FLOWSHEET DEVELOPMENT FOR THE EXTRACTION OF RARE EARTHS USING
MINTEK’S PYEARTH™ PROCESS

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ABSTRACT

The PyEarth™ process developed at Mintek has been demonstrated successfully at pilot scale. Nine tons of ore containing 2.3 mass % of total rare earth oxides (TREO) from a dormant minerallogically complex iron-rich ore deposit, was subjected to a concentrating smelting step in a 200 kW DC open-arc pilot furnace and the rare earths from the resulting slag were extracted via a hydrometallurgical route. This ore deposit is practically not amenable to extraction by conventional methods. The testwork results proved that the PyEarth™ process was able to optimize the extraction of rare earths while minimizing waste through the removal of iron as a saleable iron alloy. Reduction of iron to the alloy in the smelting step was found flexible, efficient, and controllable, while rare earths were left virtually unreduced in the slag phase. The slag chemistry is selected to optimize both the smelting and leaching performances as well as the alloy composition. Leaching of the resulting slag was found possible and most efficient in hydrochloric acid amongst other lixivants tested.

KEYWORDS

Direct smelting, Extraction, Hydrochloric leaching, Iron rich rare earth bearing ore, PyEarth, Rare earth
INTRODUCTION

Rare earth elements have been used extensively over the past decade in the manufacture of clean technology items such as wind turbines, hybrid and electric car batteries, magnetic levitation (‘maglev’) trains and many more. High-strength magnets are among the major applications of rare earths. These elements also find applications in catalysts, and in military equipment such as armoured vehicle navigation systems, submarine engines, and missile guidance systems, and even for refining jet fuel (Gupta et al., 2004).

Rare earths are found in natural deposits with varying abundance, mineralogy, and textures. The proven global rare earth oxide (REO) reserves are estimated at about 110 million tons (Wang et al., 2015), with about 43% of the global reserves located in China. Namibia has the second-largest reserves worldwide, with 20 million tons. South Africa has reserves of 1 million tons. Other rare-earth resources are located in Russia (17%), the United States of America (12%), Australia, India, Brazil, and Greenland (Gupta & Krishnamurthy, 2004).

About 130 000 t/a of rare-earth oxide (REO) is produced globally, with about 90% of this production coming from China. Notwithstanding the available large reserves worldwide, the bulk of global production of rare earths comes from the Bayan Obo in China and the Mount Weld area in Australia (Long et al., 2010; Li et al., 2014; Haque et al., 2014). These deposits are mainly ferruginous (iron-rich) rare earth-bearing ores. The Bayan Obo (China) ore contains 6% REO and 35% Fe; it is dominantly composed of discrete minerals of bastnaesite and monazite, with a minor proportion intimately intergrown within Fe oxide minerals (dominantly hematite and magnetite). The principal REE minerals at Mount Weld (Australia) include monazite, and, in parts, xenotime. The monazite would appear to be altered, secondary in nature, closely associated with Fe oxides; its average ore grade is 15.4% REO (Haque et al., 2014).

More than 50% of current world production of rare earths is currently consumed in China. Consumption of REO is projected to increase at a rate of 6–10% per annum and could reach a volume of 200 000 t/a in the very near future (Long et al., 2010; Li et al., 2014; Wang et al., 2015).

The ore from the southern African regions is dominated by Fe and Mn oxides and oxyhydroxides such as goethite and manganomelane minerals, the latter sometimes containing Ba. These minerals account for more than 65% of the ore, and contribute to a high Fe content. Rare earth element minerals are monazite, chemically weathered or altered monazite, and REE-bearing Al phosphates of the crandallite group (florensite, gorceixite, crandallite). Additionally, rhabdophane, Ce-pyrochlore, and Ca monazite (cheralite) are sometimes observed. As such, altered monazite is often finer-grained than monazite, and displays poorer liberation. Nevertheless, most of the REE minerals occur in size classes < 20–30 µm, and very fine intergrowths with the Fe-Mn oxyhydroxide minerals preclude liberation, even at very fine sizes (Chetty et al., 2017).

