Heat and Acid Leach Treatments to Lower Phosphorus Levels in Goethitic Iron Ores

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Phosphorus associated with goethite in high-phosphorus (>0.10 mass% P) iron ores was lowered to below 0.075 mass% P with a heat treatment at 300 or 350°C for 1 h followed by a sulphuric acid (H\textsubscript{2}SO\textsubscript{4}) leach. This phosphorus removal was associated with a sample weight loss of 10–20 mass% due to dissolution of iron oxides. After heating at 900°C for 1 h, a sulphuric acid leach resulted in similar phosphorus removal but with dissolution of less than 3 mass% of the sample.

The weight losses in the leach are associated with phase changes of the phosphorus-containing goethite phase during heating. Heating at 300 or 350°C resulted in conversion of the goethite into an intermediate hematite phase (protohematite), while heating at 900°C gave a dense hematite phase. Compared with goethite in the ore, the more porous protohematite phase was more soluble in the sulphuric acid resulting in dissolution of iron with the phosphorus, while the dense hematite phase was much less soluble and little iron was dissolved in the leach.

Leaching at 25 mass% solids for 3 h at 60°C, at a pH of 0.5 or lower, gave significant lowering of phosphorus levels. Leaches were with 0.1–1M H\textsubscript{2}SO\textsubscript{4}; the concentration of acid required depended on the amount of phosphorus to be removed. Recycling of the acid leach liquor four times did not show evidence for precipitation of phosphorus and resulted in leach solutions with up to 1 g/L P and 134 g/L Fe.

KEY WORDS: phosphorus; goethite; heat treatment; sulphuric acid.

1. Introduction

Phosphorus in feedstock for blast furnaces affects the quality of the steel produced,\textsuperscript{1} and while methods for removing phosphorus from molten iron exist,\textsuperscript{2} it is beneficial to operations if phosphorus levels are low prior to smelting and melting.

Market specifications for phosphorus in iron ore exported from Australia are around 0.075 mass% P.\textsuperscript{3} The bulk of current production is a blend of low-phosphorus (<0.07 mass% P) martite-hematite ores and high-phosphorus martite-goethitic ores.\textsuperscript{4} As discussed previously,\textsuperscript{5} it has been estimated\textsuperscript{6} that there are 8.02 billion tonnes of high phosphorus ore (>0.10 mass% P) in Australian deposits. The development of a successful process to remove phosphorus from the high phosphorus ore would significantly extend the reserves of high grade low phosphorus Australian ores.\textsuperscript{3}

In Australian ores from the Pilbara, most of the phosphorus observed in the iron ore is associated with goethite.\textsuperscript{4,7,8} This phosphorus is considered to be mainly located in a texturally distinct goethite phase (termed ochreous goethite) that is commonly associated with high levels of other impurity elements such as silicon and aluminium.\textsuperscript{7,8} While high levels of phosphorus have been observed in some samples of vitreous goethite, where dehydration and recrystallisation of the goethite has occurred in natural deposits, phosphorus levels in this type of deposit are generally lower.\textsuperscript{9}

Physical separation methods\textsuperscript{5,10} and simple leaching processes\textsuperscript{5,10} do not remove sufficient phosphorus from the high phosphorus ores to meet market specifications of 0.075 mass% P. However, a heat treatment makes the phosphorus available for removal in a subsequent acid leaching step\textsuperscript{2,11} as shown by the heating of Australian iron ores at 1000–1300°C in air and leaching with sulphuric acid that has been reported to lower their phosphorus levels to below 0.06 mass% P.\textsuperscript{3,12,13} Also, after heating iron ores with various additives (the best being alkaline earth chlorides) at 500–1200°C (with 900°C the optimum), an acid leach has been shown to lower phosphorus levels from 0.42 to 0.02–0.08 mass% P depending on the additive.\textsuperscript{14} In our previous work,\textsuperscript{5,10} a heat treatment at 300°C followed by a caustic leach was shown to lower the phosphorus content of high phosphorus Australian iron ores to marketable levels.

