the furnaces
owing to the bath being down to the level of the slag-door, it is
waved off by rabbling. At first sight it would appear difficult to
draw this remaining slag off without dragging out some lead, but
a very little practice enables it to be done so closely that there is
little but a thin skimming of slag remaining. In the event of a
little lead being pulled out into the pot, it is recovered from slag
pots. It is for this reason that the pots are tapped about 2 in.
from the bottom. By opening the fire door this last skim on the
lead bath quickly thickens. A shovelful of lime is thrown in to
assist. This skim is easily pulled off, and of course is held over
until next smelt. By this means a clean surface of lead is exposed,
and any zinc present would be quickly got rid of, for at this stage
the lead is at a bright red heat, and the free access of air due to
the open fire door quickly oxidizes it. So far, lead recovered by
this method has always been clean and soft, a proof that no zinc
could be present, since one per cent. of zinc gives lead a distinct
silvery colour, and makes it so hard that it cannot be rolled. The
lead bullion is tapped by driving a ½ in. steel bar, tapered to a
point, into the tap-hole, which is closed with a fire-clay plug. The
lead is run into an iron trough, which conveys it to the moulds
placed together on the floor."

Before tapping the furnace the lead-bath is well stirred, and a
sample is taken out with a ladle and granulated.

The cupelling or refining of the lead bullion is next described,
with particular details and some useful hints on the making of the
bone-ash test and regulation of blast. The process differs only in minor details from the usual operation of lead-refining, and need not be specialized in this work.

The costs at the Bonanza, Limited, for stores for cupelling and smelting for four months (June–Sept. 1902) were as follows:—

<table>
<thead>
<tr>
<th>Item</th>
<th>Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coal</td>
<td>£82 13 1</td>
</tr>
<tr>
<td>Coke</td>
<td>9 8 6</td>
</tr>
<tr>
<td>Fire-clay</td>
<td>3 0 0</td>
</tr>
<tr>
<td>Fire-bricks and slabs</td>
<td>17 15 6</td>
</tr>
<tr>
<td>Paper bags</td>
<td>1 17 6</td>
</tr>
<tr>
<td>Lead foil</td>
<td>3 0 8</td>
</tr>
<tr>
<td>Bar iron</td>
<td>0 3 2</td>
</tr>
<tr>
<td>Crucibles and liners</td>
<td>1 15 0</td>
</tr>
<tr>
<td>Caustic potash</td>
<td>0 12 0</td>
</tr>
<tr>
<td>Bone-ash</td>
<td>10 0 0</td>
</tr>
<tr>
<td>Sundries</td>
<td>1 5 2</td>
</tr>
</tbody>
</table>

**Cost per fine oz. recovered.**

£131 10 7 = 2.463d.

Lead loss estimated at 12% on 15,269 lbs. = 1832 lbs., say 1 ton at £20 = 20 0 0 = 373d.

**£151 10 7 = 2.837d.**

Gold output in period, 12,810 oz., fine.

It is claimed that lead-smelting recovers a larger amount of gold than the sulphuric acid method, and this claim seems to be justified by several large experiments on equal weights of identical gold slimes by both processes. In six trials the lead process gave 10.5 per cent. higher recovery than the acid treatment. These results are probably much higher than could be obtained in continuous working; but it is evident that if only one or two per cent. more gold can be recovered, the lead-process has fully established the claims of Mr. Tavener.

In the discussion which followed the reading of Mr. Tavener's paper, Mr. D. J. Williams suggested that the probable loss of gold by volatilization in the cupellation, caused by the presence of such a volatile metal as zinc, might be avoided by first dissolving the zinc-gold slimes in sulphuric acid, washing, drying, and then subjecting to lead-smelting. Mr. Tavener agreed with this proposed modification, and stated that he would rather receive at the furnace acid-treated precipitate, which would render the work easier and less troublesome. He further stated that with zinc present, a
certain amount of experience and skill was required, while with acid-treated material little or no experience was needed.*

That lead-smelting recovers a higher percentage of gold than the acid-treatment seems to be established on pretty conclusive evidence, but it is not quite so easy to point to the source of gold thus recovered; that is, to show where the loss takes place in acid-treatment.

**Refining with Oxygen and Air.**—Russell and Matthiessen † in 1862 made a number of experiments on the oxidation of silver and copper by passing oxygen and air through the molten metals. These were made to ascertain the cause of the vesicular structure in these metals, and not for the purpose of refining.

Donald Clark ‡ in 1904 suggested the use of both air and compressed oxygen for refining gold, but Dr. Rose§ was the first to show by actual experiment that the process was capable of commercial application. In a valuable series of experiments with impure bullion containing various proportions of gold, silver, zinc, lead, iron, copper, nickel, etc., he proved that by the use of compressed oxygen or air the whole of the base metals could be removed except a small residual part of copper, leaving a tough bullion of great fineness. The loss of gold due to volatilization, projection, etc. was extremely low, that of the silver somewhat greater.

The Rose process is based on the principle that while gold and silver are but feebly oxidized at high temperatures, zinc, lead, copper, iron, etc. are readily so, and hence are easily separated from the precious metals.

The bullion to be refined is melted in a clay-lined graphite crucible. A current of compressed oxygen from a cylinder, or of air, is passed into the molten charge through a one-eighth inch clay-pipe similar to that used in the Miller chlorine process.

The gold-zinc slimes are melted without a preliminary acid treatment. When the zinc oxide ceases to come off, a flux of sand and borax is added to the crucible to form a fusible matrix for the oxides of the base metals as they form in the charge.

When refining highly zinciferous gold precipitates it will probably be found that a preliminary acid treatment will be an advantage.

Air when used for refining could be economically and efficiently supplied from a small air-chamber into which air is forced at a pressure varying from 5 to say 20 lbs. per square inch, according

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‡ Australian Mining and Metallurgy, Melbourne, 1904, p. 493.
to the depth of the molten charge. A supply of compressed
is available at almost every modern mine. This method would
ensure a steady current of air, easily controlled.

The experiments made by Dr. Rose seem to show that either
air or oxygen refining is capable of successful application for the
treatment of the most impure mill bullion, at a cost in the case of
air less than that of other refining processes.

Zinc Fume Precipitation.—The method of application is to
agitrate a certain quantity of the fume with the gold-containing
cyanide solution in vats, allow the precipitate to settle, and
decant the clear solution.

The agitation is effected by artificial stirrers, or by compressed
air.

During the past few years this method of precipitation has
been adopted at a number of American mills, among which may
be mentioned the Homestake Mining Company, South Dakota;
the Montana Mining Company, Montana; the De La Mar Mining
Company, of Nevada; and the Consolidated Mercur Company, of
Utah. It has been used with success in New Zealand.

At the Homestake Company's cyanide works "Precipitation is
carried on by means of zinc dust in five tanks 25 ft. in diameter
and 20 ft. deep, built of Californian red-wood. The bottoms are
inclined to one side, where a small sump is provided in order to
drain the tanks perfectly. No gold storage solution tanks are used,
the solutions draining from the leaching vats directly into the
precipitating vats. The precipitation vats are pumped out by
duplex Prescott pumps, the solution going to filter-presses made
in the Homestake shops. The pulp treated by the mill is of a
highly silicious nature, often however carrying some hornblende,
the sulphides being mainly pyrite and pyrrhotite. The extraction
made is in the neighbourhood of 90 per cent., the cost being
between 45 and 50 cents per ton of material treated, probably
nearer the latter figure."

In America the price of zinc fume varies from 25s. to 29s. per
cwt., and the amount of fume used for precipitation varies from
6 to 9 oz. per ton of solution.

Charcoal Precipitation.—At many cyanide plants in Vic-
toria charcoal is being used to precipitate the gold from cyanide
solutions. The solution is passed through a series of barrels
packed with finely broken charcoal, on which the gold is deposited.
The charcoal is afterwards burnt to an ash, and the ash fluxed.
The process is too slow and cumbersome to recommend itself for
use in large plants where hundreds of tons of solution have to be
handled in the twenty-four hours.

* C. H. Fulton, The Engineering and Mining Journal, June 4, 1902.
CHAPTER XI.

THE APPLICATION OF THE PROCESS IN DIFFERENT COUNTRIES.

SOUTH AFRICA.

At the Witwatersrand Goldfields the cyanide process has been conducted on a more extensive scale than elsewhere. The ore there is principally a pyritic, silicious quartz-conglomerate, consisting of rounded or sub-angular pebbles of bluish-grey quartz embedded in a quartzose matrix. The pyrites occur in varying proportions in different mines, but the average is probably not less than 2 per cent. The gold does not exist in the quartz pebbles, but occurs disseminated throughout the matrix or in the iron pyrites.

This ore is locally known as "banket," or almond rock. It is comparatively hard and somewhat splintery, and often contains a small proportion of corundum and clay, which renders it tough and hard, and forms slimy products during the crushing.

Throughout these goldfields the universal practice at present is wet-crushing with Californian stamps, copper-plate amalgamation, concentration or classification of tailings, cyanide treatment of tailings, and chlorination or cyanide treatment of concentrates.

At most of the batteries a 30-mesh screen is used, but in a few cases a finer or coarser mesh is adopted.

The main features of the cyanide treatment at the Witwatersrand Goldfields are much the same as those practised in America and Australia. The general working details are given below in tabulated form. They are the same at all the cyanide plants, with minor differences according to the individual fancy of the chemist or metallurgist in charge of the operations, and the requirements of the ore or tailings.
<table>
<thead>
<tr>
<th>Process</th>
<th>Hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filling vats</td>
<td>12</td>
</tr>
<tr>
<td>Preliminary alkali or water wash, if necessary</td>
<td>12</td>
</tr>
<tr>
<td>Strong solution, 0.2% to 0.4% solution, 1/4 to 3/4 of ore—</td>
<td></td>
</tr>
<tr>
<td>In contact with tailings</td>
<td>12</td>
</tr>
<tr>
<td>Percolating</td>
<td>12</td>
</tr>
<tr>
<td>Dry or air-leaching</td>
<td>4</td>
</tr>
<tr>
<td>Weak solution, 0.15% to 0.2% solution, about 1/4 of ore,</td>
<td>12</td>
</tr>
<tr>
<td>Dry or air-leaching</td>
<td>4</td>
</tr>
<tr>
<td>Two weak cyanide washes, 0.05% to 0.1% washes, each about 1/4 of ore,</td>
<td>12</td>
</tr>
<tr>
<td>Two clean water washes, each about 1/6 of ore,</td>
<td>12</td>
</tr>
<tr>
<td>Discharging vat</td>
<td>8</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>100</strong></td>
</tr>
</tbody>
</table>

The total quantity of solution used, including the water washes, is about equal to the weight of the ore. The quantity of strong solution used varies according to a preliminary washing with a dilute solution has been employed or not. In the former case it is about 25 per cent. of the weight of the ore, and in the latter case about 40 per cent.

The percolation vats are charged with tailings to within a few inches of the top, and their surface is levelled. The strong cyanide solution is then allowed to penetrate the tailings until they are covered. The contents of the vat settle some inches, the amount of shrinkage depending on the depth of the vat and the percentage of moisture in the tailings.

The value of the tailings varies from 12s. to 20s. per ton, and the actual extraction amounts to 70 or 75 per cent., at a cost varying from 4s. to 10s. per ton, according to the size of the plant.

The Jumpers Deep cyanide plant and slime works are among the newest on the Rand, and represent the most advanced and up-to-date practice. The cyanide works of the Waihi Company at their new mill at Waikino are modelled on the latest South African practice and experience; and as they are fully described in the next pages, it will be unnecessary to give further details of South African works and practice, all of which are lucidly described by Mr. John Yates in his recent work on metallurgical engineering on the Rand.*

New Kleinfontein Mine.—In a paper read before the Institution of Mining and Metallurgy, London, * Mr. F. Cardwell Pengilly gives some interesting details of the successful treatment of tailings by the "direct filling process." The plant, he says, consists of 2 slime separators, 19 treatment vats, 4 extraction boxes, and 9 solution tanks. The slime separators are of the ordinary spitzkasten or pointed box style, 6 ft. square at the top and 6 ft. deep. Of the 19 treatment vats, 13 are of 200 tons capacity, and 6 of 130 tons capacity each. Around the top of each vat is a launder, which carries away the overflow into the slimes race. The 9 solution tanks have a total capacity of 636 tons of solution, and each is connected with the three centrifugal pumps worked from a motor that pumps the weak, medium, or strong solutions on to the treatment vats. The pulp or tailings, after leaving the battery plates, is lifted by means of a tailings wheel into a launder, by which it is conveyed to the slime separators. The slimes from the first separator pass into the second, whereby a quantity of fine sand is saved that otherwise would flow away. About 15 per cent. of slimes are eliminated in the separators, and the remaining pulp is run through launders into the treatment vats, which are previously filled with water. In the course of filling the vat, another 10 per cent. of slimes overflows into the slimes race, so that the resulting tailings to be treated in the vat contain but a small proportion of slimes.

The vat is filled with tailings (sand) to within 1 foot of the top, when the stream is diverted into another vat.

Method of Treatment.—Each vat, after being drained of water, is treated with an alkaline solution to neutralize the free acid in the tailings formed by the decomposition of pyritic ores. As soon as the solution draining away is slightly alkaline, the treatment by cyanide solution is commenced. Various strengths of solution are pumped into the vat, each vat receiving during its course of treatment from 200 to 275 tons of cyanide solution. The length of treatment is six days, and the amount of cyanide used is 6 lbs. per ton of tailings.

The following table shows the exact method of procedure adopted in the works in the treatment of a vat of tailings:

### APPLICATION OF THE PROCESS IN DIFFERENT COUNTRIES. 163

**VAT I. CHARGE I. CAPACITY 207 TONS.**

<table>
<thead>
<tr>
<th>Date filling.</th>
<th>Time.</th>
<th>Test.</th>
<th>Tons.</th>
<th>KCy Solution.</th>
</tr>
</thead>
<tbody>
<tr>
<td>October. Wed. 6th</td>
<td>1 p.m.</td>
<td>...</td>
<td>Filling.</td>
<td>...</td>
</tr>
<tr>
<td></td>
<td>2 a.m.</td>
<td>...</td>
<td>Filled leaching dry.</td>
<td>...</td>
</tr>
<tr>
<td></td>
<td>4 &quot;</td>
<td>Acid</td>
<td>Alkaline wash, 20 lbs.; NaHO - used, 20 lbs.; ½ bag lime.</td>
<td>...</td>
</tr>
<tr>
<td></td>
<td>9 &quot;</td>
<td>Alkaline</td>
<td>Preliminary solution, 0·16; KCy, 20 tons.</td>
<td>...</td>
</tr>
<tr>
<td></td>
<td>2 p.m.</td>
<td>...</td>
<td>0·01 changed to weak box.</td>
<td>...</td>
</tr>
<tr>
<td></td>
<td>2 &quot;</td>
<td>...</td>
<td>1st strong, 30 tons 0·3 KCy; 3 p.m., 20 tons; 7 p.m., 6 tons.</td>
<td>...</td>
</tr>
<tr>
<td></td>
<td>9 &quot;</td>
<td>0·09</td>
<td>Standing under solution for 3 hours.</td>
<td>...</td>
</tr>
<tr>
<td>Thurs. 7th</td>
<td>12 p.m.</td>
<td>...</td>
<td>Started leaching until 12 noon.</td>
<td>...</td>
</tr>
<tr>
<td></td>
<td>1 a.m.</td>
<td>0·1</td>
<td>Changed to medium box, 3 a.m., 1·3; 6 a.m., 1·6; 9 a.m., 0·18.</td>
<td>...</td>
</tr>
<tr>
<td></td>
<td>12 noon</td>
<td>0·2</td>
<td>Changed to strong precipitating box.</td>
<td>...</td>
</tr>
<tr>
<td></td>
<td>12 &quot;</td>
<td>...</td>
<td>2' strong, 30 tons; 0·3 KCy. 3 p.m., 10 tons; 6 p.m., 15 tons; 8 p.m., 1 ton.</td>
<td>...</td>
</tr>
<tr>
<td></td>
<td>8 p.m.</td>
<td>0·28</td>
<td>Standing under solution for 3 hours. Started leaching until 8 a.m.</td>
<td>...</td>
</tr>
<tr>
<td>Frid. 8th</td>
<td>8 a.m.</td>
<td>0·26</td>
<td>Medium, 20 tons; 12 noon, 5 tons; 4 p.m., 8 tons; 8 p.m., 8 tons.</td>
<td>...</td>
</tr>
<tr>
<td></td>
<td>9 p.m.</td>
<td>0·2</td>
<td>Changed to medium; 12 p.m., 8 tons; 5 a.m., 10 tons; 12 noon, 7 tons.</td>
<td>...</td>
</tr>
<tr>
<td>Sat. 9th</td>
<td>3 &quot;</td>
<td>1·8</td>
<td>4 p.m., 10 tons; 8 p.m., 7 tons; 4 a.m., 12 tons; 8 a.m., 5 tons.</td>
<td>...</td>
</tr>
<tr>
<td>Sun. 10th</td>
<td>6 &quot;</td>
<td>1·4</td>
<td>Weak, 5 tons; 12 p.m., 5 tons; 6 a.m., 5 tons; 12 noon, 5 tons.</td>
<td>...</td>
</tr>
<tr>
<td>Mon. 11th</td>
<td>6 a.m.</td>
<td>0·1</td>
<td>Changed over to weak.</td>
<td>...</td>
</tr>
<tr>
<td></td>
<td>6 p.m.</td>
<td>0·9</td>
<td>5 tons; 12 p.m., 5 tons; 7 p.m., 5 tons; 12 p.m., 4 tons.</td>
<td>...</td>
</tr>
<tr>
<td>Tues. 12th</td>
<td>6 a.m.</td>
<td>...</td>
<td>Leaching dry.</td>
<td>...</td>
</tr>
<tr>
<td>Wed. 13th</td>
<td>6 &quot;</td>
<td>...</td>
<td>Discharging.</td>
<td>...</td>
</tr>
</tbody>
</table>

Amount of KCy used, 5·9 lbs. per ton.
Precipitation of Gold in the Extractor Boxes.—The precipitation of gold from the solution coming from the treatment vats is effected, he says, by the means of zinc shavings. A certain portion of the ore coming from the mine is of a rebellious nature, containing sufficient foreign metals to retard, if not prevent, the precipitation of gold by zinc. To counteract the effects of these foreign metals, it has been proved efficient in practice on these fields to set up a lead couple, and at these works this is effected by dipping the zinc shavings in a weak solution of acetate of lead, preparatory to being placed in the extractor boxes.

In all cases it is found that zinc containing from 1 to 2 per cent. of lead gives the most satisfactory precipitation. By the use of acetate of lead a perfect precipitation is brought about; only traces of gold remaining in the solution after passing through the extractor boxes.

The actual working costs amount to 1s. 11d. per ton, including general charges and maintenance.

It has been shown in America and Western Australia that with certain classes of ore and certain local conditions, it may be more advantageous to dry-crush than wet-crush preparatory to treatment. In South Africa wet-crushing is universal. Dry-crushing in that country has never been viewed with much favour, nevertheless the experiments made by Mr. Franklin White at the Luipaards Wei Estate mine are interesting and instructive.*

The tanks used were made of steel, 25 ft. in diameter and 8 ft. deep, with the usual filter bottoms and bottom discharge doors. Three tanks were placed close to the mill and mine in a row some distance away, and a little below. The ore was treated a few days in the upper tanks and then transferred to the lower, a double treatment being thus obtained. Mr. White does not consider double treatment to be of advantage in dealing with dry-crushed material, as the ore is properly mixed in the first instance, and there is abundance of air entangled in the dry sand. He is rather of the opinion that when the damp ore is transferred to the second row of tanks, there is a tendency to pack closer and to retard filtration.

Lime was added to the ore at the rock-breaker floor, in proportions varying from 1 lb. to 2½ lbs. per ton. By this means it was thoroughly mixed in the different machines, and any lumps were broken up.

The actual cyanide treatment differed a little from the ordinary procedure, and followed lines first suggested to the author by Mr. W. R. Feldtmann.

APPLICATION OF THE PROCESS IN DIFFERENT COUNTRIES. 165

As soon as the tank in the first row was filled, a strong solution, 0·25 per cent. KCy, followed by two others, the last being 0·15 per cent., was pumped on to it. The time given to this treatment would be sixty-six to seventy hours. The solutions were not allowed to stand, but were drained off when the tank was once filled. The object of this was to allow fresh air to obtain access to the mass of damp sand. Each solution would be about 27 tons to the tank of ore (165 to 170 tons).

The ore was then transferred to the second row of tanks, having lost about 67 per cent. of its original assay value in this short time. It is probable that the abundance of air entangled in the dry sand, as compared with what would be held in a tank of sand settled with water, materially assisted the solution of the gold. Also the finer grains of the free gold would be quickly taken up.

A solution of 20 to 25 tons, not exceeding 0·20 per cent. KCy, was then pumped on to the transferred sands and drained off, the mass being allowed to remain damp for about ninety-six hours, when weaker solutions (0·15 per cent. to 0·10 per cent.) were used in continuous washes, making up a total of 75 tons per tank (second treatment). A water wash of 20 or 30 tons completed this part of the process, which would last some 275 hours.