BRIEF OVERVIEW OF RARE EARTH EXTRACTION

Conventional processes

Highly efficient separation technologies have been key to the exploitation of the rare earths. The extraction of rare earths should be sustainable, environmentally friendly, and economically viable. Conventional flowsheets used include milling, and production by physical methods of a concentrate, from which rare earths are recovered by hydrometallurgical means.

The Bayan Obo ore is milled to 90–95% passing 74 µm, and then processed through low- and high-intensity magnetic separation, and flotation of iron minerals, leaving cleaner tailings concentrating REE minerals. However, the REE are concentrated using flotation of milled ores for the Mount Weld process (Li & Yang, 2014).

Flowsheets are particularly designed to meet mineralogical challenges. The cost of the conventional extraction process increases with mineralogical complexity, as more process units would be
required to extract the rare earths and also to minimise pollution of the environment and groundwater with the radioactive wastes generated.

**PyEarth process**

Upgrading of ore from a number of deposits, particularly those from the southern African regions, by physical methods would not be economically viable because of the complexity of the mineralogy. These ores would require fine milling to liberate the rare earth oxide species, and concentration of very fine milled materials is rather challenging. Application of fine milling below 35 μm size is not usual in commercial operations because this is energy-intensive and costly. Research activities are geared towards development of more efficient flotation reagents that will enable the upgrading of the ore into a rich rare earth concentrate. For all these reasons, upgrading the low-grade (typically 2.5% REO) ferruginised southern African ores using the conventional process was found to be very challenging and inefficient. The high Fe content (> 30% Fe), and textural and mineral complexity, precluded the typical processing route of comminution, physical separation, and leaching.

A novel process, called PyEarth™, was invented by Mintek, and employs direct smelting of the ores to selectively remove iron as a saleable Fe-rich alloy whilst concentrating the REO in the slag phase. The direct smelting approach considered the general principles of the pyrometallurgical process: the thermodynamics and furnace operation. The basic premise is to selectively reduce iron oxide (Fe₂O₃, Fe₃O₄, FeO in FeO(OH)) to iron, thereby concentrating REOs in the slag, as most rare earth oxides are not readily reduced to metallic form at lower smelting temperatures of between 1450 and 1550°C. In this temperature range, the iron oxides are easily reduced by carbon, and the manganese oxides are moderately easily reduced, whereas, for example, CeO₂, La₂O₃, and Ce₂O₃ are (in increasing order) much harder to reduce.

As ores in the southern parts of Africa are very rich in iron oxides, reductive smelting of such ores could potentially result in significant upgrading of the REO content of the resultant slag.

A number of laboratory-scale smelting tests were conducted in alumina, magnesia, and graphite crucibles. The crucible tests provided information that was useful for the pilot tests that are outlined here (Bisaka et al., 2016).

- The rare earth-bearing slag is of amphoteric nature when considering the wear of alumina and magnesia crucibles. This erosion was the cause of crucible failures and contamination of the final slag product, thus affecting the REO grade of the slag.
- Clean separation between the slag and metal was only achieved at high temperatures, above 1700°C. The crucible products displayed poor separation at lower temperatures.
- Clean slag with the highest REE grade, up to 13% REE, was achieved in the graphite crucible.
- The recovery of Fe to the alloy was > 98%.
- Full recovery of rare earth to the slag was achieved.
- A graphite lining was recommended for the pilot-scale furnace.

The slag produced was submitted to various leaching tests. The leaching tests showed the following (Bisaka et al., 2016):

- Alumina contamination affected the leaching efficiency.
- HCl leaching of the clean slag produced in the graphite crucible was the most efficient, with leaching extraction as high as 98%.

The objective of the current work was to demonstrate the smelting of the iron-rich rare earth ore in Mintek’s 200 kW DC arc furnace, and the leaching of the slag produced from the pilot smelting tests.
Operating an iron-rich rare earth smelting process in a DC arc furnace requires a knowledge of the products’ properties, particularly the slag properties such as the slag liquidus temperature, viscosity, and density. These parameters contribute significantly to the metal-slag separation, flowability, and the thermal energy distribution in the furnace. As data for rare earth-bearing slags are scarce in the open literature, theoretical simulations of the slag properties were carried out to estimate the performance of the slag in a DC arc furnace.