In this paper, results for the removal of phosphorus from three iron ore samples with an acid leach after a heat treatment are reported. These results expand on our previous acid leaching results\textsuperscript{5} and enable the development of a comprehensive account of the influences of thermal treatment and acid leaching conditions on phosphorus removal from iron ore.
2. Experimental

2.1. Samples

Three West Australian high phosphorus iron ore fines (–6.3 mm) designated ore 1, ore 2 and ore 5 were used in this work. Ore 1 and ore 2 were examples of high phosphorus Brockman Iron Formation in the Pilbara and ore 5 was a high phosphorus ore from within the Yilgarn Iron Province. Representative samples for the test work were prepared by splitting a sub-sample from the ores as received, crushing and grinding the sub-sample to less than 2 mm to homogenise the sample, and then splitting small samples for individual tests from the –2 mm material.

The ores contained 0.146 mass% P (ore 1), 0.123 mass% P (ore 2) and 0.161 mass% P (ore 5) with iron contents of 62.0, 62.1 and 64.2 mass% Fe, respectively (Table 1). The XRF assays in Table 1 indicated that the compositions of ore 1 and ore 2 were similar and they contained higher levels of impurities (3.07 and 3.22 mass% SiO₂ and 2.27 and 2.18 mass% Al₂O₃, respectively) than ore 5 (0.40 mass% SiO₂ and 0.11 mass% Al₂O₃).

Quantitative XRD data for the samples indicate the iron minerals present in ore 1 and ore 2 were goethite (35–39 mass%) and hematite (55–60 mass%). A small amount of kaolinite (6 mass%), an aluminosilicate clay mineral, was also identified in ore 1 and ore 2. Ore 5 was composed mainly of goethite (66 mass%) and hematite (31 mass%) with a small amount of magnetite (3 mass%).

2.2. Methods

Small samples of the ores (1–2 g) were heated in a platinum boat in a tube furnace in air at the required temperature, usually for 1 h. After the required time at temperature the sample was allowed to cool in the furnace. When larger samples were used (70 g), the sub-samples were heated in a platinum dish in a muffle furnace in air (±10°C) for the required time. Generally, the sample was then removed from the furnace and allowed to cool in air before leaching or analysis. The heated samples were free flowing and were used for leaching without further processing.

The heated samples were leached in a glass reaction vessel fitted with a stirrer, a condenser and a thermocouple. The leachant and sample were heated to temperature with an external heating mantle and then leached for the required time. After leaching, the liquor was cooled and the leach product was filtered, washed with water, filtered and dried for analysis.

Solid samples for analysis were ground in a tungsten carbide ball-mill and elemental analyses were conducted on fused beads using a wavelength dispersive X-ray fluorescence (XRF) spectrometer using the program developed in CSIRO for iron ore analyses.

The concentration of phosphorus and iron in the leach solutions were determined using Inductively Coupled Plasma Optical Emission Spectroscopy (ICP OES).

3. Results

3.1. Effect of Heating Conditions

Heating the ores prior to a caustic leach has been shown to increase the amount of phosphorus removed. Similarly, a heat treatment prior to acid leaching increases the amount of phosphorus removed and Fig. 1 indicates that after the same heat treatment of ore 1, the products from leaches with 1M NaOH and 0.1M H₂SO₄ contained similar phosphorus levels as did those from leaches with 5M NaOH and 0.5M H₂SO₄. It can be seen that phosphorus removal increased significantly with heating temperatures of 275 to 300°C. Between 350–800°C there was little variation in the level of phosphorus removal, and greater amounts were removed when the ore was heated at 800°C and above.

The increase in phosphorus removal after heating at about 300°C and heating to above 800°C is considered to be associated with phase changes in the phosphorus-containing goethite fraction of the ores (see the Discussion).