A careful series of moisture tests and measurements of solution sumps during the treatment of four tanks (680 tons) showed that the total loss of liquid in the treatment was 96 tons, or 24 tons per tank. The moisture in the discharged residues averaged 12·3 per cent., or, say, 20 tons per tank; the remaining 4 tons would be represented by evaporation from surfaces of sumps and tanks and by leakages. As the ore contained about 3 tons of water per tank, in the form of moisture when delivered from the mill storage bin, the actual consumption of fresh liquid equals 21 tons per tank, or $\frac{1}{4}$ ton (25 gallons) per ton of ore.

The solutions running from the first row of tanks carried from 13 dwt. to 32 dwt. per ton; those from the second row 3 dwt. to 4 dwt.; and the final wash 0·8 dwt. to 2$\frac{1}{2}$ dwt.

**Summary of Cyanide Costs.**

**TRIAL CRUSHING (COARSE).**

<table>
<thead>
<tr>
<th>Item</th>
<th>£</th>
<th>d</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tanks and extractor house work, per ton</td>
<td>2</td>
<td>3·30</td>
</tr>
<tr>
<td>Clean up and smelting, etc.,</td>
<td></td>
<td>5·15</td>
</tr>
<tr>
<td>Discharging residues,</td>
<td></td>
<td>8·36</td>
</tr>
<tr>
<td>Sundries,</td>
<td></td>
<td>1·05</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>3</td>
<td>5·86</td>
</tr>
</tbody>
</table>

Actual extraction 68·91 per cent.
NEW ZEALAND.*

The principal gold-bearing formation of the Hauraki Goldfields is of volcanic origin, consisting of a great accumulation of andesitic lavas, tuffs, breccias, and agglomerates of lower tertiary age. These rocks everywhere bear evidence of having been subjected to the prolonged leaching action of thermal waters, doubtless accompanied by steam and acid vapours. They are found in all stages of decomposition or alteration; and in many mines the hard blue andesite can be seen to pass by a series of almost insensible gradations into a soft, or fairly hard, greyish-yellow or blue altered rock, to which the distinctive name propylite has been applied.

It is in this altered andesite that the veins yielding payable ore occur. The veins vary from a few inches to 40 ft. in width, but in linear extension they can seldom be traced for any considerable distance. The ore values are also irregular, and in no case, as yet, is the same vein or lode worked with payable results in two adjoining mines.

Cyaniding Ores.—In the southern portion of the Hauraki peninsula, the pay-ores consist of whitish-grey chalcedonic or cryptocrystalline quartz, often possessing a wavy, banded structure of alternating layers of grey and blue flinty quartz. They are comparatively free from base sulphides.

The gold is about 645 fine, and usually associated with silver sub-sulphide (Ag₂S) in varying proportions. It is generally extremely finely divided, being seldom visible to the eye, and in the great bulk of the Waihi ore it is impossible to raise even a colour by panning.

Prior to the introduction of the cyanide process, these ores were treated by dry-crushing and hot pan-amalgamation with chemicals, by which a recovery of 65 per cent. was effected.

When cyanide treatment was adopted, dry-crushing was naturally continued at the different mills, the dry pulverized material being charged into shallow vats and treated directly by cyanide. From 65 per cent. by pan-amalgamation, the recovery rose at a bound to 85 per cent., and in some cases to 90 per cent., and the results were considered so satisfactory that no further improvement was considered possible.

In a few years, however, it became apparent that dry-crushing possessed many disadvantages compared with wet-crushing, the principal being the cost of drying the ore, the low duty of the

* Excerpt from paper by author read at California Meeting, September 1899, American Institute of Mining Engineers.
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stamps, and the large number of vats required for leaching. In 1897 mine owners began to turn their attention to wet-crushing, and one by one, since the beginning of 1898, the different mills have been adopting wet-crushing, until at the present time dry-crushing is the exception and not the rule, as it was two years ago.

Wet-crushing Practice.—A. For ores containing a large proportion of free, easily amalgamable gold, with a small proportion of fine or "float gold" and silver sulphide, the mill practice is:—

(a.) Crushing with water.
(b.) Plate amalgamation.
(c.) Spitzlute separation of sands and slimes.
(d.) Cyanide treatment of sands and slimes by ordinary percolation.

A typical example of an ore of this class is that of the Kauri Gold Estates at Opitonui, where a new 40-stamp mill has been started. The sands and heavy slimes are subjected to the "double" cyanide treatment, but it is doubtful if the additional saving will pay for the extra labour involved. So far no provision has been made for the treatment of the fine slimes. If they are worth it, they will probably be treated by agitation and decanting.

B. For a clean ore, almost identical with that described above, but containing a small proportion of free amalgamable gold, and a large proportion of fine cyanide gold, with little or no slimes, the method of treatment at the Crown Mines is:—

(a.) Crushing with cyanide solution.
(b.) Direct cyanide treatment of mixed sands and slimes by percolation.
(c.) Plate amalgamation of free gold.

With an ore so exceptionally free from slimes, it seems that the order of treatment could be reversed with advantage, both as regards stamp duty and efficiency of amalgamation on the plates.

The Crown Mines Company was the first to adopt wet-crushing for these gold and silver-bearing chalcedonic ores, which occurred in 1897, and much credit is due to Mr. F. R. W. Daw, the superintendent, for the successful inauguration of the method.

The ore is hard and splintery, clear and pure from all impurities, and, unlike most of the ores from the neighbouring mines, contains little or no silver except what is alloyed with the gold. It is crushed in the Company's 60-stamp mill with cyanide solution in the mortars, about 2½ tons of solution being used to 1 ton of ore. A 25-mesh screen is used, and the duty per stamp is about 2 tons per day. The slimes formed in crushing are said to amount to less than 5 per cent.
The monthly output is about 2750 tons. For the month of March 2916 tons were crushed, yielding bullion valued at £5797. The bullion was worth a fraction under £2 per ounce.

The cyanide plant consists of twenty-eight leaching vats, each 22½ ft. in diameter and 4 ft. deep. Much reticence is maintained as to the exact treatment, but the main features are understood to be as follows:—

The whole of the pulp from the 60-stamps is conducted by a launder to one vat, and allowed to discharge into the centre until the vat is about half full. The pulp is then diverted to another vat, which is allowed to fill in the same manner. The mixed sands and slimes in the first vat are allowed to settle for an hour or two, after which the fairly clean top solution is syphoned off into a collecting tank, whence it is pumped up to two elevated tanks, from which the solution for the stamps is supplied. The pulp is again diverted into the first vat until the charge is complete. After settlement, the top clear solution is again drawn off. In this way three or four vats may be in course of filling at the same time.

The settlement of the slimes is effected without the aid of lime, by allowing the solution to percolate from the bottom of the vat during the periods of filling and settlement. The downward tendency of the currents promoted by the draining from below is said to cause the settlement of the finest matter within a reasonable time. This is a point that should be noted by cyaniders troubled with slimy products.

The mixed sands and slimes are treated by percolation in the ordinary way. The depth of each charge is about 30 inches, and the weight 40 tons. The strong cyanide solution is allowed to percolate from thirty to forty hours, while the weak cyanide and water washes are drawn off by the aid of an air-pump.

The syphon used in the sand vats consists of a length of 2½ in. rubber hose, to one end of which is attached a short length of wooden batten to keep it on the surface of the solution. The other end is fixed, inside the vat, to a short iron pipe passing through the side about 18 inches above the filter-cloth.

In the extractor-room there are five precipitation boxes of the ordinary pattern, divided into compartments by baffle boards; and four zinc towers, consisting of wooden boxes about 6 ft. high and 30 in. square, set on end and connected in a series like charcoal towers. The solutions flow upwards through the zinc turnings, the overflow being conducted in a pipe to the bottom of the next tower, and so on to the last.

The sands are sluiced out of the vats over a wide expanse of
amalgamated copper-plates, which catch a certain proportion of the free gold.

The actual recovery from all sources is said to vary from 84 to 87 per cent., but the costs are not obtainable.

C. For ores containing some easily amalgamable gold, and fine gold associated with pyrites and silver sulphides, the treatment used is:—

(a.) Crushing with water.
(b.) Plate amalgamation.
(c.) Spitzlute separation of fine slimes, if necessary.
(d.) Vanner concentration of sulphurets.
(e.) Cyanide treatment of sands by percolation.
(f.) Cyanide treatment of slimes by agitation and decanting.
(g.) Cyanide treatment of concentrates by agitation.

The practice at the Woodstock mill is a typical example of this treatment.

The ore is a chalcedonic and finely crystalline quartz, containing a small proportion of clayey matter and a little pyrites. It is stained a greyish and blackish-brown colour through the presence of iron and manganese oxides.

At the Company's 40-stamp mill the monthly output is about 1100 tons, the stamp duty being slightly under 2 tons of 2240 lbs. per day. For the June month, 1901, 1000 tons of ore were crushed for a return of £1362, which is equal to a value of £1, 7s. 6½d. per ton. The value of the bullion varies from 8s. to 12s. per ounce, being principally composed of silver.

The ore is crushed with water and passed over amalgamated copper-plates, from the end of which the pulp is raised by a wheel elevator to a spitzlute. The slimes from the spitzlute are conducted to a slime tank, while the sands carrying some heavy slimes are passed over vanners, which collect about 1 per cent. of rich concentrates.

The vanner tailings, composed principally of sands and heavy slimes, are led to the leaching vats, which are provided with automatic distributors. The construction of the distributors is of the simplest character, being similar to those formerly used at Waihi. They consist of a central wooden box, pivoted on a wooden pillar fixed in the centre of the vat, and from which extend seven narrow wooden launders or arms of light make and different lengths, so as to effect an even distribution of the pulp. At the end of each arm there is fixed a piece of sheet zinc to divert the stream to one side.

The whole of the pulp from the vanner ends is collected in one stream and diverted into one vat at a time until the charge is
THE CYANIDE PROCESS.

filled. During the filling the overflow carries the lighter slimes into the slime vats.

The sands and heavy slimes are treated with cyanide by ordinary percolation.

The slimes from the spitzlute, and those from the sand vats, are agitated with cyanide in vats provided with slowly revolving arms. When the gold is dissolved lime is added, and the slimes are allowed to settle, after which the clear solution is decanted off. The slimes are washed by agitating with successive portions of water and decanting.

The concentrates have a value of £30 to £40 per ton, a large proportion of the value being in silver sulphide. They are treated by agitation with a 4 per cent. solution of cyanide for thirty-six hours. Two pounds of lime are added to every ton of concentrates. The charge weighs about 1½ tons. Mr. F. Rich, the superintendent, who adopted the present treatment, informed the author that the recovery varied from 90 to 94 per cent., at a cost of 18s. per ton for labour and material.

The recovery by cyanide from all sources per ton of ore milled is said to vary from 82 to 86 per cent., at a cost of 4s. 9d.

D. For very slimy ores containing very little easily amalgamable gold, and a large proportion of extremely fine gold besides the usual silver sulphides, the treatment is as follows:—

(a.) Crushing with cyanide solution.
(b.) Spitzlute separation of sands and slimes.
(c.) Treatment of sands by percolation.
(d.) Treatment of slimes by agitation and decanting.

This method of treatment is subject to various modifications as regards mechanical appliances and methods of application, but the general principles are everywhere the same.

The procedure at the Waitekauri 40-stamp mill is as follows:—

The ore, which contains a good deal of oxidized products, is crushed with cyanide solution in the mortars. From the screens the pulp is conducted direct to the sand vats, into which it is distributed by means of revolving wooden box launders actuated from a secondary shaft. The slimes, of which there are about 33 per cent., are allowed to drain into a collecting vat, flowing through a pipe fixed in the inside of the vat. This pipe has a movable joint, and is raised by a screw as the pulp accumulates in the vat.

The collecting vat is provided with revolving arms which keep the fine slimy pulp from settling. From this vat the slimes are pumped into the slime leaching vats, which are provided with a double set of slowly revolving arms, the lower ones having rakes
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on them and the upper ones loose pieces of sacking which drag through the pulp. In these vats the slimes are treated by agitation and decanting, lime being added with each wash to facilitate settlement.

The sands are treated by ordinary percolation with first a 0.5 per cent. solution of cyanide, and then the weak and water washes.

There are 12 sand vats, 14 slime vats, and 2 slime collecting vats each 22.5 ft. in diameter and 4 ft. deep. The monthly output of the 40-stamps is about 2200 tons of 2240 lbs. The June monthly output was larger than usual, being 2543 tons, yielding 7220 ozs. of bullion valued at £6773 0s. 6d., equal to a value of 18s. 9d. per ounce. The actual recovery is said to be 90 per cent., at a cost of 5s. 6d. per ton.

The exceptionally large proportion of slimes in this ore rendered the adoption of wet-crushing a knotty and difficult problem. The increased output, higher extraction, and lower costs are proofs enough of the success of the change from dry-crushing which was effected under the supervision of Mr. G. Davey, the superintendant, without hitch, or decrease in the monthly output, a matter of no little moment in these days of heavily capitalized public companies.

Concluding Remarks.—Among the points most likely to attract the notice of cyaniders are the low stamp duty and the heavy consumption of cyanide.

The low stamp duty of the New Zealand mills has often been a subject of discussion, but no satisfactory explanation has yet been advanced. In the opinion of the author, it is due to the circumstance that the mills having been designed and erected in the first place for dry-crushing, the mortars are too narrow and restricted to give good results by wet-crushing, and until these are replaced by new mortars, specially designed for wet-crushing, it seems hopeless to look for better results. The advantages of a stamp duty of 4 or 5 tons per day instead of 2 tons are too obvious to require enumeration.

With reference to the large consumption of cyanide, it is well known that silver in all its forms requires a stronger solution to effect its dissolution than gold; and in the Hauraki Goldfields the large consumption of cyanide is due to the presence of silver sulphide (principally Ag₂S), which is generally worth saving, and to the circumstance that the free gold is alloyed with about one-third its weight of silver.

According to Elsner’s equation for the dissolution of gold by potassium cyanide, 4 lbs. of cyanide should dissolve 100 ozs. of gold, but in practice it is found that it takes nearly forty times
that quantity. To dissolve 100 ozs. of silver would require 7.5 lbs. of cyanide, according to the equation:

$$4\text{Ag} + 8\text{KCy} + O_2 + 2\text{H}_2\text{O} = 4(\text{AgKCy}_2) + 4\text{KHO}.$$ 

For the dissolution of 100 ozs. of silver existing as the sub-
sulphide \((\text{Ag}_2\text{S})\), 7.01 lbs. of cyanide would be required by the following equation:

$$\text{Ag}_2\text{S} + 4\text{KCy} = 2(\text{AgKCy}_2) + \text{K}_2\text{S}.$$ 

The potassium sulphide resulting from the dissolution of silver sulphide also tends to cause a loss of cyanide. It has been shown by Crosse and others that a trace of alkaline sulphide in cyanide solutions does not act injuriously, but the large quantity of \(\text{K}_2\text{S}\) liberated in the treatment of the silver-bearing ores of the Hauraki Goldfields must cause the precipitation of a portion of the dissolved gold in the vats. Much of this precipitated gold is doubtless redissolved by the excess of free cyanide present in the solutions, but it always requires this excess to obtain adequate extractions, thus necessitating the use of comparatively strong solutions.

One of the most perplexing features connected with the treatment of these ores is the constantly varying proportion of silver, which necessitates the use of solutions of varying strength to obtain adequate extractions, thus adding another source of anxiety to the many worries which the use of cyanide entails on even the successful cyanider.

At many of the Hauraki mines, cyanide treatment was adopted by the owners on the author’s recommendation, but only after he had made a careful investigation of the constituents of the ore, and repeated trials on a working scale, at the Government Metallurgical Works at the Thames. In other cases, the necessary experimental trials were made by the author’s assistants at the mine. In no case was the process adopted until success had been assured, a precaution which doubtless has been a potent factor in promoting the popularity of cyanide treatment in this country.

The ores of Te Aroha and Monowai are generally very refrac-
tory, containing free milling gold, mostly very fine, associated
with sulphides of silver, iron, copper, lead, zinc, and often mer-
cury. Many attempts have been made to treat them by cyanide,
but without success, and, so far as our present knowledge goes, it is doubtful if they can ever be treated successfully in the raw state by that process.

For the treatment of cupriferous ores and concentrates from
the Jubilee, Sylvia, and Monowai mines, which could not be
treated successfully by ordinary cyaniding, the author obtained
good results by first subjecting the ore to a chloridizing roast, and
then leaching out the copper chlorides with water. After an alkaline and water wash, the gold and silver contents were extracted by cyanide by ordinary percolation. During the roasting the silver sulphides present were chloridized, the chloride being easily dissolved by cyanide.

From a large parcel of Monowai ore, 92 per cent. of the gold and 85 per cent. of the silver were extracted, the composition of the ore being (F. B. Allen, M.A., B.Sc.):

<table>
<thead>
<tr>
<th>Component</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Insoluble gangue</td>
<td>90.15</td>
</tr>
<tr>
<td>Copper pyrites</td>
<td>3.78</td>
</tr>
<tr>
<td>Iron pyrites</td>
<td>4.40</td>
</tr>
<tr>
<td>Galena</td>
<td>0.25</td>
</tr>
<tr>
<td>Zinc-blende</td>
<td>0.26</td>
</tr>
<tr>
<td>Alumina</td>
<td>0.13</td>
</tr>
<tr>
<td>Water and loss</td>
<td>1.03</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>100.00</strong></td>
</tr>
</tbody>
</table>

The bullion contents of this ore were:—Gold, 1 oz. 5 dwt.; silver, 14 oz., per ton.

The Moanataiari Co.'s Works, situated at Thames, are of recent construction, and in point of completeness and modernness of equipment, among the finest in New Zealand. The plant consists of a 60-head battery, two rock-breakers, grizzlies, automatic feeders, 24 vanners, 21 Berdan pans, a complete cyanide plant for treating vanner concentrates, 9 Cornish buddles for concentrating the tailings from the vanners, and all up-to-date appliances for assaying and retorting. The cyanide works consist of three steel vats, 20 ft. in diameter and 7 ft. deep, each provided with two bottom-discharge doors, with a capacity each of 200 tons of concentrates; two zinc extractors, each 15 ft. 6 in. in length; and three concrete sumps, 50 x 11 x 6 ft. over all. Wet-crushing, concentration, and the cyanidation of the concentrates are the interesting features of the practice at these works, which were designed by the author and erected under his supervision. The cyaniding of concentrates is comparatively new in New Zealand.

Talisman Mine.—In the Ohinemuri Goldfields, the practice of dry-crushing and direct cyanide treatment has been superseded by wet-crushing, concentration, and cyanide treatment of sands, slimes, and in some cases the concentrates. At the Talisman mill, the ore was conveyed from the mine by an aerial tramway being dumped on to a grizzly, which passed the coarse stuff to a Blake Marsden crusher, whence the ore passed to a revolving
drying furnace. The stamps weighed 1000 lbs., and the mortars were provided with back and front discharge.

The pulverized ore was elevated to an ore-bin, from which it was carried to the cyanide vats by a long line of revolving screw-conveyors. The mechanical drier was a very efficient machine, drying about 12 tons of ore for every ton of firewood consumed.

This company's dry-crushing mill has been dismantled, to give place to a new 50-stamp wet-crushing plant. Wire plate-amalgamation, vanner concentration, separation of slimes, cyaniding of sands and slimes, the latter by agitation and decantation, has been introduced.

Waihi Mine.—The ore in the upper levels of the celebrated Martha lode at the Waihi mine is typical of most of the ores in this district. It consists principally of hard, splinterly, whitish-grey chalcedonic and crypto-crystalline quartz, often possessing a banded and wavy structure. It is perfectly free from all base metallic sulphides, and the amount of iron oxides present is so small that when roasted and pulverized the colour of the dust is pinkish-grey.

The value varies from £3 to £4 per ton, the precious metals existing in the proportion of about 8 or 10 oz. silver to 1 oz. gold. The free gold is alloyed with about 35 per cent. of silver, being valued at about 53s. per oz. The greater proportion of the silver exists in the form of the bluish-grey sub-sulphide known as argentite. In the surface levels, thin leafy plates of gold were not infrequently seen adhering to the surface of large cuboidal masses of quartz, but in the lower levels a colour is rarely seen, the gold existing in an extremely fine state of subdivision. Such an ore is theoretically perfect for cyanide treatment, and actual experience has proved it to be so. By the old stamp battery, and copper-plate amalgamation, the recovery amounted to only some 4 dwt.s. per ton, equal to about 15 per cent. of the value. By dry-crushing and pan-amalgamation the extraction was raised to 60 per cent., but at present the actual extraction by the cyanide process amounts to over 90 per cent. of the assay value.

The following particulars of the dry method of treatment formerly in use at the Waihi mills were supplied by Mr. H. P. Barry, the general manager, for the annual report of the New Zealand Mines Department for 1894:—

Drying the Ore.—The ore is trucked to the drying kilns which consist of open circular holes excavated in the solid rock, their dimensions being 37 ft. in depth and 20 ft. in diameter at the top, and tapering somewhat at the bottom. Each kiln is capable of holding 100 tons of ore at a charge. The lower part is lined with bricks, and finished off with a brick arch, having a door
and an iron chute for discharging the dried ore into trucks, which have access to the kiln by means of a tunnel cut in the rock.