**SMELTING TESTWORK**

**Raw materials**

Nine tons of iron-rich rare earth-bearing ore from a southern African deposit was subjected to the smelting tests in Mintek’s 200 kW DC arc furnace. The ore, with 100% passing -6 mm, was processed as received without any further size reduction or physical preparation. A sub-sample of the ore was submitted for chemical and mineralogical characterisation. The bulk chemical composition was determined by a Varian Vista-PRO CCD simultaneous inductively coupled plasma optical emission spectrometer (ICP-OES). The morphological and phase chemical compositions were determined by Zeiss MA15 scanning electron microscopy (SEM) equipped with a Bruker energy-dispersive spectrometer (EDS) and Bruker D8 advanced X-ray diffractometer (XRD), respectively. The chemical composition of the ore is given in Table 1. The proximate analysis of the anthracite is given in Table 2. High-purity CaO was used as a fluxing agent to improve the smelting process, particularly to decrease the slag liquidus temperature and viscosity—constraints that are exacerbated at small scale. Few rare earth elements with concentrations higher than 100ppm are included in Table 1. Concentrations of Th and U in the ore were 221 and 72 ppm, respectively.

| Table 1. Summary of the bulk chemical composition of the iron-rich rare-earth ore (mass %) |
| MgO | Al₂O₃ | SiO₂ | CaO | TiO₂ | MnO | FeO(OH) | P₂O₅ | La | Ce | Nd | TREE |
| % | % | % | % | % | % | % | ppm | ppm | ppm | ppm | % |
| 1.13 | 6.48 | 6.08 | 2.06 | 3.87 | 9.09 | 52.45 | 1.77 | 6,060 | 10,200 | 3,900 | 2.34 |

| Table 2. Summary of the bulk chemical composition of the anthracite (mass %) |
| Ash | Volatiles | Fixed carbon | Total sulphur |
| % | % | % | % |
| 12.2 | 4.3 | 81.2 | 0.71 |

**Pilot smelting setup**

The 200 kW DC arc furnace facility used for this smelting work is schematically presented in Figure 1. The furnace is made of a shell having an outer diameter of 1 m. The water-spray-cooled steel shell is lined with a single layer of graphite bricks, three rows high, and a hearth lined with rammable carbon-based materials. The refractory lining resulted in a furnace crucible internal diameter (ID) of approximately 0.7 m. The furnace was equipped with an alumina-lined conical roof, and a shell bolted onto a domed base. A single tap-hole was used to tap a stream of both molten slag and metal. A centrally-located graphite electrode of 40 mm diameter operated as a cathode, and the anode was constituted of steel pins buried in the hearth. The feed system comprised individual hoppers used to feed ore, anthracite, and limestone. The furnace is equipped with an off-gas system for the cleaning of process gases prior to their release to the atmosphere.
Experimental description

Various conditions were investigated throughout the 92 batches that made up the smelting testwork. Anthracite additions were gradually increased from 12.5% to 25%, and two conditions of limestone addition at 1–5% were tested. As in the laboratory-scale tests conducted previously, lime was added to improve the reduction of MnO, and to lower the viscosity of the slag. The discussions in this paper will be focused around the furnace behaviour and the analyses of the products.

SMELTING RESULTS

Chemical analyses and recoveries

The pilot process consisted of current-control furnace operation with constant power and constant feed rate. After a batch of about 120 kg of rare earth ore was fed into the furnace, the tap-hole was opened to release the molten materials. The latter were collected into a ladle, and left to cool prior to separation and sampling. The samples were sent for chemical analysis.

The molten material flowed freely when the tap-hole was opened. Clean metal and slag layers were separated after solidification in a ladle. The good metal-slag separation confirmed that acceptable slag properties were achieved. As can be seen in Figure 2, the FeO content in the slag decreased with increasing anthracite addition, while the REE concentration in the slag increased. Figure 2 also shows that when 5% limestone was added with 21.5% anthracite (in ratio to the ore), the concentration of REE increased to a greater extent. REE were typically upgraded to more than 7 – 8%, and the highest REE was slightly above 10% in the present testwork; this is 4 times the REE concentration in the ore.