Table 1. Effects of leaching conditions on impurity removal from the as received ore samples.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Leaching conditions</th>
<th>Pulp density (mass%)</th>
<th>Weight loss (mass%)</th>
<th>Leached product assays (mass%)</th>
<th>Phosphorus removed (mass%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Lixiviant</td>
<td>Temp. (°C)</td>
<td>Time (h)</td>
<td>Fe</td>
<td>P</td>
</tr>
<tr>
<td>Ore 1</td>
<td>Water</td>
<td>B Pt</td>
<td>3.0</td>
<td>0.5</td>
<td>0.7</td>
</tr>
<tr>
<td></td>
<td>0.5M H₂SO₄</td>
<td>60</td>
<td>3.0</td>
<td>0.5</td>
<td>3.2</td>
</tr>
<tr>
<td></td>
<td>1M H₂SO₄</td>
<td>60</td>
<td>3.0</td>
<td>25</td>
<td>1.4</td>
</tr>
<tr>
<td>Ore 2</td>
<td>Water</td>
<td>B Pt</td>
<td>3.0</td>
<td>0.5</td>
<td>1.3</td>
</tr>
<tr>
<td></td>
<td>0.1M H₂SO₄</td>
<td>B Pt</td>
<td>3.0</td>
<td>0.5</td>
<td>4.6</td>
</tr>
<tr>
<td></td>
<td>1M H₂SO₄</td>
<td>60</td>
<td>3.0</td>
<td>25</td>
<td>1.2</td>
</tr>
<tr>
<td>Ore 5</td>
<td>Water</td>
<td>B Pt</td>
<td>3.0</td>
<td>0.5</td>
<td>0.6</td>
</tr>
<tr>
<td></td>
<td>0.5M H₂SO₄</td>
<td>60</td>
<td>3.0</td>
<td>0.5</td>
<td>2.3</td>
</tr>
<tr>
<td></td>
<td>1M H₂SO₄</td>
<td>60</td>
<td>3.0</td>
<td>25</td>
<td>1.2</td>
</tr>
</tbody>
</table>

† Weight loss in the leach based on the solids added to the leach and the residue weight.

‡ Based on the head assay, the weight before heating and the residue assay and weight.
work\textsuperscript{5) indicated that the phase changes were complete within 30 min for a heating temperature of 300 °C, significantly less than the 1 h heating time generally used in this work.

### 3.2. Effect of Leaching Conditions

The effect of the leaching conditions to lower phosphorus levels in the ores was determined using sub-samples of the ores and samples heated at 300 or 350 °C and at 900 °C prior to leaching.

#### 3.2.1. Acid and Water Leaching of the Ores

Leaching ore 1 with boiling water removed little phosphorus and a leach with 1M \( \text{H}_2\text{SO}_4 \) at 25 mass% solids removed almost 10 mass% of the phosphorus (Table 1). With ore 2, 10 mass% of the phosphorus was removed in both the water and 1M \( \text{H}_2\text{SO}_4 \) leaches. There was little loss of weight of the samples in the water and acid leaches and the resulting leach products for ore 1 and ore 2 contained 0.136 mass% P and 0.112 mass% P, respectively. Acid leaching at a low solids pulp density of 0.5 mass% solids removed 15 mass% of the phosphorus from ore 1 and 19 mass% from ore 2 (Table 1).

Water and acid leaching of ore 5 removed little phosphorus from the sample (Table 1).

Leaching with water or acid was ineffective in mobilising silica or alumina from ore 1, ore 2 or ore 5.