**Charging the Kilns.**—The kilns are charged with alternate layers of wood and ore, the layers of wood being about 5 ft. apart. When the kiln is fully charged, the wood is lighted, and, after it is all burned up, about half the charge, that is about 50 tons, is withdrawn and another 50 tons of raw ore, together with the necessary wood, are placed on the top. After this about 50 tons of ore are withdrawn every third day, while a similar quantity of raw ore and wood is added.

The method of drying the ore is very expensive, as one ton of wood will only dry about three tons of ore. The cost of firewood at the Waihi big mill is 2s. per ton of ore dried, and the total cost of drying, including labour, is 2s. 6d. per ton.

**Crushing and Pulverizing.**—From the kilns, the dried ore is trucked to the rock-breakers, whence it passes by gravitation to the self ore-feeders. The pulverizing machinery consists of a 90-stamp battery and an Otis ball-mill, having a capacity of about 10 stamps. The ore is passed through a 40-mesh screen.

**Filling the Cyanide Leaching Vats.**—From the screens, the dry dust falls into a long, narrow trough running parallel with the stamp-motors, along which it is conveyed to the dust-bin at one end of the mill by means of an Archimedean screw. From the dust-bin the pulverized ore is lifted by a bucket-belt elevator and discharged on to an 8 in. rubber-belt provided with rope edges, and by this conveyed to, and across, the dust-hopper, which is 110 ft. long, running the entire length of the cyanide-plant house.

The dust-hopper has twenty doors for discharging the dust into the trucks, which are run straight out over the leaching vats on to travellers running on rails. The travellers are provided with hand traversing gearing, thus enabling a truck to be tipped at any part of the vat. This is an important feature, as the finely pulverized material has a tendency to pack if moved about or touched in any way after being tipped into the vat.

As a further preventative against packing, there is a small traveller fixed below the main traveller, provided with a platform at the height the ore has to be filled up to. All the trucks are tipped over this platform, which breaks the fall of the dust, and throws it in a light shower all around.

**The Cyanide Treatment.**—The following particulars of the cyanide treatment were kindly supplied to me by Mr. E. G. Banks, the chemist of the cyanide operations.

The plant consisted of thirty-eight circular leaching vats, each 22½ ft. in diameter and 4 ft. deep, together with the necessary
dissolving and solution vats, sumps, extractors, vacuum cylinders, solution and air-pumps, etc.:

<table>
<thead>
<tr>
<th>Activity</th>
<th>Hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filling vat, 30 tons, two men,</td>
<td>2½</td>
</tr>
<tr>
<td>Strong solution, 10 tons, 0·35% KCy leaching,</td>
<td>30</td>
</tr>
<tr>
<td>Weak solution, 7 tons, 0·1% KCy, with vacuum,</td>
<td>15</td>
</tr>
<tr>
<td>First water wash, 6 tons, with vacuum,</td>
<td>24</td>
</tr>
<tr>
<td>Second</td>
<td>36</td>
</tr>
<tr>
<td>Discharging vat, one man sluicing,</td>
<td>2</td>
</tr>
<tr>
<td>Taking up and cleaning filter-bottom,</td>
<td>4½</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>114</td>
</tr>
</tbody>
</table>

A vacuum of 20 in. to 23 in. is maintained to obtain the above results. The average value of the ore was about £4 per ton, and the actual extraction from 90 per cent. to 92 per cent. of the original value, at a cost of 7s. 6d. per ton, not including royalty.

Dry-crushing and direct cyaniding of the pulverized ore has been abandoned at all the Waihi Company's mills in favour of wet-crushing, concentration, and subsequent treatment of sands and slimes. The treatment of the slimes is effected by agitation and filter-pressing, combining some of the essential features of both South African and Western Australian practice.

I am indebted to Mr. G. Banks, the company’s metallurgist, for the following clear and succinct notes on Waihi cyanide practice at the different mills. Victoria mill.—The ore is pulverized by 200 stamps to pass through 40-mesh wire wove steel screens, and is then elevated by means of plunger sand pumps to launders which convey the pulp to nests of spitzluten where the separation of the sands and slimes takes place.

*Treatment of Sands.*—The sands flow to the intermediate sand collecting vats, of which there are five built of steel, 38 ft. in diameter by 8 ft. deep, fitted with an annular lauder on the outside top edge to convey the overflow water, at times containing a little slime, away to the slime thickening boxes. The sands are charged into the percolating vats by means of an automatic revolving distributor, which is moved on an overhead traveller from vat to vat as required.

The vats are fitted with Roche’s bottom-discharge doors (Pl. XII.). The filter-bed is arranged by a wooden grating covered with woolpack. Each vat holds from 250 to 300 tons of sands. After draining to get rid of surplus water, a preliminary treatment is given with weak cyanide solution, followed by a strong (0·35 per cent. to 0·45 per cent.) solution and usual washes.
The old vat-house contains ten rectangular concrete vats, five on each side. Each vat is 50 ft. by 40 ft. and 4 ft. deep.

**Slime Treatment.** — The slimes from the spitzluten and the overflow from the sand vats are mixed with lime-water (about 2 to 4 lbs. of lime per ton of slime) and conveyed to a nest of 36 V-shaped thickening boxes, in which the slimes quickly settle, and are then drawn off in a fairly thick state from the bottom of the boxes. The overflowing clear water is returned to the stamps.

The thickened slimes are collected in six steel vats 32 ft. in diameter by 14 ft. deep. In these the slimes rapidly settle, and as the clear water overflows, it is stored for re-use in the mill and spitzluten.

When the vat is filled with slime-pulp to within 2 or 3 ft. from the top, the inflowing slimes are cut off and the charge allowed to settle for about 24 hours. Whatever clear water may be on top is syphoned off, and the thickened pulp (now about 1 of slime to 1 of water) is ready for treatment in the agitators.

The agitators are built of steel, 20 ft. in diameter by 6 ft. deep, and hold about 25 tons of slime (dry weight), together with about 40 tons of cyanide solution. The strength of the solution is about 0·1 per cent. The pulp is agitated by paddles, secured to a vertical shaft, actuated by overhead worm-gearing. The speed is 8 revolutions per minute.

After 24 hours' agitation the pulp is run to a pressure tank, and from thence is forced by compressed air into Johnson 6-ton filter-presses, where the gold-bearing cyanide solution is extracted.

The zinc method of precipitation is used and does very good work, even on very dilute solutions.

Vanners are being erected to concentrate the more heavily mineralized ore. The tailings from the vanners will be mixed with the lightly mineralized ore-pulp, and pass through the course of treatment just described.

An extraction of 85 per cent. to 90 per cent. of the ore value can be recovered by this process, but no particulars of the actual recovery or costs are yet available.

**Waithi Union Mill and Cyanide Plant.** — This mill consists of 40-stamps for wet-crushing and plate-amalgamation; sixteen circular steel vats for sand treatment (each vat having a capacity of 22 tons of sand); three slime-collecting vats, 32 ft. in diameter by 14 ft. deep; two agitators, 20 ft. in diameter by 6 ft. deep; and one Johnson filter-press, with a capacity of 6 tons of slime-dry weight, per charge. In addition there are the usual ore-
breakers, spitzluten, pumps, air compressors, etc. The screens used are 40-mesh wire-woven steel, and the pulp after passing the amalgamated tables is elevated by a wheel-elevator to a height of 25 ft. and is here classified into sands and slimes by four spitzluten.

Sand Treatment.—The sands flow to the vats and are distributed by means of an automatic revolving distributor. A weak sump-wash, containing about 0·07 per cent. to 0·1 per cent. of cyanide, is run on, and this is followed by a solution 0·4 to 0·6 strong. The total time of treatment is from five to six days. The tailings are sluiced away through a bottom central discharge-door Pl. XII.

Slime Treatment.—The slimes, after the addition of lime-water (about 3 lbs. of lime per ton of ore), flow to the large collecting vats and are there thickened by natural settlement to about one part of slime to one of water. In these vats, a syphon provided with a ball-and-socket joint drains off the top clear water automatically. The clear water overflowing from the settling vats is sent to a reservoir to be pumped up for re-use in the mortar-boxes. This course is found to be necessary owing to the fact that even when the water appears to be perfectly clear it contains from $\frac{1}{2}$ grain to $2\frac{1}{2}$ grains of gold per ton, and as several tons of water are used per ton of ore, it is evident the loss would be from 6d. or 8d. up to two or three shillings per ton of ore, if the water were allowed to run to waste.

From the collecting or settling vats, the thickened slimes are drawn off to the agitators, situated at a lower level, in charges equal to 20 to 25 tons of dry slime, and are agitated for 24 hours with $1\frac{1}{2}$ tons of 0·1 per cent. cyanide solution to 1 ton of dry slime.

Waihi Mill.—This is the oldest of the Waihi Company’s mills. Here the ore for some years was dry-crushed and pan-amalgamated by the Washoe process; and afterwards dry-crushed, and the dry pulp cyanided by the direct process in the manner described by Mr. Barry in the preceding pages. Of the present treatment Mr. Banks writes in April of 1903 as follows:—

"At the old 90-stamp mill dry-crushing was stopped at the end of 1902 and alterations necessary for wet-crushing rapidly completed. By the middle of January 1903, the mill was restarted. The ore (about 220 tons per day) is stamped through 40-mesh wire-woven steel screens; passed over amalgamated tables, of which there are fifteen, each 12 ft. by 6 ft.; and then over thirty union vanners which extract about $2\frac{1}{2}$ to 4 per cent. of concentrates. After passing the vanners, the pulp is elevated by wheel elevators to a series of sand separating boxes. Here the sands and slimes are separated, care being taken to keep the sand as free from slime as possible.

"Treatment of the Sand.—This is effected in circular vats $4\frac{1}{2}$ ft.
deep and capable of dealing with 40 tons of sand per charge. The vat is first filled with water and the sand is then run in through an automatic revolving distributor. The overflow water (carrying a little fine sand and slime) runs into an annular launder and is conveyed back to the elevating wheels and joins the pulp flowing to the separating boxes. When sufficient sand has been run into the vat a 2 in. draw-off pipe, passing through the side of the vat, is lowered and the water drawn off. This water also contains a little slime and flows to the elevating wheels.

"After the charge has drained for several hours, a weak sump wash (testing about 0·08 per cent. KCy) is run on, and this is followed by a strong solution testing 0·5 per cent. KCy; when this solution has reached the bottom of the charge, percolation is stopped and the solution circulated by means of an air-lift inside the vat. After five days the strong solution is drawn off and the charge washed with weak solution and water. The tailings are then sluiced away.

"Treatment of the Slime.—After separation from the sand, the slime pulp (about 20 of water to 1 of slime) is mixed with lime-water and conducted over a nest of spitzkasten, thickening the slime-pulp to about 10 per cent. of slime, the balance of the water passing over in a clear state to be re-used in the mortars.

"The slime-pulp is now collected in one of two collecting vats, which are 14 ft. deep by 32 ft. in diameter, and fitted with an overflow launder. The pulp is run to the centre of the vat and discharged through a vertical box chute extending several feet below the top of the vat, so as to cause as little disturbance as possible. The pulp runs in until the slimes show at the top, the clear water overflowing to a reservoir for re-use. The collecting vats thicken the pulp to about 2½ of water to 1 of slime.

"The thickened slime pulp is now still further separated from the contained water by means of filter-presses, of which two are required for ‘drying’ the slimes, each dealing with about 40 tons (dry weight) of slime per day.

"The slime-cakes, now containing only 25 per cent. to 35 per cent. of moisture, drop from the presses on to a screen-conveyor which discharges into a disintegrator. This disintegrator is 7 ft. in diameter by 14 ft. deep and fitted with three sets of revolving arms bolted to a centre shaft driven by overhead gearing at the rate of 20 revolutions per minute. Cyanide solution (0·12 per cent.) flows in at the bottom of the disintegrator in such proportion that the overflowing pulp contains 1 of slime to 1½ of solution. This pulp flows through a series of four agitators each 14 ft. deep by 20 ft. in diameter, fitted with stirring gear making three revolutions per minute."
"The stirring gear consists of two arms secured one foot from the bottom of the vat to a central shaft driven by overhead worm-gearing. Effective agitation is produced by blowing compressed air through a number of pipes dipping down close to the stirring arms. The revolving arms serve to bring the pulp in contact with the escaping air, at which point fairly violent agitation is produced.

"The pulp enters No. 1 agitator at the top, flows from the bottom of No. 1 through a connecting pipe terminating half way up the side of No. 2. Similarly it passes from No. 2 to No. 3, and from No. 3 to No. 4. From No. 4, the pulp flows to a larger agitator, No. 5, which is 32 ft. in diameter by 14 ft. deep, and also fitted with stirring gear and air-agitation. The compressed air in the pressure tank after filling a filter-press is utilized by being blown through the agitators.

"The pulp is drawn off from No. 5 agitator, from time to time as required, to fill one of the four filter-presses used to complete the treatment. A press charge contains about 5½ tons (dry weight) of slime. The cakes are 3 in. thick.

| Time required to fill press at 60 lbs. pressure | Hours |
| " to wash at 75 lbs. pressure | 3 |
| " to open, discharge, and close press | 1½ |

**Total** | 5½ |

"From five to six hours can be taken as the time required to treat a press charge of Waihi slime. Of course, the more fine sand contained in the slime, the quicker a press will fill and wash. About 25 dry tons of Waihi slime can be taken as the average amount treated per press per day. The labour required is one man to each press each shift of eight hours. The cost for cloths, rubber-rings, etc., is from 2d. to 4d. per ton of slime treated.

"Extraction.—By amalgamation and concentration from 40 per cent. to 50 per cent. of the value is recovered, each process extracting about half that percentage. The sand treatment extracts about 80 per cent. of the gold and 60 to 70 per cent. of the silver contained in the sands. There is obtained by the slime treatment a recovery of 95 to 96 per cent. of the gold, and about 70 per cent. of the silver. The total extraction is about 89 per cent. to 90 per cent. of the gold and 75 per cent. of the silver.

"The consumption of cyanide per ton of ore varies from 2½ lbs. on lightly mineralized ore, to 4 lbs. on heavy sulphide ore."
WAIKINO
WAHI CO'S WET CRUSHING PLANT
as used at the
INTERMEDIATE SAND COLLECTING VATS
Bottom discharge door
APPLICATION OF THE PROCESS IN DIFFERENT COUNTRIES.

"The cost of the slime treatment is not given, but it will probably work out to about 6s. 6d. per ton of slimes treated.

"The output of the Waihi Company is about 18,000 long tons every four weeks of twenty-four working days, for a return varying from £50,000 to £52,000. The ore is pulverized in the company's mills, comprising 320 stamps, and then subjected to cyanide treatment by the various processes described above. The cyanide plants have a capacity of about 1000 tons per day."

From the foregoing description it will be seen that the slime process in use at the Waihi and Waihi Union mills is practically the same as that in use at Kalgoorlie in Western Australia. On the other hand, the process at the Waihi Company's Waikino mills is almost identical with that in use at Glencairn Main Reef, Johannesburg, differing only in the final stages in the adoption of filter-presses to separate solutions from the slimes instead of decantation which is almost universal in South Africa.

**Waihi Tailings.**—The treatment of the Waihi Company's tailings is very instructive. The tailings are the residues resulting from the pan-amalgamation of the dry-crushed ore before the introduction of the cyanide process. The ore was crushed through a 60-mesh screen, and pan-amalgamated in charges. The residues were discharged from the settlers into large dams, where they were allowed to settle. They mostly consisted of fine sands and a good deal of slimes. They contained no base metallic impurities, and the gold existed principally in the form of amalgam.

Some 25,000 tons of these tailings were successfully treated by the Cassel Gold Extracting Company, whose works have recently been acquired by the Waihi Gold and Silver Mining Company, who treated the remainder of the tailings on their own account. The plant consists of eight leaching vats, each 22½ ft. in diameter and 4 ft. deep, together with all the necessary appliances.

The details of the cyanide treatment adopted for the treatment of these tailings are given below in tabulated form:—

**Cyanide Treatment of Waihi Tailings.**

<table>
<thead>
<tr>
<th>Activity</th>
<th>Hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filling leaching vat, 30 tons, three men</td>
<td>8</td>
</tr>
<tr>
<td>Preliminary lime or water-wash, 6 tons, with vacuum</td>
<td>6</td>
</tr>
</tbody>
</table>

**Leaching—**

<table>
<thead>
<tr>
<th>Solution Type</th>
<th>Quantity</th>
<th>% KCy</th>
<th>Hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>Strong solution</td>
<td>8 tons</td>
<td>0.6%</td>
<td>30</td>
</tr>
<tr>
<td>Weak solution (from strong sump)</td>
<td>4 tons</td>
<td>0.2%</td>
<td>12</td>
</tr>
</tbody>
</table>
Washing, using Vacuum—

<table>
<thead>
<tr>
<th>Step</th>
<th>Quantity</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>First weak cyanide wash (from weak sump)</td>
<td>4 tons</td>
<td>12</td>
</tr>
<tr>
<td>Second</td>
<td></td>
<td>12</td>
</tr>
<tr>
<td>Third</td>
<td></td>
<td>12</td>
</tr>
<tr>
<td>Fourth, water wash</td>
<td></td>
<td>12</td>
</tr>
<tr>
<td>Discharging vat, one man sluicing</td>
<td></td>
<td>4</td>
</tr>
</tbody>
</table>

Total: 108

Remarks.—The tailings were generally very clean, and the preliminary lime or water wash was not always applied. The strong solution was allowed to stand in contact with the tailings about four hours before the percolation was commenced. The average value of the tailings was about 24s. per ton, and the actual extraction about 75 per cent., at the cost of 8s. per ton.

Try Fluke Mine.—At the works of this Company, at Kuaotunu, the ore was wet-crushed through a 40-mesh screen, and passed over amalgamated copper-plates. The tailings were run directly from the plates into settling-pits. When the pits were full, the slimes were removed from the lower end, and spread out to dry in the sun. When dry, they were broken up, and then were filled into the leaching vats together with sand, in the proportion of one truck of slimes to two trucks of sand.

The sand and slimes were thoroughly mixed in the vat before the solution was put on. The Company’s metallurgist informs the author that the average time of treatment was as follows:

<table>
<thead>
<tr>
<th>Operation</th>
<th>Hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filling vats, 20 tons</td>
<td>6 to 8</td>
</tr>
<tr>
<td>Strong solution, 5 tons, 0·6% KCy, standing in contact with tailings</td>
<td>8 to 12</td>
</tr>
<tr>
<td>Percolating</td>
<td>24 to 30</td>
</tr>
<tr>
<td>Weak solution, 5 tons, 0·2% KCy</td>
<td>5 to 6</td>
</tr>
<tr>
<td>Weak cyanide washes, five of 5 tons each, 0·1% of KCy</td>
<td>25 to 30</td>
</tr>
</tbody>
</table>

Totals: 68 to 86

The ore consisted of grey and yellowish-brown quartz, sometimes containing a considerable proportion of iron and manganese oxides, the latter generally predominating. The greater part of the gold was excessively fine, being locally known as “float gold.”

The average value of the tailings was about 20s. per ton; and the actual extraction by cyanide about 75 per cent., at a cost of 7s. 6d. per ton. When the tailings were of higher value than usual, they were turned over in the vat after the last washing and
washed again. In this case it was found that the extra extraction more than paid for the extra labour.

At the Kapai-Vermont mine, which adjoins the Try Fluke, the same ore was dry-crushed in a ball-mill, and then subjected to direct cyanide treatment with the most satisfactory results, the actual extraction generally exceeding 85 per cent. of the assay value. In this mine, shoots of very rich ore have been frequently met with, containing a considerable proportion of comparatively coarse gold. With such ores the strong cyanide solution was circulated through the leaching vats until an adequate extraction was obtained.

**Waitekauri Mine.**—At the Golden Cross section of the Waitekauri Gold Mining Company's Special Claim, near Waihi, the ore, before the introduction of wet-crushing at the new mill, was dry-crushed with stamps to pass through a 40-mesh screen, and treated directly with cyanide. As a small proportion of the gold was coarse, the tailings were passed over amalgamated copper-plates, 30 ft. long and 3 ft. wide, set with a fall of 1 in 12.

The details of the cyanide treatment of this ore may prove useful, and are given in the following tabulated statement:

<table>
<thead>
<tr>
<th>Operation</th>
<th>Hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filling vats, three men, 22 tons</td>
<td>3 1/2</td>
</tr>
<tr>
<td>Strong solution, 9 tons, 0.45% KCy</td>
<td>48</td>
</tr>
<tr>
<td>Weak solution, 9 tons, 0.2%</td>
<td>18</td>
</tr>
<tr>
<td>First weak cyanide wash, 5.5 tons, 0.05 to 0.15%</td>
<td>18</td>
</tr>
<tr>
<td>Second &quot; 5.5 &quot; &quot;</td>
<td>18</td>
</tr>
<tr>
<td>Third &quot; 5.5 &quot; &quot;</td>
<td>18</td>
</tr>
<tr>
<td>Fourth, water wash, 3.0 &quot; &quot;</td>
<td>18</td>
</tr>
<tr>
<td>Discharging vat, one man sluicing</td>
<td>14 1/2</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>156</strong></td>
</tr>
</tbody>
</table>

The average value of the ore treated during 1894 was £4, 15s. per ton; and the actual extraction varied from 91 per cent. to 93 per cent., at a cost of 8s. 6d. per ton.