The evolution of the alloy analyses through the testwork is presented in Figure 3. The concentration of Fe in the alloy decreased while Mn increased. At low reduction of Mn, Fe was in the range of 96 to 98%. The highest Mn in the alloy was closer to 5%. Lower reduction of Mn was observed in this testwork as compared to the crucible tests where the Mn in the alloy was more than 10% (Bisaka et al., 2016). The low levels of MnO reduction in the pilot tests were attributed to the low operating temperatures achieved in the pilot tests. The operating temperature of the furnace was 1600°C towards the last batches.
By comparison, the laboratory-scale temperatures at which high reduction of MnO was achieved ranged between 1600 and 1800°C (Bisaka et al., 2016).

![Figure 2](image1.png)

**Figure 2.** Evolution of slag FeO, REE (mass %) throughout the testwork

![Figure 3](image2.png)

**Figure 3.** Evolution of the alloy composition (mass %) throughout the testwork

**Estimation of slag properties pertinent to operation**

The rare earth oxides in the material have very high melting points, in excess of 1900°C. These oxides also readily form phases with a perovskite structure with, for example, Al₂O₃, hence it is not expected that these oxides ever melt at process temperatures of 1500–1600°C. Separation of REOs is carried out by way of removal of all other gangue around phases associated with REOs as much as possible. In this case, FeOₓ and MnOₓ are both reduced to metal and leave the REO structure behind.
The slag formed in this process is relatively complex to study, based on the fact that it contains many components for which limited information is available for the estimation of slag properties (in this case liquidus and viscosity). Major components expected in the slag were 5–15% FeO, 20% MnO, 15–20% Al₂O₃, 20–30% SiO₂, 1–5% MgO, 5–20% CaO, 1–10% BaO, and 9–12% TiO₂. Other components include 2–3% SrO, as well as the rare earth oxides. The liquidus temperature of the slag was initially estimated using FactSage 6.4, using all the components in the slag excluding SrO, BaO, and the REOs. Data for BaO-solutions became available in FactSage 7.1, released in January 2017, and all of the calculations were repeated using the new database.

The testwork was grouped into four broad categories:

- No flux, with 12 – 19.5% anthracite, which yielded a slag with %FeO + %MnO ~ 35%
- No flux, with 21.5% anthracite, which yielded a slag with %FeO + %MnO ~ 30%
- Additions of 21.5% anthracite and 5% limestone
- Additions of 25% anthracite and between 1 and 5% limestone.

For the bulk (>85%) of the slag consisting of FeO, MnO, Al₂O₃, SiO₂, MgO, TiO₂, and BaO, the liquidus temperature was estimated for the different conditions tested. It was found that the liquidus temperature of the bulk slag (excluding SrO and rare earth oxides), under reducing conditions, was between 1400 and 1450°C for slags where the %FeO + %MnO was between 30 and 35%, while for slags with very little (< 5% FeO, and/or < 5% MnO), the liquidus temperature was slightly higher at 1520–1530°C.

The viscosity of the bulk slag was estimated using the Viscosity module in FactSage™ 7.1. Components in the calculation were FeO, MnO, Al₂O₃, SiO₂, MgO, TiO₂, and BaO. It was assumed, as a first approximation, that BaO has a similar effect on the viscosity of the slag as CaO. Reduction of TiO₂ to Ti₂O₃ was ignored. Experience at Mintek has shown that, in general, slags with a viscosity lower than 4.5 poise are readily tappable from the DC furnace. Slags with high amounts of FeO and MnO had viscosity values of 0.6–1.7 poise for temperatures from 1500 to 1700°C. Slags with low FeO values had significantly higher viscosities (4.7 to 20.9 poise), which is part of the reason why CaO was used as flux to compensate for the loss of FeO in the slag.