#### 3.2.2. Heating at 300 or 350 °C Followed by an Acid Leach

Acid leaching after a heat treatment gave better phosphorus removal with the amount of phosphorus removed increasing with increasing acid concentration. For ore 1, heated at 300°C, the results in Table 2 show that a leach with 0.1M \( \text{H}_2\text{SO}_4 \) gave a product with 0.108 mass% P, while a leach with 0.5M \( \text{H}_2\text{SO}_4 \) gave 0.061 mass% P. Similarly, increasing acid concentration increased phosphorus removal from ore 2 heated at 300 °C (Table 3) and ore 5 heated at 350 °C (Table 2). A leach of ore 5, the ore with the highest initial phosphorus content, with 0.5M \( \text{H}_2\text{SO}_4 \) gave a product with 0.118 mass% P while a leach of ore 1 under the same conditions with only 0.1M \( \text{H}_2\text{SO}_4 \) gave a product with 0.108 mass% P.

Increasing the solids pulp density in the leach required higher acid concentrations to achieve a similar levels of phosphorus removal. For heated ore 2, a leach with 0.1M \( \text{H}_2\text{SO}_4 \) at 0.5 mass% P while a leach with 0.5M \( \text{H}_2\text{SO}_4 \) gave 0.061 mass% P. Similarly, increasing acid concentration increased phosphorus removal from ore 2 heated at 300 °C (Table 3) and ore 5 heated at 350 °C (Table 2). A leach of ore 5, the ore with the highest initial phosphorus content, with 0.5M \( \text{H}_2\text{SO}_4 \) gave a product with 0.118 mass% P while a leach of ore 1 under the same conditions with only 0.1M \( \text{H}_2\text{SO}_4 \) gave a product with 0.108 mass% P.

### Table 2. Effects on impurity removal from the ore samples of a heat treatment at 300 or 350°C for 1 h and a leach with sulphuric acid for 3 h at 0.5 mass% solids.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Heating temp. (°C)</th>
<th>Leaching conditions</th>
<th>Weight loss\textsuperscript{2} (mass%)</th>
<th>Leached product assays (mass%)</th>
<th>Phosphorus removed\textsuperscript{1} (mass%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Lixiviant</td>
<td>Temp. (°C)</td>
<td>Fe</td>
<td>P</td>
</tr>
<tr>
<td>Ore 1</td>
<td>300</td>
<td>0.1M ( \text{H}_2\text{SO}_4 )</td>
<td>60</td>
<td>6.3</td>
<td>63.4</td>
</tr>
<tr>
<td>Ore 5</td>
<td>350</td>
<td>0.25M ( \text{H}_2\text{SO}_4 )</td>
<td>60</td>
<td>4.85</td>
<td>66.3</td>
</tr>
</tbody>
</table>

\textsuperscript{2} Weight loss in the leach based on the solids added to the leach and the residue weight.

\textsuperscript{1} Based on the head assay, the weight before heating and the residue assay and weight.

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higher levels of phosphorus removal and higher sample weight losses from ore heated at 300°C. A leaching time of 3 h was used for most tests.

Figure 2, a graph of phosphorus removed during leaching against the sample weight loss in the leach, shows that the amount of phosphorus removed from samples heated at 300 or 350°C increased with increasing weight loss. Extrapolation of the lines in Fig. 2 suggests that total phosphorus removal would be associated with a weight loss of around 40 mass%. For ores 1 and 2 this is close to the 35–39 mass% goethite in the ores, suggesting that the phosphorus in ores of this type is mainly in the goethite which is soluble in acid after heating. Ore 5, from a different province than ores 1 and 2, contains 66 mass% goethite and the results lead us to suspect that this sample contains different forms of goethite with the more soluble forms containing higher levels of phosphorus.

Generating leach products with 0.075 mass% P from ores heated at 300 or 350°C resulted in dissolution of iron oxides in the sample and iron levels of around 25 g/L Fe in the leach liquors (ore 2) with an associated sample weight loss of 10–20 mass%. In practice, to limit iron losses when leaching ores heated at 300 or 350°C, the liquors may need to be processed to recover iron from solution.