**WESTERN AUSTRALIA.**

**TREATMENT OF SULPHO-TELLURIDE ORES.**

At Kalgoorlie, the chief mining centre in the State, the gold occurs in the oxidized surface ores in a free state, and in the unoxidized ores in combination with tellurides and sulphides. In all cases it exists in an extremely fine condition in talcose
calcareous ores that possess a great propensity to form slimes. The successful treatment of these ores at first presented a difficult problem to metallurgists, and after repeated failures along old lines, a process was developed that in some respects possesses some peculiar features. And the early difficulties were not confined to the mere treatment of the rebellious ore. The climate was tropical and dry, water scarce and often brackish, and the distance from the seaboard great, and over arid plains. But these difficulties have now been overcome. A State railway connects the distant mining centres with the capital, and a State water supply, the greatest undertaking of the kind in the world, provides an abundance of pure water, carried in a pipe-line for a distance exceeding 300 miles.

The telluride ores are very brittle, and consequently a larger proportion of the gold goes into the slimes than into the sands. This circumstance and the slimy character of the ore led to the adoption of the cyanide process and filter-press treatment of the slimes. The object at many of the mills is to slime as high a proportion of the ore as possible.

The Diehl and Riecken processes have been installed at several mines with, it is said, satisfactory results. A detailed description of them will be found in Chapter XIII.

In some of the mills the slimes are drained or partially dried in presses, disintegrated and agitated with cyanide solution, and afterwards pressed and washed in presses. In other mills the dissolution of the gold is effected directly in the presses.

I am indebted to Mr. F. B. Allen, M.A., B.Sc., Director of the Kalgoorlie School of Mines, for the following interesting details of the general methods of treatment adopted at that mining centre:—

**General.**—The essential features of the treatment are dry-crushing, or crushing with cyanide solution, concentration, roasting or not according to circumstances, fine sliming, leaching of slimes with cyanide, and filter-pressing. The concentrates are roasted, ground in pans, and partly treated by amalgamation and partly by cyanide.

The roasting is generally effected in Edwards or Brown straight-line furnaces. The former roast from 14 to 16 tons per day, at a cost of 4s. to 4s. 6d. per ton; and Brown, 30 tons per day to 0·1 per cent. sulphur as sulphide, for 7s. to 9s. per ton.

Ore containing over 3 per cent. of moisture is dried before dry-crushing in ball-mills. No. 5 Krupp, running at 25 revolutions per minute with 15 H.P., crushes 25 tons through a 40-mesh screen for 1s. 2d. per ton, and of this from 65 to 75 per cent. will pass a 100-mesh sieve. The Griffin mill, when crushing to a 15-mesh, will form a product of which 75 per cent. will pass through a 100-mesh sieve, at an approximate cost of 2s. per ton.

**Great Boulder Proprietary.**—The sulphide ore is partly
crushed wet with stamps and amalgamated in Wheeler pans, and partly dry-crushed in Griffin mills, roasted, fed into mixers which supply Wheeler pans with the pulp, which is further ground and amalgamated with mercury without the use of copper-plates.

Cyanide Process.—The pulp from the continuous overflow of the pans is led into settlers, first passing, in the case of sulphide ore, over canvas tables to eliminate concentrates. About one-third of the gold contents are recovered by amalgamation. The furnaces used are Edwards (16 tons per day), fed with producer gas, and they roast the ore down to 0.11 per cent. sulphur as sulphide, and 2.0 per cent. sulphur as sulphate. Push conveyors are used throughout, and samples are taken automatically as the ore is discharged from the elevators.

The fine slimes from the settler go to a settling tank, whence they are lifted up and passed through spitzluten, the heavier particles being returned to the pans. The slimes, 1 to 1, are agitated and passed to montejus or pressure-tanks, and then to filter-presses.

The four Dehne presses hold 4 tons each; the five Martin presses 3.4 tons, forming 3 in. cakes. They are worked by hydraulic pressure.

The solutions are clarified by Excelsior presses, passed through three zinc boxes, and the gold-slimes treated with sulphuric acid and melted in tilting furnaces.

The residues are dumped by a Ledgerwood hoist on to a 60 ft. heap.

Kalgoorlie Mine.—Part of the ore is dried in a White-Howell drier, the drier ore passing direct to large 200-ton storage bins.

The ore is automatically fed into six No. 5 Krupp ball-mills, crushed through a 35-mesh screen, so that about half will pass through a 120-mesh, and then led into a 400-ton bin, whence by roll feeders it is fed into nine Edwards roasters, each treating 15 tons per 24 hours.

The roasted ore falls on to a push-conveyor and is carried to a bucket-elevator, which lifts it to a mixer. The pulp from this is separated by a series of conical spitzkasten into slimes, which pass on through several pyramidal spitzkasten to have the surplus water removed; while the sands and concentrates run over copper-plates 10 ft. long and Halley tables, from which the concentrates are ground in Wheeler pans. The sands are sent to three 100-ton vats, drained of water, and bottom-discharged into steel vats below. Ordinary cyanide treatment occupies three weeks, and the cyanide solutions as they come off are returned to

* Three-hearth Merton furnaces are now being used.
the top of the sands by an air-lift, which keeps the solutions constantly circulating.

The slimes from the spitzkasten thickened up to 1 in 1 flow to a set of five pressure tanks $5\frac{1}{2} \times 13\frac{1}{2}$ ft., where they are agitated with compressed air and cyanide solution for four hours. One tank is being filled while another is being emptied and the others working, the air passing from one to the other. The agitated pulp is then filter-pressed. Cyclone settling cones are used for settling the dust, as well as an air-lift for the pulp.

**Golden Horseshoe.**—The oxidized ore is screened through 2 in. grizzlies, the coarser lumps passed through No. 3 Gates crushers, and the broken ore stored in a 200-ton bin, whence it is trucked to Challenge ore feeders supplying a 50-stamp battery of 1000-lb. stamps. The ore is wet-crushed through a 24-mesh woven wire-screen at the rate of 4 or 5 tons per 24 hours, the greater part passing a 100-mesh screen. Free gold is amalgamated both inside the boxes and outside on copper-plates, which are followed by concentrators of the Wilfley type. The coarse sands are ground and amalgamated in a series of pans followed by settlers and spitzkasten. The fine sands thus obtained are lifted by a 42 ft. tailings-wheel to vats fitted with Butters distributors. The slimes pass over 30 ft. of canvas tables for the further elimination of fire concentrates. Double treatment of the sands, occupying nine or ten days, is practised. The fine concentrates, etc., are collected and sent to the smelter.

The slimes are pumped direct, without agitators or montejus, to nine Dehne presses, each carrying fifty 3 in. cakes, pressed, washed with cyanide solution and water, and discharged. Excelsior presses are used for clarifying the gold solutions, which are then passed to four zinc extractor boxes.

Sulphuric acid treatment with filter-pressing is adopted for treatment of the zinc slimes, which are finally melted in a tilting furnace.

**Sulphide Ore.**—The new battery is of 50 stamps, each 1250 lb. The ore, after passing the Gates crusher, is brought by a Robins belt to Challenge feeders and crushed wet through a 30-mesh screen, passed over copper-plates, and lifted by a tailings-wheel to a series of grading boxes supplying Wilfley tables. The slimes which overflow from the graders, together with that from the Wilfleys, are lifted again to three hydraulic classifiers, the slimes passing to settling tanks and the sands to five flint-mills which crush them to 220-mesh, and then are again elevated to the graders and finally passed to the settling vats. The thickened slime produced in these vats is run into steel agitators, agitated with cyanide for 24 hours, pressed in six 5-ton presses, and the
waste material dropped on to Robins belt conveyors for removal. Rich sulphide ore is sent to the smelters.

**Great Boulder Main Reef.**—The Sulphide ore is treated without drying by breaking in a No. 3 Gates crusher, delivering it by a Challenge feeder on to a Robins belt, and then to two Krupp ball-mills, where it is crushed to 30-mesh, a large portion, especially when the ore is schistose, passing 100-mesh.

The crushed ore is roasted in a Richards shaft-furnace 65 ft. high, with eleven floors, and rabbled by hand, especially on the lower floors. In this furnace 35 tons per day are treated. There are also three Edwards furnaces, each of a capacity of 12 or 15 tons, which are bricked up to a constant angle.

The hot roasted ore falls into a launder carrying dilute cyanide solution, and is raised by a tailings-wheel to a spitzkasten. The sands are ground fine in Wheeler pans containing mercury, but no plates, and with a continuous overflow into another spitzkasten, from which the sands again pass to the pans, while the slimes join the first slimes, which are pumped into agitation vats. These vats are 21 ft. in diameter and 6 ft. deep. The vats are allowed to fill in 7 hours, the solution gaining in cyanide during the last foot of filling from cake cyanide then added.

After 10 hours' agitation the pulp is discharged or run off into montejus and filter-pressed in 3 in. cakes (Dehne).

The solutions are clarified by passing through a small press, and led into zinc boxes. The gold slimes are washed, pressed, treated with sulphuric acid, dried in a large iron muffle, and smelted in Cornish furnaces.

<table>
<thead>
<tr>
<th>Process</th>
<th>Hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dehne: filling in</td>
<td>1 1/2</td>
</tr>
<tr>
<td>Leaching and washing</td>
<td>1 1/2</td>
</tr>
<tr>
<td>Discharging</td>
<td>2 1/2</td>
</tr>
</tbody>
</table>

**Lake View Consols.**—Sulphides, roasted.—From Gates No. 5 crusher the ore passes to 400-ton ore-bin, and thence by aerial tram to ball-mill bin. There are four Krupp mills with a capacity of 130 tons per day, 65 per cent. of which will pass a 150-mesh screen.

Ore is fed by conveyors to four Brown straight-line furnaces 180 ft. x 10 ft., roasting 30 tons per day down to 0·2 per cent. of sulphur as sulphides. The roasted ore is elevated into 50-ton agitators into which 0·15 per cent. KCy is running, and the resulting sands are transferred into ten leaching vats, while the slimes pass on to three agitators for further agitation, before being passed through montejus and Dehne presses. Then follows zinc precipitation and sulphuric acid treatment.
Sulphides—Diehl Process.—The ore is wet-crushed, amalgamated on plates. The sandy concentrates from Wilfley tables are roasted in Edwards furnace, restamped, and ground in grit-mills down to a fine pulp. The pulp, after having been thickened by passing through spitzkasten, is treated in closed steel vats by agitation for 20 hours with bromo-cyanogen. Then follows filter-pressing and zinc precipitation.

The cost of treatment by Diehl process in 1901 amounted to 30s. per ton, including royalty 1s. 9d., water 4s. 3d., bromo-cyanogen 4s. 6d., cyanide 3s. 6d.

South Kalgoorlie.—Oxidized Ore.—The sequence of operations is as follows:—Gates crusher, Cornish rolls, Griffin mills (wet), copper-plates, grinding pans, canvas tables, spitzkasten, concentration of slimes, slimes agitation, Dehne filter-pressing.

Sulphide Ore.—Griffin mills (dry); two Brown straight-line furnaces, 150 ft. x 10 ft., with an 80-ton capacity. The roasted ore is elevated and mixed with cyanide solution, 1 to 1, and then treated in vats by the Riecken process. The vats are four in number, 11 ft. deep and about 13 ft. x 8 ft., with sloping sides and rounded bottom, holding about 18 tons of ore. KCy, 0·05 per cent. Electricity—amperage, 200; voltage, 2 to 3.

The current is supplied through iron anodes between the paddles of the agitator, which are kept moving at about 12 revolutions per minute for 18 hours. The current passes off by the copper-plates suspended along the sides, which are kept bright by a constant stream of mercury fed from pipes just above the level of the pulp, and moving backwards and forwards across the plates. The mercury is lifted by an air-lift after being drawn off below, for continuous circulation. The copper-plates are easily removable by tackle, and with the mercury from the well yield amalgam.

The pulp is drawn off by a bottom valve, and led into a storage vat 12 ft. x 6 ft., and kept agitated. Thence it is drawn off as required through montejus and filter-pressed.

The raw ore has the following composition:—

<table>
<thead>
<tr>
<th>Component</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silica</td>
<td>42·45</td>
</tr>
<tr>
<td>Alumina</td>
<td>15·77</td>
</tr>
<tr>
<td>Ferric oxide</td>
<td>9·98</td>
</tr>
<tr>
<td>Iron (in pyrites)</td>
<td>3·15</td>
</tr>
<tr>
<td>Sulphur (in pyrites)</td>
<td>3·60</td>
</tr>
<tr>
<td>Lime</td>
<td>8·94</td>
</tr>
<tr>
<td>Carbon dioxide</td>
<td>10·44</td>
</tr>
<tr>
<td>Magnesia</td>
<td>3·98</td>
</tr>
<tr>
<td>Moisture</td>
<td>0·40</td>
</tr>
<tr>
<td>Combined water, alkalies, and tellurium</td>
<td>1·29</td>
</tr>
<tr>
<td></td>
<td>100·00</td>
</tr>
</tbody>
</table>
APPLICATION OF THE PROCESS IN DIFFERENT COUNTRIES. 189

After roasting, analysis shows sulphur as sulphates, 2·82 per cent.; sulphur as sulphide, 0·32; sulphur volatilised, 0·46. Total sulphur, 3·60 per cent.

In a recent report on the Riecken process at the South Kalgoorlie mine, Mr. R. Hamilton gives the costs as follows:-

1. Coarse crushing and transport to mill, . 2 7 per ton.
2. Fine pulverising, including proportion of general charges and power, . 5 3 ,,.
3. Roasting and conveying ore, including proportion of general charges and power, . 7 0 4 ,,.
4. Agitation with cyanide, and electrical precipitation treatment, . . . . 8 5 1 ,,.
5. Filter-pressing and washing, . . . . 6 3 ,,.

Total charges, . . £1 9 7 4 ,,.

The average value of the ore for January, February, and March was £3, 17s. 6d. per ton, and the value of residues 2·125 dwt. of gold, equal to a value of 8s. 10 1/4d. per ton, representing an extraction of 88·4 per cent.

COSTS, FEBRUARY 1903.

Ivanhoe Mine.

Probably lowest costs on the field:—

Mining, . . . . . . 13 10·73
Reduction, . . . . . . 11 9·65
Development, . . . . . . 7 11·85
Capital, . . . . . . 2 3·03
General, . . . . . . 3 0·55

38 11·81 per ton.

11,098 tons for 10,318 ozs. of gold, valued at £43,800.

Lake View Mine.

Stoping, . . . . . . 9 5·6
Diehl process, . . . . 17 11
General, . . . . . . 3 6·3

30 10·9 working costs.

Mine development, . . . . 13·9 approx.
Additions to plant, . . . . 4·9 ,,.

Kalgoorlie Mine, 46s. per ton (higher than usual).
NEW SOUTH WALES FILTER-PRESS PRACTICE.

The following interesting details are extracted from a paper recently written by Mr. I. W. Rock.* The plant to which these details refer is one capable of dealing with 200 tons of dry slimes per day. A summary of the process is as follows:—

1. The dry slimes are discharged into a mixer, in which they are ground up, while cyanide solution is added, the mixture outflowing into a storage tank, whence it is elevated into agitator vats by means of a centrifugal pump. 2. Mixture agitated for a period ascertained by experiment, by some mechanical means, thus ensuring intimate contact between the metallic and chemical particles. 3. Contents run into montejus, which consist of cylindrical vessels provided with inlet and outlet valves, and also connections to the receiver of an air compressor. 4. Forcing the mixture by means of compressed air from the montejus into filter-presses, the gold-bearing solution escaping through the filter-cloths of the presses, while the solid material remains in the frames, forming large thin cakes. 5. A further extraction from these cakes by “washing” or forcing, at a high pressure, a weaker solution of cyanide through them, and, if found necessary, a second washing with water only. 6. Getting rid of the exhausted slimes by opening the presses and discharging the cakes into trucks, which convey them to a dump. 7. Clarifying the extracted gold solutions flowing from the presses, by running them through sand traps and thence into tanks, pumping them into elevated tanks, in order to obtain regular pressure, and passing them through a finer class of filter-presses, in order to retain any impalpable solid matter which would foul the zinc extractor boxes. 8. Passing the outflow from the presses into the extractor boxes and treating therein in the usual manner, the outflow being of course returned to sumps for strengthening and re-use.

Mixers.—These are made as a standard article by the engineering trade, and consist of a steel cylinder 6 feet in diameter by 5 feet deep, lined with an inserted cone, which has openings in it near the top of the outside shell, and at the bottom a four-bladed propeller on a vertical shaft, with mitre wheels and driving gear. The propeller not only cuts up the slimes and mixes them with the solution, but it drives the mixture upwards through the openings and up to the outlet, whence it overflows into the storage tank, on its way to the agitators.

Agitators.—These consist of steel vats 16 feet high and 8 feet diameter, open at the top, and with the lower portions tapered to openings which connect with both outlet and circulating pipes.

APPLICATION OF THE PROCESS IN DIFFERENT COUNTRIES. 191

To each of these agitators there is connected a centrifugal pump, whose sole duty is to draw the contents of the vat from the bottom and redeliver it into the top. This mode of mixing the particles and bringing them into intimate contact has been found highly efficient and economical in regard to time as compared with arms or paddles adopted in horizontal agitators previously adopted.

Centrifugal Pumps.—These circulating pumps, and the one for elevating the mixture into the agitators, are the only ones used for dealing with gritty material in the process. They have the usual water-pressure chambers in connection with the glands, but as water would dilute the mixture and upset the proportion, they are fed with cyanide solution under pressure from an elevated tank, supplied by a small pump.

Montejus.—These are each large enough to receive the contents of an agitator, made after the style of a compressed-air receiver, buried vertically in the ground. Inlet and outlet valves, of chemical and grit-proof make, are fitted on them; compressed-air supply and exhaust, test cock and pressure gauge, are fitted on each. In addition, there is a small air-agitation connection, so that any mixture remaining for a time in one is kept alive and the deposition of solid matter prevented.

Filter-Presses.—These are generally made by or after the pattern of Dehne, the German engineer, the usual size having fifty 40-inch × 40-inch × 2-inch frames, the total contents being 75 cubic feet. In addition to the fittings supplied by makers, there are several others which are necessary and have to be added, such as drip trays, slimes and leak launders, and several other minor accessories. Of course the discharge shoots for delivering the cakes of spent slimes, as they fall out of the frames into the trucks beneath, form part of the setting of the presses in the building prepared for them.

Wash Solution Pump.—This may be of any description of high-pressure pump with cast-iron fittings. The pressure for filling the presses from the montejus may be reckoned at up to 80 lbs., that of the wash-water at 100 lbs. to the square inch. It may be noted here that both after filling the presses and after washing the contents, compressed air is admitted for a short period to drive out any moisture.

Clarifying Presses.—These are also filter-presses, but of a much lighter description, the frames being usually made of wood. They are supposed to be self-cleansing, by reversal of the flow through them, and are so with some materials, but other slimes are so gluey that it is found necessary to have a spare set of filter-cloth frames, which can be substituted for the foul ones in
a few minutes, the latter being taken away and scrubbed. These presses are very delicate, and the pulsation of a pump destroys their efficiency. It is therefore necessary to pump the gold solution into a tank about 18 feet high, whence it flows quietly through the presses, and thence to the extractor boxes.

The items of storage vats, sumps, etc., need not be particularly specified, and it need hardly be added that the adjuncts of sufficient steam power, compressed air, and supply of water are necessary.

Practical Work.—If a plant as above described is doing its ordinary work, the routine would be as further described. Sidetipping trucks are delivered at regular intervals alongside the mixer, and the contents dipped bodily out, the fall being broken by a shoot, sufficiently flat to require the attendant to use a shovel and in some degree regulate the feed. He also, through practice, knows how much cyanide solution to run in from a supply cock, so that a fairly constant flow of mixture escapes into the storage tank.

When one of the agitators is empty, a signal is given to refill, and the elevated centrifugal does this in a few minutes from the storage tank, the circulating pump running all the time. After sufficient agitation, the man in charge of the machinery room opens the inlet valve and fills one of the montejus, the air escape cock being open. When the charge is all run in he closes the inlet valve and opens the little agitation cock until he receives a signal that a press is to be filled. When he gets that he closes the above, opens the outlet valve, and then the compressed air cock; the pressure gauge rises, and in a few minutes half the contents of the montejus has been transferred to the press in the room above. In this room the signal to fill was given when any one of the presses had been emptied and reclosed, ready for filling. All the outlet cocks on the frames are open, and the gold solution at once flows from them until the press is full of solid matter. Notice is then given to close the valves previously open and start the wash water pump; the taps are closed, the wash water outlet opened, the flow therefrom directed for a little time into the same launder as the gold solution ran into, and thereafter into a main leading to a sump for re-use. Pumping is then stopped, a little air blown through to complete the process, and the press unscrewed, the drip trays being previously removed. One after another the dummies, as the slimes-bearing frames are called, are pulled forward, the cakes either fall or are pushed out into the trucks below, each frame is scraped clean with a scalpel, and the press tightened up again for a fresh charge.