It was also predicted that the addition of CaO to the slags low in FeO would lead to the precipitation of CaTiO₃ (a perovskite type structure). SEM-EDS analysis of the slag samples (alluded to in the leaching section of this paper) confirmed that REOs were found primarily on their own as needles in the slags with no flux, and as partitioned between Ca-Ti-perovskite and REO needles in slags where CaO was used as flux. This is not unexpected, as Ca is known to be able to easily occupy the “A” site in the perovskite structure (ABO₃, where A = Ca, Ce, Sr, etc., B = Ti, Nb, etc.) (Hendrick et al., 1997; Feng et al., 2016).

**LEACHING EXPERIMENTAL WORK**

Leaching raw material

Two slag samples generated in the smelting testwork were subjected to leaching tests. Sample A was a blended sample produced in batches 44 through 49, under the condition with 21.5% anthracite and no flux additions to the smelting recipe. Sample B was produced in batch 87, where 25% anthracite and 5% limestone were added to the smelting recipe. The chemical analyses of these slag samples are given in Table 3. Sample B contained more REE as a result of improved MnO reduction, and it also had a higher CaO concentration. Sample A and Sample B were classified as high Mn and low Mn slag, respectively. The chemical analysis of the Th and U were at 738 ppm and 240 ppm in Sample A and 718 ppm and 248 ppm in Sample B, respectively. At these levels, the samples are classified as radioactive, and require proper procedures to be in place when handling them, in order to limit exposure.
Table 4 presents the XRD analysis of Sample A and Sample B, particularly the main mineralogical phases. Both slag samples contained higher concentrations of spinel phase which characteristically contain no REE. Sample B contained a higher concentration of perovskite phase than Sample A. Sample A contained a higher amount of pyrophanite due to higher concentration of MnO in the slag. The REEs in Sample A were contained predominantly in the needle-shaped Ca silicate phase, whilst in Sample B they occurred predominantly in the perovskite phase.

Table 3. Base metals, REE, Th and U analysis in the head “slag” samples

<table>
<thead>
<tr>
<th>Sample</th>
<th>Al</th>
<th>Ca</th>
<th>Fe</th>
<th>Mg</th>
<th>Mn</th>
<th>Si</th>
<th>Ti</th>
<th>La</th>
<th>Ce</th>
<th>Pr</th>
<th>Nd</th>
<th>Sm</th>
<th>Gd</th>
<th>REE</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>%</td>
<td>%</td>
<td>%</td>
<td>%</td>
<td>%</td>
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<td>%</td>
<td>ppm</td>
<td>ppm</td>
<td>ppm</td>
<td>ppm</td>
<td>ppm</td>
<td>ppm</td>
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</tr>
<tr>
<td>A</td>
<td>8.49</td>
<td>3.15</td>
<td>6.44</td>
<td>2.55</td>
<td>11.2</td>
<td>10</td>
<td>5.64</td>
<td>15038</td>
<td>26277</td>
<td>2667</td>
<td>9034</td>
<td>1374</td>
<td>1365</td>
<td>5.96</td>
</tr>
<tr>
<td>B</td>
<td>10.8</td>
<td>9.53</td>
<td>4.33</td>
<td>1.95</td>
<td>6.68</td>
<td>8.03</td>
<td>7.52</td>
<td>19737</td>
<td>33510</td>
<td>3483</td>
<td>12971</td>
<td>1871</td>
<td>1721</td>
<td>7.96</td>
</tr>
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</table>

Table 4. XRD analysis of the head samples

<table>
<thead>
<tr>
<th>Compound</th>
<th>Formula</th>
<th>Sample A</th>
<th>Sample B</th>
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</thead>
<tbody>
<tr>
<td>Spinel</td>
<td>(Mg, Mn)Al₂O₄</td>
<td>xxx</td>
<td>xxx</td>
</tr>
<tr>
<td>Perovskite</td>
<td>Ca(TiO₃)</td>
<td>xxx</td>
<td>xxx</td>
</tr>
<tr>
<td>Pyrophanite</td>
<td>(Mn,Mg)(TiO₃)</td>
<td>xxx</td>
<td></td>
</tr>
<tr>
<td>Feldspar</td>
<td>(K,Na,Ba)AlSiO₃O₈</td>
<td>xxx</td>
<td>xxx</td>
</tr>
<tr>
<td>Pyroxene</td>
<td>(Ca,Na)Fe(Si₂O₆)</td>
<td>xx</td>
<td></td>
</tr>
<tr>
<td>Quartz</td>
<td>SiO₂</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Barium Niobium Titanium Oxide</td>
<td>Ba₃(Nb₄,Ti)O₉</td>
<td>x</td>
<td>x</td>
</tr>
</tbody>
</table>