Ore 1 and ore 2 had appreciably higher silica and alumina levels than ore 5 and the assay data in Tables 2 and 3 for the ores heated at 300°C for 1 h indicate that, while little silica was removed in the acid leach, 30 mass% of alumina could be removed from ore 1 and 10 mass% from ore 2 with a 1M H2SO4 leach of ores heated to 300 or 350°C prior to leaching.

In a commercial operation, acid usually would be added at several stages during the leach rather than as a single addition at the start as was done for the leaches presented above. When acid was added during the leach to control the pH at a specified value, similar trends to those discussed above were observed as shown in Table 4. With a pH value of 0.5 or lower, a substantial reduction in phosphorus levels was achieved.

3.2.3. Heating at 900°C Followed by an Acid Leach

Heating at a higher temperature of 900°C resulted in a dense hematite phase being formed from the goethite.16) The results in Table 5 for ore 5 heated at 900°C show increasing phosphorus removal with increasing acid concen-
tation in the leach and suggest that a leaching temperature of 60°C is sufficient to reduce phosphorus levels in the leach product, as was observed with the lower heating temperatures.

Results for heating ore 1 and ore 5 at 900°C and leaching with sulphuric acid at 0.5 mass% solids (Table 5) and 25 mass% solids (Table 6) show significantly higher levels of phosphorus removal than when the ores were heated at 300 or 350°C. For example, with a heating temperature of 900°C and a leach at 0.5 mass% solids with 0.5M H₂SO₄, phosphorus levels in ore 5 were lowered to 0.076 mass% P compared with 0.118 mass% P with the lower heating temperature of 350°C (Table 3).

About 50 mass% of the alumina was removed by acid leaching of ore 1 or ore 2 heated at 900°C, but there was little silica removed in the acid leach (Tables 5 and 6). The significantly lower weight loss during leaching of ores heated at 900°C reflects the lower solubility of the dense hematite phase compared with the ores heated to 300 or 350°C (Fig. 2). For example, a leach residue with 0.075 mass% P was achieved for the ores with a weight loss of less than 2.5 mass% (Tables 5 and 6), while with the lower heating temperature of 300°C the weight loss was 10–20 mass%, and less phosphorus was removed (Tables 2 and 3).

### 3.3. Acid Recycle Tests

In a commercial operation, it can be expected that the

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**Table 4.** Effects on impurity removal from heated ore samples of a sulphuric acid leach at a constant pH.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Heating temp. (°C)</th>
<th>Target pH</th>
<th>Weight loss¹ (mass%)</th>
<th>Leached product assays (mass%)</th>
<th>Acid added² (kg/t)</th>
<th>Phosphorus removed³ (wt %)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Fe</td>
<td>P</td>
<td>SiO₂</td>
</tr>
<tr>
<td>Ore 1</td>
<td>300</td>
<td>1.0</td>
<td>0.9</td>
<td>62.0</td>
<td>0.146</td>
<td>3.07</td>
</tr>
<tr>
<td></td>
<td>300</td>
<td>0.5</td>
<td>14.0</td>
<td>63.6</td>
<td>0.140</td>
<td>3.06</td>
</tr>
</tbody>
</table>

* Samples were heated at 300°C for 1 h (ore 1 and ore 2) or 350°C for 1 h (ore 5) and leached at 80°C for 3 h at 25 mass% solids with the pH maintained at the selected value by addition of sulphuric acid. Acid addition was required throughout the leach.

¹ Weight loss in the leach based on the solids added to the leach and the residue weight.

² Based on the weight before heating, and the volume and concentration of sulphuric acid added during the leach.

³ Based on the head assay, the weight before heating and the residue assay and weight.