The treatment of the gold solution is pretty well automatic; its
flow through the clarifying presses has been already described, and its final delivery into the zinc boxes of the extractor-house. It will be understood that the diagram in no way indicates any arrangement of the plant. Accessibility and ample space are particularly necessary, and some details of the plant which are purchased as articles ready for use from the makers require considerable alteration to make them suit the above requirements.

Other States in Australia.—In Victoria, South Australia, and Queensland, the cyanide process is employed almost exclusively for the treatment of sands and old accumulations of tailings. Nowhere has any distinctive feature been developed except in Victoria, where charcoal is much used for precipitation purposes instead of zinc. This is really a survival and adaptation of the charcoal precipitation process formerly employed in Victoria in the chlorination process.

UNITED STATES.

The cyanide process has been successfully introduced in the states of California, Colorado, Idaho, Montana, Nevada, Utah, New Mexico, and Washington, and Black Hills district, South Dakota. Generally speaking, the adoption of the process in the States has been slow, the primary cause for this being doubtless due to the complex character of the ores. The progress of the process, however, during the past year or two has been very marked.

The process is one presenting many difficulties, especially with ores containing base sulphides, and up to the present time American metallurgists have been content to feel their way on safe ground.

The cyaniding of tailings is conducted on the same lines as elsewhere, while the slimes problem seems to have been successfully solved by the adoption of agitation followed by decantation.

The treatment of the sulpho-telluride ores of Cripple Creek districts by cyanide after roasting has been attended with much success. Up to the present time, the electrical precipitation of gold from cyanide solutions has had little or no application on a working scale. For the treatment of high-grade slimes, there may possibly be scope for the filter-press process practised in Western Australia.

The new Homestake plant, with a capacity of 1200 tons per day, and the Smuggler-Union of 600 tons per day, are among the largest in the States. Actual working details of cyanide treatment in the States are seldom available, and the following notes will be read with much interest.
Colorado.—*Camp Bird Mills, Ouray.*—I am indebted to Godfrey Doveton for the following instructive particulars of the cyanide treatment of tailings at this mill. The tailings contain a small percentage of copper, which has led to the adoption of some interesting modifications of the usual practice. Mr. Doveton says the treatment of the tailings presents no new feature, but as much of the gold is partially enclosed in the coarser particles of sand, a rather lengthy treatment is required.

The tanks are charged with Butters distributors, and the slimes overflow from three slime gates placed at the sides of the vats, and are conducted to the slime dam and settled. Slime gates are preferred to a circular launder, and as a somewhat better classification is effected, a better leaching is the result.

A sizing test of a large number of vat samples showed the following results:—

<table>
<thead>
<tr>
<th>Retained on 40 mesh,</th>
<th>. . .</th>
<th>5 per cent.</th>
</tr>
</thead>
<tbody>
<tr>
<td>60</td>
<td>. . .</td>
<td>20</td>
</tr>
<tr>
<td>80</td>
<td>. . .</td>
<td>13.13</td>
</tr>
<tr>
<td>100</td>
<td>. . .</td>
<td>15.00</td>
</tr>
<tr>
<td>120</td>
<td>. . .</td>
<td>12.80</td>
</tr>
<tr>
<td>150</td>
<td>. . .</td>
<td>17.50</td>
</tr>
<tr>
<td>Passed 150</td>
<td>. . .</td>
<td>15.62</td>
</tr>
</tbody>
</table>

Total . . . 99.05

The ore was crushed in stamper batteries through a 35-mesh wire-wove screen.

On assay, the material found on the 40 and 60 meshes was found to run considerably higher than the finer material, and the bullion button was very much finer than that resulting from the assay of the finer product.

The vats are sampled at the distributor nozzles as a check upon the vanner tailings at the stamp mill. When filled the charge is also sampled with a borer, some 40 to 50 bores being taken from a 500-ton vat. An acidity test for free and combined acid is made on the head sample, and the requisite quantity of lime is found, and added on top of the charge, and well mixed by shovelling over the top layer of material.

*Details of Treatment of 100-ton Charge.*—The tank being partially drained and lime added, 20 tons of a weak solution are run on, containing about 0.05 per cent. KCy. This solution, which also contains a considerable percentage of copper cyanide, acts as a food for the cyanicides, and also, owing to the presence of much cupro-cyanide, dissolves much of the copper contained in the charge, thereby leaving the material in a better condition for the actual
working solution. No gold is dissolved as a rule by the preliminary weak solution, and as the whole of the available cyanide is destroyed, it is allowed to drain off and run down the sluice launders, to be saved for sluicing purposes, etc.

It should be mentioned that on rare occasions, when a portion of the gold contents is amenable to treatment, from 10 to 25 cents value of gold is removed by the first solution.

The weak solution is immediately followed by a succession of 10-ton charges of solution of 0.25 per cent. free KCy, until the outgoing solution rises to about 0.20–0.225 per cent. KCy. Usually 60 tons of solution are run on, and by the time the outgoing solution is up to near standard strength, 65 to 70 per cent. of the gold is dissolved. The charge is now allowed to macerate from 8 to 16 hours, and is subsequently rapidly drained, sampled, and shovelled to the tank below for the second treatment. During the shovelling the charge sample has been assayed, and the gold yet capable of extraction ascertained. Usually almost all the soluble AuKCyo compound has been carried out by the charges of solution. Should the assay show that much of the previously insoluble gold still remains in the ore, standard solution is added in the lower vat. Should the gold contents be unusually high, 0.3, 0.35, or 0.4 per cent. solution is used for the saturation of the shovelled charge. If, however, a fairly good extraction has been obtained in the upper vat, 0.25 per cent. solution is used, and when we are crowded with tailings weak washes are applied immediately after the saturation.

Mr. Doveton continues:——“Our usual practice is, however, to saturate with 0.3 per cent. solution, and allow to macerate from 4 to 6 hours, then drain rapidly, and apply a couple of washes of 0.3 per cent. solution of 10 tons each, drain partially dry, and give 25 tons of weak solution (0.08–0.125 per cent.). Before the last of the weak washes has disappeared below the surface, the charge is allowed to macerate two hours. This ensures all of the dissolved gold being carried out.

“The charge is now drained dry on surface, and water washes added till cyanide values fall to 0.04 per cent. The liquor is now transferred from the weak gold tank to the waste gold tank, and the solution, low in cyanide, is passed through the waste zinc boxes, and thence out of the mill. The gold value of the solution at the time of transfer from weak to waste zinc boxes is about 40 cents per ton, and much of this is precipitated in the waste zinc boxes.

“At the time of sluicing, the charge is fairly dry, containing about 12 per cent. of moisture, and the outgoing solution usually contains about 0.02 per cent. of free KCy, but only.
little double cyanide. As is usually the case, the first weak solution flowing from the charge is high in double cyanide, which we are now about to recover by our recovery process.

"It is noticeable that but little gold is carried out by the solution till the outgoing strength is almost up to standard strength of cyanide.

"A series of solution assays were made on a number of vats during treatment. Samples of the outgoing solution were taken hourly, and the samples for each six hours assayed together. It was found that up to 0·05 per cent. KCy the solution carried scarcely any gold at all. Between 0·05 and 0·1 per cent. the value increased from 40 cents to 150 cents per ton; from 0·1 per cent. to 0·15 per cent. KCy it was about 225 cents; and at 0·2 to 0·25 per cent. KCy the value remained about 5 dollars per ton of solution. The value seems to remain pretty constant till the solution falls to 0·2 per cent. KCy, when a gradual decrease sets in, and at 0·05 per cent. KCy the value is usually 80 cents per ton, and at 0·02 per cent. KCy the value is seldom more than 25 cents.

"There is a considerable quantity of copper in the solution, but nevertheless we have a most perfect precipitation. Our aim is always to precipitate the copper on the zinc with the bullion, thus keeping the copper contents of the solution at a constant figure.

"Our strong sump assays for over a year have seldom gone as high as 40 cents, the average being from 15 to 20 cents. Our weak sump assays show but an average of usually from 2½ to 4 grains of gold per ton by the assay of very large samples, say 50 assay-tons.

"The precipitation is usually practically perfect the week following the clean-up. With our waste zinc boxes we are not quite so successful. The copper contents of the incoming solution are usually high, and the gold value only some 25 to 45 cents per ton. The available cyanide is very low, varying from a trace to 0·025 per cent., with the total cyanides 0·05 per cent.

"By coating the zinc with mercury by immersing it in a weak solution of mercuric cyanide—made from mercuric chloride and KCy—we are enabled to make a very fair precipitation.

"When the weak solution runs rather lower than usual in gold value, more of the copper is precipitated on the zinc, and a somewhat less perfect precipitation of the gold takes place. The addition of concentrated cyanide solution at the head of the bin does not seem to help the precipitation of the bullion at all, but greatly accelerates the plating of the copper. The practice of strengthening the solutions entering the zinc boxes has been entirely discarded; it has been found here most advantageous to
promote the precipitation of the copper along with the bullion, thus preventing the solution becoming overcharged with copper. Occasionally, after placing fresh zinc in the last compartments of the boxes, it will be perfectly coated with copper in about eight hours, because the solution, when it reaches the last compartment or two, is very low in bullion value. However, in the course of 24 hours, when perhaps a higher bullion value is entering the boxes, the gold will be plated over the copper, the precipitate being a very lustrous black.

"Should the coppery zinc show no signs of precipitating bullion, precipitation is readily aided by the use of a few Winchester quarts of strong mercuric cyanide solution, which is allowed to drip into the compartment affected. When the zinc becomes gray in colour from the amalgamation of the deposited mercury with the coppery zinc, the addition of mercuric solution is discontinued, and in a few hours the zinc will become black from bullion precipitate.

"The evolution of hydrogen is much increased, should the solutions contain more copper than usual, and on several occasions, when the solution entering the boxes carried as much as 0.045 per cent. Cu, and the usual bullion contents, 4 or 5 dollars per ton, the evolution of hydrogen was so great as to lift the zinc partially out of the boxes; however, a good bullion precipitation was obtained, the solution leaving the boxes worth only 5 grains of gold per ton. An example of this was noticed recently, when a complete analysis of a working solution showed the presence of a small quantity of manganese, about 0.00778 per cent., estimated on the evaporation of 2 litres of solution by Volhardt's method. No sulphates or magnesia were found, but lime was present to the extent of 0.01387 per cent. of Ca. There was found a little lead, considerable alumina, no ferrocyanide, but a large amount of sulphocyanide, of the latter about 0.1078 per cent. There was a little iron present, but in what combination could not be determined.

"No alkaline sulphide has yet been detected in the solutions here, and it would seem that when zinc is used as the precipitating agent no alkaline sulphide can exist, the zinc being readily precipitated as sulphide from the ZnK₄Cy₄.

"The oxygen is frequently estimated in the solution by a modification of Thresbis' iodimetric method. The estimation is very valuable, as it enables us to ascertain whether our solutions are sufficiently aerated or not. Numerous oxygen tests showed the presence of from 2 to 7.36 mgrms. of oxygen per litre of solution.

"Experiments and numerous estimations were made to test the merit of employing jets of compressed air in the sumps and
storages, should it be found that the solutions received insufficient oxygen during their passage from the end of the zinc boxes to the leaching vats. However, we found that in the majority of the cases the solutions were capable of dissolving but a very little more oxygen on aeration, and the greater portion of that dissolved was diffused again on standing for any length of time.

"The weak solution was found, on the average, to contain a little more dissolved oxygen than the strong, the increase being most noticeable when the weak solution leaves the zinc boxes. Here the weak solution containing 0.08 per cent. KCy and 0.11 per cent. total KCy contained 0.64 mgrm. per litre of dissolved oxygen, while the strong solution contained only 0.4 mgrm."

Cost of Cyanide Treatment at Camp Bird Mills.
5000 tons Tailings monthly.

<table>
<thead>
<tr>
<th>Description</th>
<th>Cents per ton</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cyanide, 1 lb. per ton,</td>
<td>27.00</td>
</tr>
<tr>
<td>Zinc, 0.55 lb.</td>
<td>5.00</td>
</tr>
<tr>
<td>Lime, 1.8 lb.</td>
<td>1.30</td>
</tr>
<tr>
<td>Assays</td>
<td>3.60</td>
</tr>
<tr>
<td>Labour</td>
<td>18.15</td>
</tr>
<tr>
<td>Power—steam,</td>
<td>6.00</td>
</tr>
<tr>
<td>Sand-pumps, maintenance,</td>
<td>4.32</td>
</tr>
<tr>
<td>Electric light,</td>
<td>1.20</td>
</tr>
<tr>
<td>General repairs,</td>
<td>0.40</td>
</tr>
<tr>
<td>Supervision, etc.,</td>
<td>4.60</td>
</tr>
</tbody>
</table>

Total,                     71.57

Doveton gives the total charges as about 3s. per ton of ore treated, including 2.4 cents per ton for sulphuric acid treatment of gold slimes.

Smuggler-Union Mine.—The cyanide plant at this mine was erected last year. It has a capacity of 600 tons per day. Zinc precipitation is used.

Dorcas Pneumatic Cyanide Mill, Florence.—This plant has a capacity of 120 tons per day. The treatment here is of some interest, as compressed air is used to aerate and agitate the solutions and pulp in the leaching vats. The following description is extracted from a paper by Dr. Wells.*

The ore, averaging 20 dollars in value, passes through crushers and coarse rolls to the sampling room, thence through a revolving dryer and through two sets of finishing rolls which crush it to

* The Engineering and Mining Journal, Jan. 4, 1902.
24-mesh size. The finely crushed ore is then roasted in a Holt- 
hoff-Wethey furnace, the roasted product passing to the leaching 
tanks.

There are six of these tanks, 30 ft. in diameter, 4.5 ft. deep, and 
fitted with air pipes in the bottom for the introduction of air 
during the leaching. The air is supplied at a pressure not 
exceeding 5 lbs., about 1 cubic foot of air per minute to each ton 
of ore being sufficient for agitaiton and oxygenation.

The treatment generally lasts five days, and leaves a value of 
about 1 dollar per ton in the ore. The residue is then sluiced 
out and concentrated on Wilfley tables. The tailings after this 
treatment average about 40 cents per ton.

The tanks are filled by a conveyor, and as soon as the bottom 
of the tank is covered with pulp a solution containing 10 lbs. 
KCy per ton is run in gradually, the pulp at the same time 
continuing to flow in until the tank is full. The air is then 
turned on gradually, and is kept on until the pulp shows an extrac- 
tion of at least 90 per cent. Whenever the air comes up un-
evenly through the charge the ore has to be stirred by men with 
iron rods. This generally requires about 30 hours. The air is 
then shut off and the pulp allowed to settle for one hour. 
Percolation is then begun and the strong solution run off as 
quickly as possible, followed by a weaker one of 5 lbs. KCy per 
ton. This operation is continued until the tests of the solution 
show only traces of gold. Water is then added to displace 
the KCy solution. The loss in cyanide is stated to be less than 
1 lb. per ton. The dust resulting from the dry-crushing is 
collected and treated with cyanide, without previous roasting. 
From dust assaying 51.20 dollars per ton the tailings only contained 
80 cents per ton. It is added to the roasted ore in the leaching 
tanks, 3 tons to each tank, spread evenly on top of the charge.

_Cripple Creek Telluride Practice._—The country rock of this 
district is andesite breccia, phonolite, and decomposed granite. 
On the surface the ores are oxidized and carry iron peroxide, 
manganese oxide, and oxide of tellurium. Below water-level, the 
gold occurs in the minerals calaverite and sylvanite, and is always 
associated more or less with iron pyrites.

The gold in the surface ores is free, but not amenable to 
amalgamation, being coated with metallic oxides. It is, however, 
easily extracted by cyanide.

The unoxidized telluride ores have to be subjected to a pre-
liminary dead roast before cyanide treatment.

The reduction of the ore is generally effected with Krom 
rolls, instead of the mills used at Kalgoorlie, the ore being crushed to 
pass through a 40-mesh screen. The roasted ore is leached by
percolation in vats with two solutions of cyanide. The stronger, containing from 0.5 to 0.75 per cent. of potassium cyanide, is allowed to percolate for 50 hours, after which the weak solution is added. The time of treatment varies from 70 to 100 hours.

Precipitation is effected with zinc.*

The treatment of the sulpho-telluride ores of Western Australia is obviously a more difficult problem than that of Cripple Creek ores. At Kalgoorlie, the ore is talcose and is highly calcareous, and forms so large a proportion of slime that ordinary percolation is impossible, and decantation too slow and costly. The successful solution of the difficulty was found in the use of filter-presses, in which the dissolution of the gold is effected under pressure, and by means of which the slimes are easily separated from the gold-containing solutions and washes.

As at Cripple Creek, the oxidized ores are treated in the raw state, while the undecomposed ores are subjected to a dead roast before cyanide treatment. With the adoption of filter-presses, it was soon recognized that the greater the proportion of slimes the better, and to attain that object, pulverizing mills, instead of rolls, have come into general use.

California.—California King Mines.—The plant at this mine has a capacity of 1000 tons per day. The ore is crushed by rolls to 20-mesh size and then treated in ten cyanide vats, each 40 ft. in diameter and 5 ft. deep, provided with bottom-discharge doors. Zinc precipitation is used.

There are many cyanide plants scattered throughout the State, including the large 140-stamp cyanide mill of the Golden Cross Mines at Hedges, but no particulars of these are available.

Montana.—Dr. Wells says that cyaniding has made great progress in this State. Many new plants have been erected, some of them of large size.

The process as carried on in this State is much the same as elsewhere. In many of the mills, the method is wet-crushing, amalgamation on plates, followed by cyaniding of tailings. In other cases the ore is crushed in rolls and cyanided direct. With slurry ores, the practice is to agitate and decant.

Nevada.—The sun-drying slime treatment adopted at the Dexter plant at Tuscarora has already been referred to in the chapter dealing with slime treatment. Of the other plants in operation in this State some are treating tailings, while others are dealing directly with dry-crushed ore.

New Mexico.—The Cochiti Company of Bland has a successful

* Prof. Furman, Mines and Minerals, January 1897.
method of treatment which possesses some novel features.* Here 98 per cent. of the dust will pass a 100-mesh screen. The process used is agitating and settling, the agitation being effected by steam and compressed air, the latter being under 60 lbs. pressure. Steam is used to heat the charge, as it was found that the air on expanding cooled it too much. The mixture of air and steam is admitted through 1 in. horizontal iron pipes in the bottom of the tank. The pipes have 0.0625 in. holes in the underside for the escape of air and steam. In 24 hours 94 per cent. of the total values are in solution, using a solution of cyanide of 3 lbs. per ton, but only 80 per cent. is recovered, due to the solution going to waste in the slimes. To neutralize the acidity of the ore and to aid the settling of the slimes, lime is suspended in a wire basket in the upper part of the tank. The cost of treatment is given as 1 dollar per ton.

Utah.—Mercur is one of the greatest centres of the cyanide process in the United States, the different plants having a capacity of over 2000 tons per day. The Mercur ore has already been described at the end of Chapter VI. when dealing with Coarse Crushing and Cyaniding.

The Golden Gate Cyanide Works.—These are said to be the largest and best equipped in the United States.† They were constructed in 1898, and are built on a hillside with eight levels. In order to get the ore to the top of the works it has to be hoisted on an incline 800 ft. long. The mill is 294 ft. wide, and 420 ft. in length up and down the slope. The difference in elevation from top to bottom is 145 ft. The retaining walls, which are 2 ft. wide at the top, and have a batter of 1 ft. in 12, required over 50,000 cb. yards of rubble masonry. The various floors were constructed by blasting out the hillside. The broken stone thus obtained was used for the retaining walls and filling behind them. The mill is driven by power transmitted electrically a distance of 35 miles at a tension of 40,000 volts. The loss of energy in transmission is said to be only 5 per cent.

At the works the 40,000 volt 3-phase current is transformed to one of 220 volts of 2-phase. The current is delivered at a contract price of 60 dollars per H.P. The first section of the mill contains the coarse crushers, and in the second are the dryers. In the third section is the fine crushing machinery, which consists of four sets of 26-in. rolls, and three sets of 36-in. Berthelet apparatus are used for sizing. There are six elevators, with a lift of 60 ft. The fourth, fifth, and sixth sections contain the roast-

* Hunter, Engineering and Mining Journal, 19th Jan. 1901.
† Excerpt from Bosqui's Cyanide Process, p. 174.
ing furnaces, which are of Brown's straight-line design, four in number. Those intended for arsenical ores are estimated to have a daily capacity of 75 tons, while those for talcose ores are rated at 150 tons. The ore is stirred by the rabbles once each minute. One man attends to two furnaces. The gases are carried from the furnaces through 6 by 8 ft. flues into the main dust chamber, which connects with a steel stack, 8 ft. in diameter and 85 ft. high, located on the hill above the buildings. The top of this stack is 275 ft. above the lowest level of the building. The leaching department, which constitutes section 7, is 60 by 294 ft. It has two floors, the main floor supporting ten tanks 25 by 50 ft. and 5 ft. deep (presumably rectangular), and three solution tanks 20 ft. in diameter and 12 ft. deep. The tanks are supported by masonry piers. They are charged by hand from cars run on bridges over the tanks. The eighth section of the mill, which is 50 by 70 ft. and two stories in height, is the precipitation department. It contains three precipitation tanks, each 14 ft. in diameter and 8 ft. deep. The tailings from the leaching tanks are discharged into cars which are run to the waste dump. The building is constructed of steel.