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**Table 5.** Effects on impurity removal from the ore samples of a heat treatment at 900°C for 1 h and a leach with sulphuric acid for 3 h at 0.5 mass% solids.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Leaching conditions*</th>
<th>Weight loss‡ (mass%)</th>
<th>Leached product assays (mass%)</th>
<th>Phosphorus removed³ (mass%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Lixiviant</td>
<td>Temp. (°C)</td>
<td>Fe</td>
<td>P</td>
</tr>
<tr>
<td>Ore 1</td>
<td>Water</td>
<td>60</td>
<td>1.0</td>
<td>65.1</td>
</tr>
<tr>
<td></td>
<td>0.1M H₂SO₄</td>
<td>60</td>
<td>2.1</td>
<td>65.6</td>
</tr>
<tr>
<td></td>
<td>0.5M H₂SO₄</td>
<td>60</td>
<td>5.4</td>
<td>66.2</td>
</tr>
<tr>
<td>Ore 5</td>
<td>Water</td>
<td>60</td>
<td>0.5</td>
<td>68.8</td>
</tr>
<tr>
<td></td>
<td>0.1M H₂SO₄</td>
<td>60</td>
<td>1.6</td>
<td>68.8</td>
</tr>
<tr>
<td></td>
<td>0.25M H₂SO₄</td>
<td>60</td>
<td>1.3</td>
<td>68.9</td>
</tr>
<tr>
<td></td>
<td>0.5M H₂SO₄</td>
<td>60</td>
<td>2.3</td>
<td>69.3</td>
</tr>
<tr>
<td></td>
<td>1M H₂SO₄</td>
<td>60</td>
<td>2.5</td>
<td>68.9</td>
</tr>
<tr>
<td>Ore 5</td>
<td>1M H₂SO₄ R T</td>
<td>1.4</td>
<td>68.5</td>
<td>0.093</td>
</tr>
<tr>
<td></td>
<td>1M H₂SO₄</td>
<td>60</td>
<td>2.5</td>
<td>68.9</td>
</tr>
<tr>
<td></td>
<td>1M H₂SO₄ B Pt</td>
<td>26.1</td>
<td>68.5</td>
<td>0.050</td>
</tr>
</tbody>
</table>

* BPt = Boiling Point and RT = Room Temperature (around 21°C).

¹ Weight loss in the leach based on the solids added to the leach and the residue weight.

² Based on the head assay, the weight before heating and the residue assay and weight.
leach liquors would be recycled several times to leach fresh heated ore to minimise water requirements for the plant and reduce acid consumption. In order to determine whether phosphorus salts might precipitate from solution in subsequent leaches, the heated ores were leached with sulphuric acid and then the leach liquor was used for another four (recycle) leaches. Results for the recycle leaches on ore 2 heated at 350°C, and ore 5 heated at 900°C, are presented in Table 7.

Ore 2, was leached at 60°C with recycled liquor to which concentrated sulphuric acid was first added to return the pH of the liquor to a value of pH 0.1. The results in Table 7 show that four leaches had an average product phosphorus level of 0.078 mass% P with an average weight loss of 8.1 mass% and the phosphorus levels in solution increased to 1 g/L P without any indication of phosphorus precipitation. An iron level of 134 g/L Fe was measured in the final solution sample, corresponding to an average of 26.8 g/L Fe dissolved in each leach (Table 7). The average acid addition for each of the recycle leaches was 120 kg H₂SO₄/t ore.

After heating at 900°C, recycle leaches on ore 5 at room temperature with the pH returned to 0.1 prior to each leach gave an average phosphorus content of 0.099 mass% P in the leach products, with an average weight loss of only 0.65 mass% and again the solution assays did not indicate any precipitation of phosphorus during the leaches up to a concentration of 1 g/L P in solution. The average solution assay for iron for each leach was low (0.39 g/L Fe), reflecting the small weight losses in the leach (Table 7). The average acid addition for each of the recycle leaches was 90 kg H₂SO₄/t ore.

It is interesting to note that a lower acid consumption was measured for the ore heated at 900°C compared with that for the ore heated to 300°C. It is anticipated that the acid consumption will be significantly lower if the recycle tests were conducted at a pH of 0.5 that has been shown to give acceptable phosphorus removal (Table 4).