**South Dakota.**—The following details of cyanide plants and practice are extracted from a paper contributed by Mr. Chas. H. Fulton, M.E., to the *Engineering and Mining Journal*, January 4, 1902.

**The Homestake Company's 1200-ton Tailing Plant at Lead.**—This plant takes the tailings pulp from the Golden Star, Eighty, and Highland stamp mills. The pulp is conveyed to the cyanide plant in a 10 in. cast-iron pipe. On its way it passes through the classifier house, where some of the slimes are separated by means of eight large sheet-iron cone-shaped classifiers. These cones have no upward or rising current, but as the ore pulp is charged by pipe at the centre, the sands settle to the bottom and are discharged, while the slimes overflow at the periphery. The sands are then carried to the cyanide plant, where they are fed to seven additional cones and treated as above described. The slimes overflowing at the periphery are again discarded, while the sands are discharged to four-compartment jigs of the Harz pattern, which act as classifiers. One cone feeds four jigs. The jiggling is done on beds of pyrites concentrates, relatively coarse screens being employed, except in the last compartment. The stroke varies in the different compartments from \( \frac{1}{2} \) in. to about \( \frac{3}{4} \) in. The last compartment is widened out in order to compensate for the increased amount of discharge water which accompanies the slimes during their carriage from the tanks. The continuous hatch discharge is taken by open launder to the revolving pipe distributors, which
fill the leaching tanks, the peripheral overflow of the tanks still carrying off slimes. Lime in the form of an emulsion is added to
the pulp launder which takes the jig discharge.

There are fourteen leaching tanks, 45 ft. in diameter, 9 ft. deep, built of California red-wood. The tanks are placed seven in a row. There is one revolving distributor of the Butters type for each row, being moved from tank to tank by overhead track, as required. To get an even overflow at the periphery of the tank, the tops of the staves are grooved to a depth of \( \frac{1}{2} \) in., and a soft pine tongue is inserted, projecting from \( \frac{3}{4} \) to \( \frac{3}{4} \) in. above the top of the staves. This is then easily planed, and can be kept perfectly level. It is also readily renewed.

It is the object to charge just as much slime with the sands as possible, and still leach in a reasonable time. For this purpose the adjustments are made on the jigs and distributors. The value of the pulp as it leaves the stamp mills is from 1.25 dollars to 1.50 dollars per ton. The amount crushed in the mills approximates 2000 tons per day, while the cyanide mill treats about 1200 to 1300 tons per day. The difference, or 700 tons, is discarded as slimes which, while carrying considerable value, cannot be treated in the condition they are in, and must be discarded. The proportion of slimes separated out is about 35 per cent. The sands are leached directly in the filling vats.

The Cyanide Plant of the Wasp No. 2 Mining Company.—This plant on Yellow Creek, near Kirk, is a dry, coarse crushing plant, capacity 90 tons per day. The ore is quartzite, the gold being on the cleavage planes, and in the seams and cracks, so that coarse crushing only is necessary to make it available for solution. The value varies from 3 dollars to 12 dollars per ton, the main bulk averaging about 4 dollars. It is mined very cheaply (85 to 90 cents) by open-cut mining. The mill is built in terraces on a very steep hillside, and the ore is moved through almost entirely by gravity. It is hauled from the mine 750 ft. distant in trains of four cars, and charged into the main bin, from which it passes to a No. 2 D. Gates crusher over a 4 by 8 ft. grizzley having 1\( \frac{1}{2} \) in. squares between bars. The undersize and the crushed ore from the Gates go to the storage bin, from which it is fed by Tulloch feeders into one set of coarse Gates rolls 14 by 24 in., 80 revolutions per minute. The rolls discharge to a stationary inclined 2-mesh screen 7 ft. long, 1 ft. wide, the oversize from which passes through the finishing rolls 14 by 24 in., operated at 100 revolutions per minute, and meets the undersize, the combined product being elevated by the one-bucket elevator in the mill to the shaking finishing screens, situated on a level with the coarse rolls. This screen is an inclined shaking 2\( \frac{3}{4} \)-mesh screen, 16 ft. long and
2 ft. wide, the lower half being stationary. Nearly all of the finished product passes a 6-mesh screen, the oversize from which is returned by gravity to the finishing rolls.

The ore is charged from the finished product bin into the four leaching tanks of Oregon fir; each tank is 16 ft. diameter, 7 ft. deep, and of 55 tons capacity. For this purpose a 16 in. belt conveyor is used at a speed of 600 ft. per minute. A large 100-ton tank has recently been added, which gives the mill the capacity mentioned above. Tanks are charged in from two to two and a half hours, the cyanide solution being added after the tank is about one-half filled. The strength of the strong solution is 6 lbs. cyanide per ton of solution. About 15 lbs. of strong solution are run on, and usually allowed to stand three to four hours, then it is drained, a weak solution being added at top to replace it. After this complete replacement it is allowed to stand a short time and then circulated for the balance of the leaching. The weak solution contains from $2\frac{1}{2}$ to 3 lbs. cyanide per ton. About 50 tons of weak solution are used, followed by from 6 to 9 tons of wash water. To neutralize the acidity of the ore and give a protective alkalinity, 6 lbs. of lime per ton are added before the ore is dumped into the first bins. The cost of cyaniding is from 85 to 90 cents per ton. The precipitation is accomplished by zinc thread in specially constructed barrels, and very good results are obtained, the tailings solution having a value of from 4 to 10 cents per ton. The extraction made is from 80 to 90 per cent. of the values, and the consumption of the cyanide amounts to about $\frac{1}{2}$ lb. per ton of ore. The precipitates are treated by the usual sulphuric acid method. Zinc dust precipitation in the tanks, pumping the resultant mixture through filterpresses, was tried at the mill, but discarded on account of the difficulty in the clean-up.

MEXICO.

El Oro Mine.—This mine is situated about 100 miles northwest from the city of Mexico. The ore is gold-bearing quartz, containing a small amount of silver. The country-rock is slate, which is covered with heavy flows of andesite.

The most important lode is the “San Rafael,” which strikes a few degrees west of north, and dips westward at an angle of 63°. The average width of this great lode is some 50 feet, and it has been traced for more than a mile in length. The ore is somewhat similar to that of the “Martha” lode at Waihi.
APPLICATION OF THE PROCESS IN DIFFERENT COUNTRIES. 205

Up till the beginning of 1906 only oxidized ore had been treated. It contained a small percentage of pyrites.

The treatment is amalgamation on plates in the stamp mills, and cyaniding of tailings which are classified in cone pulp-thickeners giving two products, namely, sands and slimes. The sands are subjected to fine grinding in tube-mills, further separated into sands and slimes, which are treated separately with cyanide.

For the above and following particulars I am indebted to an instructive paper by G. Caetani and E. Burt * on "Fine grinding of Ore by Tube-mills, and Cyaniding at El Oro, Mexico."

In Mill No. 2, the newest of the two mills, the ore is crushed by 100 stamps, each weighing 1140 lbs., and dropping 6 in. 100 times per min.; the discharge is 2.5 in. through 25-mesh brass wire screen (0.74-mm. aperture). The duty per stamp is 4.7 tons per 24 hours of actual work. The ratio of mill-water to ore is from 9 to 10.

The ore is dumped from skips into ore-bins, crushed through gyratory and jaw crushers, and thence conveyed to the mills by cars, or by belt-conveyors. The mills are of the standard Allis & Chalmers pattern. The ore is stamped through from 25- to 35-mesh screens, and runs over amalgamated plates, which catch from 13 to 18 per cent. of the gold and from 1 to 5 per cent. of the silver. The bullion from the retorted amalgam contains about equal parts by weight of silver and gold.

Owing to the scarcity of water in the district, the mill-water, after separation from the slime, has to be pumped back to the large storage-tanks. About 340 tons of mill-water pass daily into the cyanide solutions, being carried in as moisture in the slime, which contains 53 per cent. of water, and in the sand, which contains 15 per cent. of water. The discharged slime carries 40 per cent., and the sand 15 per cent., of water; therefore, about 75 tons of weak solution have to be discharged daily.

The pulp from the plates is classified by two 4.5-ft. cones put in series with two 2-ft. cones, called "pulp-thickeners," giving two classified products, namely, an overflow containing about 81 per cent. of slime, and representing 52 per cent. of the total mill-output, and a discharge containing 9 per cent. of slime.

The discharge of the pulp-thickeners is run through tube-mills, where from 40 to 50 per cent. of the sand is slimed, according to the various conditions under which the tube-mills are working.

The pulp discharged by the tube-mills joins the overflow from the four cones and forms a final product, containing, on an average, 69 per cent. of slime. The product is then classified, the final result being to separate two mill-products—sand and slime; the former containing about 20 or 25 per cent. of slime, and representing about 35 per cent. of the rock crushed, and the latter containing about 5 or 10 per cent. of sand.

**Tube-mills.**—At present there are three tube-mills in use at the new plant constructed by the Fried. Krupp Aktiengesellschaft Grusonwerk, of Magdeburg, Germany, and the principal data concerning them are as follows:—

<table>
<thead>
<tr>
<th>Tube-mill.</th>
<th>No. 3</th>
<th>No. 4</th>
<th>No. 5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Internal diameter of drum</td>
<td>47 in.</td>
<td>59 in.</td>
<td>59 in.</td>
</tr>
<tr>
<td>Internal length of drum</td>
<td>19 ft. 6 in.</td>
<td>23 ft.</td>
<td>26 ft.</td>
</tr>
<tr>
<td>Revolutions per minute</td>
<td>32</td>
<td>27.5</td>
<td>28.5</td>
</tr>
<tr>
<td>Pounds of pebbles in mill when half full</td>
<td>11,700</td>
<td>21,800</td>
<td>24,700</td>
</tr>
<tr>
<td>Capacity per 24 hrs., tons</td>
<td>100–120</td>
<td>130–150</td>
<td>150–180</td>
</tr>
<tr>
<td>Cost of mill</td>
<td>$2940</td>
<td>$3850</td>
<td>$4750</td>
</tr>
</tbody>
</table>

Bulk of pebbles = 100 lbs. per cu. ft.

The tube-mills are set on concrete foundations and held in place by anchor-bolts. Experience thus far at El Oro indicates that the foundations cannot be made too solid. The mills, when in motion and heavily loaded with pebbles, bear very heavily on the foundations, which, owing to the mechanical arrangement of the mills, cannot be made on the top as wide as is desirable. The cast-iron bed-plates have a tendency to dig into the cement, and it is advisable to enlarge the bearing-surface of the bed-plates by placing larger cast-iron plates under them. A piece of rubber belt placed below these plates will be found to be advantageous. The vibrations produced by the spur-gears are violent, and the anchor-bolts easily crystallize and break; therefore pockets are advisable. The driving-pinion ought to revolve in such direction as to lift the gearing of the tube-mill, and press down on its own bearing. On the contrary, it is continuously pulling on the anchor-bolts of the boxes, thus tending to loosen the nuts and to break the bolts.

Of the three tube-mills in use, No. 3, the smallest, has given most satisfaction and least trouble. Its capacity is not much below that of the other two; but the cost of running it is much
smaller. The main advantage is that the mill is comparatively light and compact, and therefore seldom gets out of order. The stopping of a tube-mill means a great loss of money, hence reliability is of greater importance than capacity.

The pebbles used are the so-called Danish pebbles, costing £7, 13s. 4d. per ton at the mill, or over three farthings per lb.

Tube-mill No. 3 shows an average consumption of 3·7 lbs. of pebbles per ton of sand passing through. Tube-mill No. 5 shows a consumption of 3·9 lbs., and No. 4 of 9·2 lbs. The average consumption for the last four months has been 5 lbs. per ton.

The liners are made in the foundry of the company, and last as long as the imported ones, and cost about one-half as much. The consumption of liners is approximately about 1·2 lbs., per ton of sand ground.

During the month of August, 5170 tons of sand were run through the tube-mills, out of 10,773 tons crushed by the stamps, and 72 per cent. of these 5170 tons were returned to the tube-mills to be reground. The total amount of sand ground was 8890 tons.

The cost of grinding by the tube-mills was calculated per ton of sand passing through the tube-mills during a four months' run.

<table>
<thead>
<tr>
<th></th>
<th>No. 3</th>
<th>No. 4</th>
<th>No. 5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total cost per ton</td>
<td>s. d.</td>
<td>s. d.</td>
<td>s. d.</td>
</tr>
<tr>
<td>(Steam)</td>
<td>1 4½</td>
<td>1 6½</td>
<td>1 7½</td>
</tr>
<tr>
<td>Total cost per ton</td>
<td>1 1</td>
<td>1 1½</td>
<td>1 2½</td>
</tr>
<tr>
<td>(Elect. power)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

It is found that the cost of grinding fine sand is greater than the cost of grinding coarse sand, owing to the reduced efficiency of the mills, and the greater wear and tear of pebbles and liners, in the former case.

Slime Treatment.—The slime-pulp, overflowing from the hydraulic classifiers, is composed of about 1 part of slime to 12 parts of mill-water. The pulp flows directly to the steel treatment-tanks, 12 ft. in height and 34 ft. in diameter, entering at the centre through a wooden box, so arranged as to break the force of the flow. Caustic lime is added to the mill-pulp, just before it enters the tube-mills, in the proportion of 12 lbs. of lime per ton of ore. The slime settles completely, and the clear mill-water overflows and is pumped back to the mill storage-tanks.

Each slime-charge consists of from 62 to 85 tons (average 76
tons) of dry slime; and it takes about 12 hours to collect the charge when 100 stamps are crushing through 30-mesh, and when one tube-mill alone is used for regrinding the sand; while it takes about 9 hours when 40-mesh battery-screens are used, and when two tube-mills grind and regrind the coarse sand; and it takes only 5½ hours when crushing through 25-mesh, and keeping three tube-mills at work.

When a charge is completed, the slime is allowed to settle for 8 hours, at the end of which a 76-ton charge will have caked down to a depth of 50 inches. The pulp has an average specific gravity of 1·39, and contains about 53 per cent. of water.

The clear mill-water is decanted as the settling takes place, thus avoiding any delay when the settling is considered to be complete. Lately, a larger percentage of very fine sand has been sent over with the slime, causing a far greater rate of settling of the slime.

The pulp is then set in agitation by a mechanical agitator, consisting of a horizontal wooden cross revolving in the centre of the tank, 14 in. from the bottom, and at a speed of 7 rev. per minute. Two arms of this cross are 15 ft. long, and two are 7 ft. long. When the whole pulp is once set in rotation it takes about 6 horse-power to keep up the movement.

Ten minutes after the agitation has begun, a valve at the bottom edge of the tank is opened, and the pulp put into circulation by a centrifugal pump, which elevates the pulp and throws it back to the centre of the tank. Compressed air is fed near to the inlet of the pump to aerate the pulp. These centrifugal pumps do excellent work; but the cost of operating them is high, as they consume 14 horse-power. The pump has to run 14·25 hours for the complete treatment of each charge of slime. Reckoning the h.p. year at £30, the total cost of the pump-agitation is 7d. per ton of slime.

When the pulp has been thoroughly mixed, it is sampled and the specific gravity is determined, so as to calculate the weight of each charge. Cyanide solution is then added to the pulp, and the treatment of the slime carried on.

Up till July 1905 the treatment consisted of four washes, but is now somewhat modified.

Experimental working tests were made to ascertain the action of lead acetate and mercuric chloride in accelerating the dissolution of the gold and silver in the slime charges. In every test it was found that the added salt had a beneficial action, no doubt due to the desulphurizing of alkaline sulphides formed by the dissolution of the silver sulphites contained in the ore.

*Modified Slime Treatment.*—The experimental tests proved that
by using lead acetate or mercuric chloride at the beginning of the first wash the extraction of the gold could be almost completed in 12 hours, and that 60 per cent. of the final extraction of the silver could be obtained during the same period of time.

The tests also showed that the silver-extraction continues during the following operation of washing-out the values by repeated settling and decanting, almost independently of the fact whether the slime is agitated or not.

It was, therefore, decided to modify the slime-treatment on slime-plant No. 2. The outline of the treatment is as follows:—

After decanting the mill-water from the already settled slime-charge, the pulp is stirred up and mixed with about its equal weight of unprecipitated solution, so as to bring the total ratio of solution to slime to about 2:5 to 3. While the solution is flowing into the tank, the final strength of the solution is brought up to from 0.045 to 0.05 per cent. of KCy, by hanging the necessary amount of NaCy in a sack just under the flow of clear solution entering the tank. When there is an excess of strong solution in the sand-treatment plant, as after heavy rains, it is usual to pump some strong solution into the slime-tanks, so as to bring up the strength of the weak solution. Lead acetate (0.4 lb. per ton of slime) is dissolved at the same time and by the same method.

After the required volume of solution has been added to the charge, which takes about 1 hour, the double agitation by pump and stirrer is kept up for 7 hours. Then fresh, unprecipitated solution is added, so as to fill the tank completely. The agitation is then stopped, and the slime allowed to settle. After decanting the first solution as closely as possible, four additional washes are given to the charge as rapidly in succession as the setting of the slime will allow; and, with the last one, the pulp is pumped over to the deep settlers.

The actual extractions obtained by this modified treatment on a charge of 61.5 tons were as follows:—

<table>
<thead>
<tr>
<th>Component</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gold</td>
<td>92.1 per cent.</td>
</tr>
<tr>
<td>Silver</td>
<td>76.3</td>
</tr>
<tr>
<td>KCy consumed per ton of slime</td>
<td>0.90 lb.</td>
</tr>
<tr>
<td>Lead acetate per ton of slime</td>
<td>0.46 lb.</td>
</tr>
<tr>
<td>Or mercuric bichloride</td>
<td>0.20 lb.</td>
</tr>
<tr>
<td>Quantity of solution used</td>
<td>813 tons</td>
</tr>
<tr>
<td>Quantity of solution precipitated</td>
<td>459 tons</td>
</tr>
<tr>
<td>Total time of treatment in slime-tank</td>
<td>50 hours</td>
</tr>
<tr>
<td>Lime used per ton of ore crushed</td>
<td>12 lbs.</td>
</tr>
</tbody>
</table>
To diminish the amount of solution to be precipitated in the zinc-boxes, the first two solutions are unprecipitated solutions from the deep settlers. The value of such solutions only averages 7½d. in gold and much less in silver per ton, and therefore does not affect sensibly the final value of the last wash-solution.

It was customary at El Oro to keep the ratio of solution to slime as high as possible during the treatment. But the tests described in the preceding pages have proved that a rapid and good extraction can be obtained just as well when this ratio is as low as 2½. The advantage of using a small bulk of solution during the first treatment is, that less cyanide and less lead acetate are needed to bring the solution to the desired degree of concentration.

Naturally, in washing-out the values, the ratio of solution to the slime has always to be kept as high as possible, to get a good extraction, because the capacity of the plant is not great enough to allow a great number of washes to be used.

The total cost of slime-treatment at the new mill (No. 2) for the months of July and August in 1905 were as under:—

<p>| | | | | |</p>
<table>
<thead>
<tr>
<th></th>
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<th></th>
<th></th>
<th></th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>July</td>
<td>5</td>
<td>3½</td>
<td>per ton of slime.</td>
<td></td>
</tr>
<tr>
<td>August</td>
<td>5</td>
<td>1</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Palmarejo Mine.**—The ore at this mine consists of a silicious matrix, containing a small percentage of pyrites. Manganese oxides and calcite are present in varying proportions, also traces of antimony, arsenic, and bismuth, with occasionally a little copper and zinc. It contains from 2 to 5 dwts. of gold and from 15 to 30 ozs. of silver per ton, and may be described as an argentiferous gold ore. The silver exists chiefly as argentite.

The ore, in its physical character, mineral constituents, and valuable contents, is almost identical with the argentiferous gold ores which are so successfully cyanided in the Auckland district, New Zealand, more especially those in the Crown, Talisman, Woodstock, and Komata and Waitekauri mines.

These particulars are gleaned from an instructive paper by T. H. Oxnam * on "Cyaniding Silver-Gold Ores of the Palmarejo Mine, Chihuahua, Mexico," from which it is seen that the ore is successfully treated by wet-crushing in a 50-stamp mill, and concentration, followed by the cyaniding of sands by percolation, and of slimes by agitation and decantation.

BRITISH COLUMBIA.

Athabasca Mine, Nelson.—In a paper read before the Mexican meeting of the American Institute of Mining Engineers in 1901, Mr. E. Nelson Fell gives some interesting information on the treatment of tailings at this mine. The ore, he says, consists of a quartz gangue, containing a little lime and variable quantities of the sulphides of iron, lead, and zinc. The following figures, giving the analyses of the ore before milling, and of the tailings after milling, which constituted the material to be cyanided, are based on the daily samples taken during February 1901.

*Analyses of Ore and Tailings.*

<table>
<thead>
<tr>
<th>Element</th>
<th>Ore Per cent</th>
<th>Tailings Per cent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zn</td>
<td>1·93</td>
<td>0·91</td>
</tr>
<tr>
<td>Fe</td>
<td>7·04</td>
<td>2·65</td>
</tr>
<tr>
<td>Pb</td>
<td>1·63</td>
<td>0·20</td>
</tr>
<tr>
<td>CaO</td>
<td>4·97</td>
<td>2·46</td>
</tr>
<tr>
<td>S</td>
<td>5·99</td>
<td>1·71</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>3·20</td>
<td>3·20</td>
</tr>
<tr>
<td>SiO₂</td>
<td>74·20</td>
<td>85·00</td>
</tr>
<tr>
<td>Au</td>
<td>1·68</td>
<td>0·32</td>
</tr>
<tr>
<td>Ag</td>
<td>1·32</td>
<td>0·38</td>
</tr>
</tbody>
</table>

Oz. per ton. Oz. per ton.

The analysis shows this to be an ore well adapted for cyanide treatment.

The plant was located to receive the tailings direct from the mill, in two distributing-tanks, 14 ft. in diameter and 10 ft. in height. The tanks are fitted with annular launders around the rim, and are filled with water before the admission of the tailings; while the overflow is carried off in annular launders.

A slimes-arrester is fixed in each vat, consisting of a sheet of iron, 10 in. wide, fitted inside each vat, about 1 in. from the staves, and extending all the way round. This ring of sheet iron is thus 2 in. less in diameter than the diameter of the vat, and is held in position by eight iron brackets so arranged that the ring can be raised entirely above the level of the tank, or lowered until the upper edge is about level with the top of the staves, according to the nature of the ore being treated. The obvious intention of using this contrivance is to retain as much of the slimes with the sands as possible.
The slimes-arrester described by Mr. Fell is exactly the same as that attached to the J. C. Fraser and Price's continuous grinding and amalgamating pans, which were extensively used in Australia and New Zealand for the treatment of mill tailings before the introduction of the cyanide process.

The overflow takes place in the space between the ring and the staves, and having such a great length of outflow, is very even and quiet.

The plant, which has a capacity of 50 tons per day, consists of five leaching-vats, each 18 ft. in diameter and 4 ft. deep, fitted with centre discharge doors, and two collecting-vats, each provided with the ordinary revolving reaction distributors. The latter are 14 ft. in diameter and 10 ft. deep, and so situated above the leaching-vats, that each one can be discharged by shovelling through side-doors into any one of three adjacent leaching-vats.

Besides these there is a strong gold tank, a weak gold tank, a waste-water vacuum tank, and two series of zinc extractors, each consisting of twelve iron boxes, which are square and have each a capacity of one cubic foot of zinc-shavings. The boxes are movable, and each is independent of the other. The two gold tanks are 10 ft. in diameter and 6 ft. deep, and the two sumps 12 ft. in diameter and the same depth.

Mr. Fell supplies the following working details:

"This description shows the actual details of treatment, and the results obtained from the treatment of Lot 59, which gives a fair example of the treatment.

"June 26th, at noon, turned on strong solution (0·24 per cent.) until same stood 6 in. deep on the sands; allowed to stand 4 hours. At 4 P.M. opened outlet-cock and allowed solution to run into gold-tanks; at 4.30 P.M., as the last of the solution was passing off, took sample, which assayed "nothing" in cyanide and 9·30 dollars in gold. Closed outlet-cock, admitted fresh charge of strong solution, and allowed to stand 8 hours. At 12.30 (midnight), June 27th, opened outlet-cock, and at 1 A.M. took sample, as before, which assayed 0·06 cyanide and 28·94 dollars in gold. At 1 A.M. turned on weak solution (0·08 cyanide), allowing same to drain through without interruption till 4 A.M. Sample at 4 A.M. assayed 0·10 cyanide and 8·08 dollars in gold. Shut off outlet-cock and allowed solution to stand until 1.30 P.M. Opened outlet-cock and admitted fresh solution (0·06 cyanide) and allowed same to run through, admitting fresh solution as required till 12.30 (midnight), June 28th. Sample taken at 10 P.M. assayed 0·07 cyanide and 0·62 dollar in gold. Closed outlet-cock and allowed to stand until 5 A.M. Opened outlet-cock at 5 A.M. and allowed fresh solution to run through until 11 A.M. Sample
taken at 6 a.m. ran 0·06 cyanide and 0·42 dollar in gold; and sample taken at 11 a.m. ran 0·6 cyanide and 0·21 dollar in gold. At 11 a.m. turned in water-wash till 2 p.m. Sample taken at 12.30 p.m. ran 0·06 cyanide and 0·21 dollar in gold; and sample taken at 2 p.m. ran 0·06 cyanide and 0·21 dollar in gold. At this point the process was declared finished; the wash was drained to waste; and the tailings were discharged.

"Assay of the tailings before treatment gave 13·02 dollars, and after treatment 2·07 dollars in gold. Percentage of recovery, 84·1. Time occupied, 2 days 2·5 hours."

If the actual recovery is equal to the extraction obtained by the assay difference, the results obtained must be considered very satisfactory, and with prolonged leaching could doubtless be increased.

**THE CYANIDE PROCESS IN INDIA.**

Laurence Pitblado states* that in the Kolar field there are at present six cyanide works in operation. The ores of the field are very simple, consisting mainly of pure quartz, with only a small percentage of iron pyrites. The material treated is tailings from the stamp mills. Those first worked in the Mysore plant averaged 4·5 dwt. gold, and yielded 65 per cent., with an average consumption of 1 lb. cyanide per ton. In 1897 a month’s test with 40-mesh screens in the batteries gave the following result: — 90·65 per cent. amalgamated in batteries and on plates; 74 per cent. of the value in the tailings recovered by cyanide lixiviation, making a total extraction of 97 per cent. of the ore as delivered to the mill. The cost of cyaniding at the present small plant of the Mysore Company is £2, 10s. 5d. per ton, exclusive of royalty and depreciation, but in the new 4000-ton plant which is being erected alongside the heap of accumulated tailings that is to be worked, it is believed that the cost will not greatly exceed 2s. At the Champion Reefs mill, where 20-mesh screens are used in the batteries, the extraction from the tailings is about 56 per cent. with a consumption of 1 lb. of cyanide per ton.

In refining the precipitate in this district it is first passed through a 30-mesh screen, drained, dried, roasted with or without a small percentage of nitre, and fused directly in plumbago crucibles. At the Mysore works the precipitate is retorted before roasting, yielding about 100 lbs. of mercury per month. The presence of mercury in the zinc boxes generally leads to the production of much-floured and brittle zinc. In fluxing and smelting

* * Journal of the Society of Chemical Industry, Feb. 28, 1898."
directly, the retorted slimes the bullion assayed 56·4 per cent. gold, 3 per cent. silver, 2·4 per cent. lead, 19·6 per cent. copper, 18 per cent. zinc, and 0·1 per cent. nickel. The slag contained a good many shots of metal. In roasting with nitre a slag freer than the above was produced, and a bullion assayed 49·5 per cent. gold, 4·6 per cent. silver, 4·29 lead, 38·21 per cent. copper, 2·10 zinc, and 0·22 per cent. nickel. These results being unsatisfactory, the following practice was adopted at the Mysore works. The retorted and dried slimes are mixed with 10 per cent. of nitre and roasted at a bright red heat. When cold they are boiled with dilute sulphuric acid (1:2), which dissolves the copper. The dried and washed product is fluxed with about 35 per cent. borax, 15 per cent. soda, and 10 per cent. sand, giving a slag free from shots of metal and a bullion assaying 81·3 per cent. gold, 6·9 per cent. silver, 2·71 per cent. lead, 6·78 per cent. copper, 0·4 per cent. zinc, 0·12 per cent. nickel.
CHAPTER XII.

THE SIEMENS-HALSKE PROCESS

The distinguishing features of this process are the use of extremely dilute solutions of cyanide and the electrical precipitation of the gold. Since the introduction of the cyanide process, the precipitation of the gold by metallic zinc has always been regarded as a weak point; and metallurgists have devoted much time in the endeavour to discover an efficient substitute for it.

Electrical precipitation naturally engaged the attention of many investigators. In 1893, the author, assisted by Mr. F. B. Allen, M.A., B.Sc., conducted a number of experiments with electrical precipitation to determine the method of precipitation to be adopted at the School of Mines cyanide plant. Many different modifications were tried. With some, the precipitation from solutions of ordinary working strength was very satisfactory; but, with all, the precipitation of the gold from dilute solutions, such as those corresponding to weak cyanide washes, was always very imperfect and accompanied by decomposition of the water.

In the Siemens-Halske process this difficulty is overcome by causing a slow artificial circulation of the cyanide solutions in the extractor.

The plant and operations connected with the leaching of the gold are the same as those described in the preceding chapters, the only difference being in the extractor-house.

The electrical precipitation of gold has been introduced with marked success at a number of cyanide plants at the Witwatersrand Goldfields, and its use is extending. Up to the present it has not been introduced to New Zealand or Australia, and so far very little has been written about it. For the following details of the process I am indebted to the papers of Mr. Charles Butters and Mr. A. Von Gernet, read before the Chemical and Metallurgical Society of South Africa, and published in the South African Mining Journal.
Discovery of the Process.—Mr. Von Gernet said the electrical precipitation of gold extracted from ores by cyanide has been in use in Europe and Asia as far back as 1888. In 1887, Dr. Siemens found that the gold anodes used in electroplating at his works in Berlin lost weight when standing idle in the cyanide solution, without any electric current passing through the bath. This, in connection with the well-known fact that gold was soluble in aqueous solutions of cyanide, first induced him to try the use of that solvent for the extraction of gold from ores.

In the same year he built a small plant to make experiments on concentrates produced in Siebenburgen. The gold was precipitated both by electrolysis and zinc filings. It was found, however, that the zinc method gave good results only from comparatively strong solutions, while the electrical precipitation was effected with both dilute and strong solutions, and its efficiency was not affected by the presence of caustic soda.

Dr. Siemens therefore decided to use electrolysis only, and early in 1888 he commenced operations on a large scale. Engineers were sent to different countries, two going to Hungary, one to America, and one (Mr. Von Gernet) to Siberia.

The operations were generally successful, and in May 1894, a plant, capable of treating 3000 tons of tailings per month, was erected at the Worcester mine, near Johannesburg. During 1895 the process was adopted by some eight or ten large mining companies, including the Metropolitan, May Consolidated, Croesus G.M. Co., No. 4 Central Works, and Robinson Slime Works; and already it is a formidable rival of the McArthur-Forrest zinc-precipitation process.

Action of the Electric Current on the Cyanide Solution of Gold.—The electric current decomposes the auro-potassic solution, depositing the gold on the negative pole and liberating the metalloid at the positive pole. In a fixed time a given electric current will deposit a certain quantity of metal, which quantity will vary for different metals in direct proportion to their electro-chemical equivalents. This law holds good only for solutions strong in metal; but with very dilute solutions, as in use in the cyanide process, the current does not find sufficient of the metallic compound present at the electrodes, and consequently decomposition of water also takes place; for this reason, to make the precipitation as efficient as possible, constant diffusion of the solution is required.

The artificial circulation of the solution is most economically and conveniently obtained by allowing a slow but steady flow through the precipitation boxes. It is of the highest importance to give a very large surface to the electrodes, since a more efficient
precipitation is obtained by doubling the number of plates than by increasing the current tenfold.

The Cathode or Negative Electrode.—To obtain a satisfactory cathode, a metal must be used which will fulfil the following conditions:—

1. The precipitated gold must adhere to it.
2. It must be capable of being rolled out into very thin sheets to save unnecessary expense.
3. It must be easy to recover the gold from it.
4. It must not be more electro-positive than the anode, in order to prevent return currents being generated when the depositing current is stopped.

The most suitable metal was found to be lead, which, in the form of foil, meets all the requirements, and is therefore used in the Siemens-Halske process.

The Anode or Positive Electrode.—The requirements of the anode are no less important. By the action of the current a metalloid is liberated at the positive electrode, and the latter, when a metal, begins to oxidize. Carbon could be used, but it will not withstand the action of the current, soon crumbling into a powder which decomposes potassium cyanide. Besides, when this finely-divided carbon is in suspension, it cannot be removed from the solution by filtration.

When zinc is used as an anode, it forms a white precipitate of ferro-cyanide of zinc by the reaction of zinc oxide upon ferrocyanide formed during leaching. In the same way, iron anodes form Prussian blue by the reaction of oxide of iron and ferrocyanide. In consequence of this reaction, the amount of ferrocyanide in the cyanide solution does not increase.

The cyanide can be recovered from the Prussian blue, formed at the iron anodes, by dissolving it in caustic soda, then evaporating the solution, and finally smelting with potassium carbonate. Mr. Von Gernet states that this process has been tried on a small scale, about 50 lbs. at a time, with the result that a nice clean potassium cyanide was obtained. In the treatment of clean tailings, this regeneration of cyanide is not of great importance; but with concentrates, or tailings, which decomposes the cyanide solutions with formation of ferro-cyanide, it will effect a considerable saving.

Electric Current Required for Precipitation.—Only a very weak current is required to precipitate the gold from cyanide solutions, a density of 0.05 ampere per square foot being sufficient. With cathodes 1 ½ in. apart, 7 volt is sufficient to produce this strength of current.
The advantages gained by using such a weak current are:—

1. The gold is deposited hard on the lead-foil.
2. The iron anodes are preserved for a long time, as their waste is in proportion to their current strength. In a plant treating 3000 tons per month, 1080 lbs. of iron are destroyed in that period.
3. Little power is required. 746 Watts equal 1-horse power. A 3000-ton plant requires 2400 Watts, equal, theoretically, to 3½-horse power, and actually requiring about 5-horse power.

The Advantages of Electric Precipitation.—The principal advantages claimed for this process are as follows:—

1. That the precipitation operates independently of the amount of cyanide or caustic soda present in the solution. Therefore, in the treatment of tailings, very dilute solutions can be used, the only limit being a sufficient quantity of cyanide to dissolve the gold satisfactorily.

A solution containing 0·03 per cent. of cyanide will dissolve gold as effectively as a solution containing 0·3 per cent., provided a longer time is allowed for treatment. In the first case, the decomposition of cyanide is much less than in the second, resulting in a corresponding economy.

2. However acid the solution may be when entering the extractor, the precipitation takes place equally as well as it does when the solution is neutral or alkaline.

3. No complications arise from the formation of lime, alumina, or hydrate of iron, which sometimes cause trouble in the zinc process of precipitation.

4. With ores or tailings containing copper, the extraction of the gold will be the same, but the decomposition of cyanide less than when using stronger solutions.

5. The successful treatment of slimes.

The Actual Working of the Process.—The first practical demonstration of this process on a commercial scale took place at the cyanide works of the Worcester Gold Mining Company, near Johannesburg, under the supervision of Mr. A. Von Gernet.

The plant consists of five leaching vats placed on a row of stone piers, with a single tunnel beneath. Each vat is 20 ft. in diameter, with 10 ft. staves, and has a capacity of 100 tons of tailings.

Between the vats and the electric extractors there are placed two tanks, 16 ft. in diameter, with 6 ft. staves, forming two intermediate reservoirs, which enable the flow through the pre-
cipitation boxes or extractors to be kept constant and steady, a matter of great importance.

A better method to secure an even flow is to pump all the solution into a small raised tank, provided with an overflow into the intermediate tank and a delivery pipe to the precipitation boxes. The small tank is always kept full to overflowing, so that it delivers under a constant hydraulic head.

Beyond the precipitation boxes there are two sumps, 20 ft. in diameter and 6 ft. deep, from which the cyanide solutions are returned to the leaching vats.

Two collecting vats, 20 ft. in diameter and 8 ft. deep, receive the tailings from the 25-stamp battery.

The Electric Precipitation Boxes.—There are four precipitation boxes, constructed of wood, each 18 ft. long., 7 ft. wide, and 4 ft. deep. Each box contains 89 iron-plate anodes, 7 ft. by 3 ft. by \( \frac{1}{4} \) in., cased in canvas to retain the small quantity of Prussian blue produced; and 88 cathodes of lead-foil stretched on iron wires fixed on a wooden frame. Each frame contains three strips, 3 ft. by 2 ft., so that, counting the double surface of each lead-sheet, there are altogether about 3000 square feet of cathode surface, the current density being 0.05 ampère per square foot. Copper wires are fixed along the top of the sides of the boxes, and convey the current from the dynamo to the electrodes.

The boxes are made of 3 in. material throughout, with stiffening pieces across the sides and bottom. The divisions are of wood, or are formed by raising some of the iron plates about an inch above the level of the solution, while others rest right down on the bottom, the joints being made water-tight by means of wooden fillets caulked with hemp packing. By this means a series of compartments is obtained, similar to those in a zinc precipitation box, the difference being that the solution passes alternately up and down through successive compartments. The rate of flow is about one foot per minute.

The Clean-up.—The boxes are kept locked, being opened once a month for the "clean-up," which is conducted as follows:—The frames are taken out singly, and the lead-foil is removed and replaced by fresh lead-foil, the whole operation taking but a few minutes for each frame. The lead, which contains from 2 to 12 per cent. of gold, is then smelted into bars and cupelled.

The gold is deposited on the lead sheets as a thin bright yellow film, which adheres firmly to the lead. The consumption of lead at the Worcester Works is 750 lbs. per month, equal to 1\(\frac{3}{4}\)d. per ton of tailings.

The working expenses for treating 3000 tons per month were as follows:—
| Filling and discharging leaching vats, | \( d \) | 10·00 per ton |
| Cyanide, \( \frac{1}{4} \)-lb., | | 6·00 " |
| Lime, | | 1·20 " |
| Caustic soda, | | 0·50 " |
| Lead, | | 1·10 " |
| Iron, | | 2·20 " |
| White labour, | | 5·20 " |
| Native labour and food, | | 1·90 " |
| Coal, | | 4·60 " |
| Stores and general charges, | | 3·30 " |

**Total,** | | 36·00 per ton of 2000 lbs. |

The cost of treatment per ton of 2240 lbs. would be 3s. 4·32d.

The tailings assayed from 6 dwt. to 8 dwt. of gold, and the residues, after treatment, from 1 dwt. to 2 dwt. per ton. The average actual extraction was about 74 per cent.

The solutions, after leaving the precipitation boxes, still contained gold, the strong solution showing by assay 4 dwt. 8 grs., and the weak solution 10 grs. per ton of solution. On the average, the strong solutions carried from 4 dwt. to 5 dwt., and the weak from 10 grs. to 1 dwt. of gold per ton of solution.

From November 1894 to May 1895, the Metropolitan Company treated 26,900 tons of tailings for 4845 ozs. of gold, at a cost of 2s. 8d. per ton. At the May Consolidated the working expenses amounted to about 2s. 4d. per ton, excluding the royalty for the use of the process, which amounts to 3 per cent. The extraction amounted to over 80 per cent. of the original assay value.

**Details of the Treatment.**—The time occupied in leaching and washing, together with the quantity of the solutions, are given in the following tabulated statement:

| Alkaline wash, 10 tons, | | 3 |
| Strong cyanide solution, 70 tons, 0·05 to 0·08 per cent. KCy, applied in 14 separate portions of 5 tons each, | | 65 |
| Weak cyanide solution, 21 tons, 0·01 per cent. KCy, applied in 3 portions of 7 tons each, | | 18 |
| Water washes, total 11 tons, pumping dry and discharging, | | 22 |

**Total,** | | 108 |

The working of this process gives rise to the production of a number of valuable commercial bye-products, including copper, lead, litharge, and paint.
CHAPTER XIII.

OTHER CYANIDE PROCESSES.

The Diehl Process.—This is an adaptation of the Sulman-Teed process. It embraces the following essential stages *:

1. Crushing and sliming the ore.
2. Treating the slimes in agitators with a solution of potassium cyanide in combination with cyanogen bromide.
3. Filter-pressing the sludge.
4. Precipitating the gold from solution by means of zinc.

According to the character of the gold and its associates, amalgamation and concentration can be added to the process.

The most direct advantage of this process is, that the ore is treated in a "raw" condition.

At Hannan Star mill, where the process was first introduced, the ore is dry-crushed in two No. 5 Krupp ball-mills provided with 30-mesh screens. The crushed ore is elevated, mixed with water, and classified into sands and slimes. The sands are conducted to copper-plates on which from 10 to 15 per cent. of the gold is saved. From the plates the pulp travels by gravitation to the fine-milling department, where it is ground into slimes in a large Krupp tube or flint mill, which is an 18 ft. long steel cylinder, 4 ft. in diameter, charged with 4 tons of flint balls. The sand is fed into one end and issues at the other of such a fineness that the whole product will pass through a sieve with 200 mesh per lineal inch.

From the grinding mill the pulp is led to settling vats, from which the surplus water is returned to the mixing machine. The thickened pulp is now led into agitators, where it is treated with a solution of cyanide of potassium and bromide of potassium.