4. Discussion

The results presented here indicate that phosphorus levels can be lowered in goethitic iron ores by leaching with sulphuric acid after a heat treatment. The final phosphorus levels achieved for each ore were related to the temperature at which the ore was heated, the concentration of acid in the leach, the temperature of the leach, the amount of phosphorus required to be removed from the ore, and the solubility of other phases in the ore. Leaching at the higher pulp density required a higher acid concentration to meet equivalent residual phosphorus levels.

The heating treatment results in dehydration of the
goethite and its conversion, through several intermediate phases, to a dense hematite phase. A schematic sequence of the conversion has been presented\textsuperscript{16} with decomposition of synthetic goethite commencing at 150°C and transforming first into ‘protohematite’ (in a temperature range of 250–400°C), then to ‘hydrohematite’ (430–700°C) and ultimately to hematite (above 800°C). In the samples used in this study, XRD patterns indicated that the conversion of goethite to protohematite had commenced at 250°C and was substantially complete after 30 min at 300°C.\textsuperscript{5}

For nickel laterite ores containing nickel associated with the goethite component of the ore, it has been found\textsuperscript{17} that dissolution of iron and nickel was significantly increased in acid leaches after heating the ores at temperatures of 340–400°C. It was proposed\textsuperscript{18} that the increase in metal dissolution was partly due to an increase in surface area from the development of micropores as a result of dehydroxylation of the goethite during the heating, and partly to an overall increase in structural disorder.

Removal of phosphorus from goethitic iron ores after heating at 300–350°C is anticipated to follow a similar mechanism. The increase in both the porosity and the structural disorder of the iron phase on heating results in increased reactivity of the phosphorus-containing iron oxide phase. The high weight loss during leaching (Fig. 2) indicate that this phase is dissolved in the acid along with the phosphorus. On heating at 900°C the goethite is converted to a dense hematite phase that has a low solubility in acid, and this results in dissolution of phosphorus with little weight loss.

Ore 1 and ore 2 contained appreciable levels of silicon and aluminium. While silicon was not removed in the acid leaches, up to 50 mass% of the aluminium was removed. This suggests that some of the aluminium may be associated with phosphorus in the goethite, as proposed by MacRae and co-workers.\textsuperscript{18}

Obtaining leach products with 0.075 mass% P from ores heated at 300 or 350°C resulted in dissolution of iron oxides in the sample with iron levels of around 25 g/L Fe in the leach liquors (ore 2), an associated sample weight loss of 10–20 mass%. Leaching ore heated at 900°C under similar acid conditions resulted in better phosphorus removal with little dissolution of iron oxides and a lower acid consumption. The economics of a process with a lower heating temperature but requiring higher acid consumption and treatment of the increased amounts of iron in solution, compared with a higher heating temperature and little iron dissolution during the leach will be a significant consideration in determining the best commercial operating conditions.

5. Summary

The results of the effects of thermal transformations of goethite in iron ores on the removal of phosphorus from Australian iron ores with a sulphuric acid leach have been presented in this paper. The study demonstrated that a low temperature heat-treatment at 300 or 350°C and a sulphuric acid leach process reduced the phosphorus levels in the ore to below 0.075 mass% P, but this was associated with appreciable levels of iron in the leach liquor and high acid consumption. Heating to about 350°C is considered to result in conversion of the phosphorus-containing goethite to a hematite intermediate phase, protohematite, that was readily soluble in the sulphuric acid along with the phosphorus.

Increasing the heating temperature to 900°C improved phosphorus removal in the leach. Heating at 900°C transformed the goethite to a dense hematite phase that has a low solubility in the acid and enabled phosphorus removal with little dissolution of iron and lower acid consumption.

The best operating conditions are likely to depend on balancing the benefits of a lower heating temperature with dissolution of appreciable amounts of iron oxides compared with a higher heating temperature but with low levels of iron in the leach liquors levels.

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