When the agitator has received its full charge, a strong solution of potassium cyanide is added. For slimes containing 1 to 3 oz. of gold per ton, we have found it sufficient to add so much cyanide that there will be 4·4 lbs. KCy per ton of dry material. After the

sludge has been agitated for 1 to 1½ hours, the solution of bromide of cyanogen is allowed to flow in, the quantity added being 1·1 lb. per ton of dry material. The agitator is kept going for 24 hours from the time the KCy solution was charged into it. In case the sludge should contain more than 3 oz. per ton, it may be advisable to add, after 6 to 8 hours' agitation, a further quantity of KCy and BrCy, to ensure a good extraction. On the other hand, if the sludge contains less gold than 1 oz. per ton, the quantity of KCy and BrCy can be considerably reduced.

About 2 hours before the agitator is ready to discharge to the filter-press, lime is added to the sludge in quantity varying from 1 lb. to 4 lbs. per ton of dry slimes. In the most cases, I think, 3 to 4 lbs. is used. A cleaner precipitate is thereby obtained in the zinc boxes.

After agitation, the pulp is pressed in Dehne's filter-presses, in which the dissolved gold is washed out. After washing, the dry cakes of slime are thrown out on the dump.

The gold in the solutions is precipitated with zinc turnings in the ordinary way. It is claimed that an extraction of over 93 per cent. can be obtained. The consumption of water is very small in this process, which is an important factor on a goldfield, where water is so scarce. All the available water is salt or brackish, which doubtless tends to retard close concentration.

Mr. Feldmann gives the following summary of the costs for the month of July 1901, when 2210 tons of ore were treated at Hannan's Brown Hill, at Kalgoorlie, by this process:

<table>
<thead>
<tr>
<th>Summary</th>
<th>Per ton.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>s.    d.</td>
</tr>
<tr>
<td>Milling</td>
<td>4     0·59</td>
</tr>
<tr>
<td>Concentration</td>
<td>1     7·19</td>
</tr>
<tr>
<td>Treatment of concentrates</td>
<td>1     4·08</td>
</tr>
<tr>
<td>Extraction</td>
<td>17    1·02</td>
</tr>
<tr>
<td></td>
<td><strong>24   0·88</strong></td>
</tr>
</tbody>
</table>

The Schilz Process.—This process is based on the use of barium peroxide, BaO₂, which it is claimed under certain conditions parts with one-half of its oxygen. It is further claimed that the normal oxide of barium, which is left after the treatment, performs further functions—inasmuch as it decomposes the sulphate of iron in the solution, forming sulphate of barium and oxide of iron, both insoluble; it removes sulphocyanides, and it dispenses with the use of lime and answers the purpose of the
same, with other minor advantages in addition. On the other hand, the process requires much longer time to effect a satisfactory extraction than in ordinary cyaniding, thus necessitating a considerable increase in vat capacity.

Herr Schilz states that to ensure success the peroxide must be well sprinkled with the tailings whilst the vat is being filled, so that a good mixture may be obtained. A rather stronger cyanide solution than usual is run in, and the whole allowed to stand undisturbed for at least three days. The longer the time the better is the extraction, and in the case of concentrates it should be at least a week. After running off the gold solution the residue is washed with four weak cyanide solutions, the first of which should remain six hours. Then, again, a strong solution should be applied, followed by weak solutions.

The quantity of peroxide depends less on the percentage of gold than on the length of time during which the solution remains in contact with the charge; and, further, the more pyrites present the greater the consumption of peroxide. The vat should not be filled to the brim with the tailings, as the mass swells up by the evolution of gas, sometimes as much as a foot. Thus the liberated oxygen should remain undisturbed as long as possible in contact with the charge, the greatest solvent action of the cyanide being after a few days. The quantity of peroxide required varies with the material under treatment.

In actual practice the quantity varies from $\frac{1}{3}$ lb. to 1 lb. per ton of ore treated. Delivered on the Rand, the cost of barium peroxide is £50 per ton.

A working trial of the process at the City and Suburban mine, according to the report of the manager, showed a gain of about 13 per cent. in extraction at a small additional cost—that is, a rise from 76 to 89 per cent. The treatment of the pyrites concentrates gave also satisfactory results.

The inventor makes the following claims for his process:

1. $\text{BaO}_2$ decomposes in contact with moistened tailings into $\text{BaO}$ and $\text{O}_2$, and supplies the cyanide of potassium with oxygen required for the dissolution of gold. This decomposition is performed in the cold.

2. $\text{BaO}_2$ supplies the oxygen which is required for the decomposition and dissolution of the reduced pyrites, and thus assists in liberating the enclosed gold.

3. $\text{BaO}_2$ in decomposing supplies $\text{BaO}$, which, being a strong basis, exerts a purifying and clearing influence upon the cyanide solutions, thus increasing their solving power.

4. $\text{BaO}_2$ renders superfluous the use of lime, the now existing use of which has in its train so many injurious secondary effects.
TREATMENT OF CUPRIFEROUS ORES AND TAILINGS.

Copper carbonates and sulphides are readily soluble in solutions of cyanide, and when present in ores, even to the amount of one-half per cent., they cause an excessive consumption of available cyanide, besides rendering the precipitation of the gold on zinc very imperfect.

Many valuable gold ores contain sufficient copper to render the extraction of the gold by ordinary cyaniding impracticable. For this cause the successful treatment of such ores is a metallurgical problem of increasing importance.

Among the methods which have been proposed for the treatment of cupriferous ores are the following:—

The Park-Whitaker Cyanide Process.—This process is intended for the treatment of cupriferous ores and concentrates which cannot be treated successfully by the ordinary cyanide processes, on account of the solubility of copper ores in cyanide solutions.

In this process the ore is subjected to a chloridizing roasting, after which the soluble copper chlorides are removed by leaching with water. An alkaline wash is then applied, and the gold and silver extracted with a dilute solution of cyanide.

During the roasting the silver sulphides present are converted into chloride, which is readily dissolved by cyanide. The dissolved copper is recovered by passing the solutions through iron turnings or scrap-iron.

Experiments on a working scale were made by the author on a parcel of ore from the Monowai mine, N.Z., with most successful results.

Scrymgeour's Process.—The copper is first dissolved out of the ore by a cupriferous cyanide solution, and the excess of copper in the solution recovered by electrolytic precipitation, a partial regeneration of the cyanide being said to take place. The residual sands are then cyanided in the usual way, and the gold recovered from the solution by electrolytic precipitation.

Hirsching's Process.†—The copper ore is dissolved out by means of ammonia, the sands washed, and then treated by cyanide as usual. The copper and ammonia are recovered from the solutions by distilling off the ammonia and leaving the copper behind as an oxide precipitate.

Bertram Hunt's Process.‡—The ore or sands are treated direct by a solution of potassium cyanide to which ammonium

* Engineering and Mining Jour., 20th Dec. 1902, p. 816.
hydrate has been added. The gold is thus extracted together with some copper, and the metals are recovered by electrolytic precipitation, the gold, silver, and copper falling to the bottom of the vats as a sludge. Zinc precipitation may also be used.

**Jarman-Brereton Process.**—In this process the authors recommend the use of KCy plus ammonium chloride for the extraction of gold in the presence of copper carbonate.

It is claimed that in extracting gold from oxidized cupferous ores, the consumption of solvent would not be large, and the solutions less cupferous than those obtained in ordinary cyanide treatment, thereby greatly modifying, if not wholly overcoming, the precipitation difficulty when using zinc shavings.

This and other ammonium processes was made the subject of exhaustive investigation by Professor Jarman and his colleague Mr. Le Gay Brereton in 1904. The results of their experiments were placed before the Institution of Mining and Metallurgy * in February 1905, and elicited a valuable contribution on the chemistry of ammonium processes from H. Livingstone Sulman.† The practical utility of all the ammonia-copper-cyanide processes is somewhat problematical.

**The Pneumatic Cyanide Process.**—In this process the dissolution of the gold is accelerated partly by jets of compressed air, which cause a continuous and gentle agitation of the pulp, and partly by the aeration of the solution caused by the passage of the air through the charge.

The compressed air is conducted to the charge through a coil of perforated pipe placed on top of the filter-cloth.

This process is identical with the Park-Horn process, for which provisional letters patent were obtained in New Zealand in 1895. Tests were made by Mr. George Horn and the author at Kuaotunu goldfield on excessively fine slimes. The dissolution of the gold was rapid and almost complete, but the mechanical difficulty of separating the gold-containing solutions from the slimes gave so much trouble that further attempts were abandoned. For the treatment of sands, ordinary percolation was found to give satisfactory results without artificial aeration.

**Gilmour-Young Process.**—This process has been operated on argentiferous gold ores in Central America. It is essentially an amalgamation process, closely following the Washoe method of treatment. The addition of a caustic alkali, copper and zinc amalgam, and mercury to the pulp containing cyanide solution must promote many complicated reactions of doubtful utility.

ELECTRO-CHEMICAL SLIME PROCESSES.

These are based on the dissolution of the gold with cyanide, and the electro-precipitations of the gold on mercury, which also serves to amalgamate any gold too coarse to be dissolved by cyanide.

The Riecken Process.—This process has lately been installed at the South Kalgoorlie mine, Kalgoorlie, and since December 1900 is reported to have treated 5000 tons of slimes. At present the capacity of the plant is being doubled. The following details of the process are extracted from a description supplied to the Australian Mining Standard in 1901.

In this process the pulverized and, if necessary, roasted ore is agitated in a vat of special construction with cyanide solution; a current of electricity is passed through the resulting pulp, by which the gold dissolved in the cyanide solution is deposited on the amalgamated sides of the vat as amalgam, while at the same time any coarse particles of gold in the ore too large to be dissolved by the solvent are mechanically amalgamated. Thus, in one operation, in a single vat, are simultaneously performed—(1) amalgamation of the coarse gold; (2) solution by cyanide of the fine gold; (3) electrical precipitation of this dissolved gold. The pulp, being thus deprived of its precious metals, is at once discharged from the vat to residues dams, and the treatment is finished.

The apparatus in which this operation is performed may be regarded as a huge electro-depositing cell, which is an iron vat with vertical ends, inclined sides, and a rounded bottom. Its dimensions are 13 ft. 4 in. long, 8 ft. 3 in. wide at the top, and 11 ft. deep. It holds a charge of 18 tons. Agitation of the pulp is effected by paddles attached to a horizontal shaft working through stuffing boxes. The paddles are 2 ft. 8 in. long, 5 in. wide, and spaced 12 in. from centres. The ends of paddles reach about 2 in. from the bottom of vat.

The sides and bottom of vat are lined with movable amalgamated copper-plates, $\frac{1}{10}$ in. thick, which form the cathodes or negative poles.

When in operation a thin sheet of mercury constantly descends over the copper-plates from a $\frac{3}{4}$ in. pipe perforated with $\frac{1}{16}$ in. holes, spaced 6 in. centres. A slow reciprocating motion is imparted to these perforated iron pipes, causing them to swing through a chord of 6 in. about twenty times per minute. The mercury finds its way to the bottom of the vat, whence it is drawn off by a trap and elevated by an air-lift to the upper receptacle, whence it again flows into the vat, thus ensuring a continuous circulation.
OTHER CYANIDE PROCESSES.

The anodes are bars of iron 1 in. thick by 3 in. wide, suspended from two girders and curved so as to be parallel to the cathodes. There are twelve cathodes in the vat.

The agitator running about twelve revolutions per minute is found sufficient to keep the pulp from settling. The salt in the local water renders the conductivity of the pulp excellent, so that a current of very low potential is quite sufficient to effect the precipitation of the dissolved gold. The current required for each vat of 18 tons is about 250 amperes of a potential of 2.5 volts, equal to about six-sevenths of an electrical horse-power, the cost of which per ton is merely nominal.

The ore is first pulverized until all the gold is liberated from its matrix. If necessary the ore is roasted. In order to obtain good results it is merely necessary that the gold should be amalgamable, or in such a fine state as will permit of dissolution by cyanide solutions. Necessarily each particular ore requires a certain specific preliminary preparation, determinable by experiment, to obtain the best results.

The pulp, containing approximately equal weights of ore and solution, is discharged into the vat previously described, and which is called the "electro vat." The agitator is set in motion, the electric current from the dynamo is started, and the flow of quicksilver is maintained by means of the air-jet, as already described. The solution is made up of potassium cyanide to the average strength of 0.075 per cent. With a high-grade ore of 2 oz. or over the above-described operation is continued for about eighteen hours, while with low-grade slimes, carrying but 3 dwt. or 4 dwt., it may be complete in about six hours. After a proper time has elapsed the discharge-valve at the bottom of the vat is opened and the pulp is allowed to flow to the residues dam; the vat is refilled as before, and the operation indefinitely repeated. If water is scarce, the supernatant liquor, after the pulp has settled in the dam, may be pumped back into the mill to make pulp for succeeding charges.

The clean-up is effected by withdrawing one copper plate at a time, and replacing it with a spare one. The amalgam produced is very fine, and contains about 27 per cent, of fine gold. In this process sodium is added to the mercury automatically to keep it active and bright.

The Rieckeen process dispenses with percolation, filter-pressing, or decantation for the removal of the dissolved gold from the ore, and entirely does away with zinc-precipitation and clean-up, the complete treatment of the ore, after its mechanical preparation, being effected in one operation in one vat, in the minimum of time, the gold being recovered as amalgam, requiring only retort-
ing and melting. The future applications of this process will be watched with much interest by metallurgists.

The Keith Electro-Cyanide Process.—The Keith electro-cyanide process is the invention of Dr. Keith, an American electrician. The process consists of two parts. First, the process for dissolving the gold out of the crushed ore; and second, the recovery of the gold from the solution. Dr. Keith’s improvement in the dissolving process consists in adding to the solution of potassium cyanide a certain amount of cyanide of mercury. In practice, he states that he finds the best results are obtained when the solvent contains 0·05 per cent. of potassium cyanide and 0·025 per cent. cyanide of mercury. This mixture of cyanides, it is claimed, operates very much faster than the simple potassium cyanide.

The process for the recovery of the gold from the solution is an electrolytic one. The gold and the mercury are deposited together upon amalgamated copper-plates. The amalgam so deposited is scraped off and the gold recovered by distilling off the mercury in the usual way. The anode is not allowed to dip into the cyanide solution, but is placed in a separate compartment and surrounded with a solution of an alkaline salt, so that the cyanide does not become decomposed. The electro-motive force of the current need not be more than half a volt.
CHAPTER XIV.

ANTIDOTES FOR CYANIDE POISONING.

All cyanides are deadly poisons; but the aqueous solutions used in practice are so dilute that there is little or no danger from the prussic acid evolved from them if the buildings are properly ventilated.

Acids react on cyanides, liberating prussic acid gas, which causes almost instant death when inhaled in a pure state. When diluted with air, it causes faintness, dizziness, and a depressing frontal headache.

Even very dilute solutions of cyanide are poisonous when taken internally, and, when they come in contact with the skin, produce, in some persons, an eruption of painful red boils. In cases where the hands and arms must be brought into contact with the solution, rubber gloves, reaching over the elbows, should be provided for the workmen. Kaffir workmen are said to suffer no inconvenience whatever from the contact of their skin with cyanide solutions.

Considering the extensive use of cyanide, the number of fatal accidents is remarkably small. Up to the present time only a few fatal cases have been recorded.

In a cyanide plant, poisoning may be apprehended from two principal sources, namely:—

(a.) From hydrocyanic acid liberated in vat-house.
(b.) From poisonous gases liberated during acid treatment of slimes.

In South Africa, Australia, and countries where the vats are not covered by roofs, danger from prussic acid vapours, liberated by the action of mineral acids or atmospheric carbonic acid, is practically unknown. Even where the vats are enclosed in a shed, the risk can be reduced to a minimum by the free circulation of fresh air.

The author has observed that the presence of HCy vapours is
always more noticeable in agitation than in percolation plants, the obvious reason being that agitation is generally adopted for the treatment of pyritic ores and concentrates, while the strength of the solutions used is nearly always high.

In cases of cyanide poisoning by inhaling the fumes of hydrocyanic acid, a German chemist recommends the use of hydrogen peroxide, $\text{H}_2\text{O}_2$, which forms with HCy the harmless compound oxamide, the reaction being represented by the following equation:

$$2\text{HCN} + \text{H}_2\text{O}_2 \rightarrow \text{CONH}_2.$$

This is said to be the most reliable and satisfactory remedy known at the present time.

In a case of HCy poisoning at the Crown Deep, the effect of the gas was immediate, one of the workmen falling as if dead. The same effect was observed at the N.Z. Crown mines, where a foreman fell into a cyanide vat without previous warning and died immediately.

**Danger during Acid Treatment of Slimes.** The slimes generally contain a small proportion of insoluble cyanide salts, which yield hydrocyanic acid when the sulphuric acid is poured on them. To guard against this danger, respirators should be used by the workmen who have to stoop over the dissolving tubs.

In cases of poisoning, subcutaneous injections of $\text{H}_2\text{O}_2$ are said by Mr. T. L. Carter of the Crown Deep mine to enable the patient to soon come to.

In ores containing arsenic, most of which are more or less soluble in cyanide, there is a danger of arsenic being precipitated with the gold in the zinc boxes. During the acid treatment of such slimes, the deadly poison arseniuretted hydrogen would be liberated by the action of the sulphuric acid on the zinc.

The symptoms observed in the case of the North Pole Company’s mill, where the superintendent and foreman both died, while many others were affected, were first nausea, then extreme langour, with pains in the legs, followed by discoloration of the skin in patches assuming the hue of tan; the whites of the eyes became yellow as in jaundice; finally, the passing of blood instead of urine to such an extent that the fluid coagulated in a few hours, the patients apparently dying from internal mortification.

The arsenic being inhaled, passes from the lungs through the whole system, and rapidly attacks the tissues of the body, precluding any relief by means of antidotes.

Where the acid treatment is used, the zinc for precipitation purposes should be free from arsenic; and in all cases the dissol-
tion of the zinc should be conducted in a special chamber or cupboard connected with a chimney having a good draught.

In cases of internal poisoning, vomiting should be induced at once by emetics or physical means.

Freshly precipitated carbonate of iron, obtained by mixing equal quantities of sodium carbonate and ferrous sulphate, is recommended for internal use.

It was lately reported in the press that Johann Antal, a Hungarian toxicologist, had found that a solution of cobalt nitrate was a perfect antidote for prussic acid poisoning. Recent investigations, however, have shown that cobalt salts exert a toxic action on animals when injected subcutaneously, finally leading to death; and for this reason nitrate of cobalt cannot be recommended for human subjects.

Cyanide Sores.—A percentage of the workmen engaged in the "clean-up" of the zinc extractors are affected with painful sores in those parts of their hands or arms which come in contact with cyanide solutions. Why some men should be exempt and some afflicted in this way is not very clear, but it is probably connected with constitutional causes.

Writing on this subject, Mr. A. Watt supplies the following instructive notes*:

"These painful affections may arise from two principal causes: first, from dipping the hands or arms into cyanide baths to recover articles which have dropped into them—a very common practice and much to be condemned; and second, from the accidental contact of the fingers or other parts of the hand, on which a recent cut or scratch has been inflicted, with cyanide solutions. In the former case, independent of the constitutional mischief which may arise from the absorption by the skin of the cyanide salts, the caustic liquid acts very freely upon the delicate tissue of the skin, but more especially upon the parts under the finger nails. We have known instances in which purulent matter has formed under the nails of both hands from this cause, necessitating the use of the lancet and poulticing. Again, when cyanide solutions come in contact with recent wounds—even very slight cuts or abrasions of the skin—a troublesome and exceedingly painful sore is sure to result, unless the part be at once soaked in warm water; indeed it is a very good plan, after rinsing the part in cold water, to give it a momentary dip in a weak acid pickle, then soak it for a few moments in warm water, and after wiping the part dry with a clean rag or towel, apply a drop of olive oil and cover up with a strip of thin sheet of gutta-percha."

* Watt, Electro-Deposition, p. 611.
Antidote for Cyanide Poisoning.—In order to minimize the danger attending cyanide poisoning, the necessary antidotes should be provided in every cyanide plant, and kept in a closed cabinet with a glass front, placed in some conspicuous and easily accessible part of the works, known to all the workmen.

The cabinet should have the words Antidotes for Cyanide Poisoning marked in clear letters near the top, and written or printed instructions how to apply the remedies pasted inside the cabinet.

The Committee of the Chemical, Mining, and Metallurgical Society of South Africa, appointed to investigate cyanide poisoning, agree in recommending as an antidote the following:—

1. Thirty c.c. of a 23 per cent. solution of ferrous sulphate.
2. Thirty-three c.c. of a 5 per cent. solution of caustic potash.
3. Two grammes of powdered oxide of magnesium (light).

They recommend, therefore, that in every cyanide-room there should be kept three boxes, containing—

1. A metal receptacle to hold about a pint, and a spoon.
2. A hermetically sealed phial, containing 30 c.c. of a 23 per cent. solution of ferrous sulphate.
3. A phial containing 30 c.c. of caustic potash.

They also suggest, further, that the ferrous sulphate be kept in a blue phial, and the caustic potash in a white phial; then, in the event of a case of cyanide poisoning occurring, all that the nearest employee would have to do would be to empty the contents of a blue phial, a white phial, and a packet into the metal receptacle, stir it with a spoon, and give it to the sufferer.

The box of remedies should never be removed from its place, but always kept complete and ready for an emergency. It is advisable to keep the remedies in duplicate in case of the accidental breaking of a sealed bottle.

In all cases of cyanide poisoning medical aid should be summoned without delay.
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