Experimental description

The feed solids were slurried in deionised water to 20% (m/m) pulp density and heated to the required temperature. HCl was used to control the slurry to the required pH or to obtain the required acid addition. Process parameters such as pH, temperature, and oxidation-reduction potential (ORP) as well as reagent additions, were recorded at hourly intervals. At the end of the test, the slurry was filtered. A sample of the unwashed leach residue was removed for the determination of moisture content. The remainder of the filter cake was weighed and washed three times by repulping in pH 3 acidified, deionised water at a ratio of 2.5 times the wet cake mass to liquor ratio. The washed cake was weighed, and a representative sample taken to determine the moisture content. A representative sample of the dried sample was taken and assayed for total REE, U, Th, and base metals.

LEACHING RESULTS

Extraction efficiencies

The testwork program for this study was aimed mainly at evaluating the effect of various acids and acid strengths on the recovery of REE, as well as the level of impurity contamination. Only the results for the leaching using hydrochloric acid (HCl) are presented in this paper. The solid based REE and impurity leach efficiencies obtained in the leach tests are presented in Table 5.
Table 5. The solid based REE and impurity leach efficiencies of sample A and B

<table>
<thead>
<tr>
<th>Test description</th>
<th>pH1</th>
<th>pH2</th>
<th>1M</th>
<th>2M</th>
<th>3M</th>
<th>pH1</th>
<th>pH2</th>
<th>1M</th>
<th>2M</th>
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<td>60</td>
<td>60</td>
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<td>60</td>
<td>60</td>
<td>60</td>
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<tr>
<td>TREE concentration, mg/L</td>
<td>11418</td>
<td>9834</td>
<td>7127</td>
<td>12736</td>
<td>17194</td>
<td>2456</td>
<td>2212</td>
<td>1469</td>
<td>2630</td>
<td>2981</td>
<td>11407</td>
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<tr>
<td>MREE concentration, mg/L</td>
<td>628</td>
<td>536</td>
<td>359</td>
<td>690</td>
<td>935</td>
<td>159</td>
<td>140</td>
<td>76</td>
<td>164</td>
<td>194</td>
<td>793</td>
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<td>HREE concentration, mg/L</td>
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<td>601</td>
<td>411</td>
<td>780</td>
<td>1043</td>
<td>448</td>
<td>404</td>
<td>211</td>
<td>476</td>
<td>540</td>
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<td>TREE leach efficiency, %</td>
<td>97</td>
<td>86</td>
<td>41</td>
<td>80</td>
<td>94</td>
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<td>86</td>
<td>40</td>
<td>79</td>
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<td>15</td>
<td>8</td>
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<td>HREE leach efficiency, %</td>
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<td>87</td>
<td>39</td>
<td>82</td>
<td>94</td>
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<td>44</td>
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<td>120</td>
<td>264</td>
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<td>455</td>
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<td>122</td>
<td>269</td>
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<td>Acid consumption, kg/t solids</td>
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<td>298</td>
<td>102</td>
<td>240</td>
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<td>436</td>
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<td>5.4</td>
<td>4.9</td>
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<td>6.1</td>
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<td>6.8</td>
<td>14.3</td>
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<td>Mn concentration, g/L</td>
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<td>10.3</td>
<td>9.4</td>
<td>12.4</td>
<td>14.7</td>
<td>8.3</td>
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<td>0.7</td>
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<td>6.4</td>
<td>0.5</td>
<td>5</td>
<td>13.7</td>
<td>17.7</td>
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<td>4.7</td>
<td>4.1</td>
<td>5.4</td>
<td>6.9</td>
<td>7</td>
<td>6.1</td>
<td>5.8</td>
<td>7.8</td>
<td>9.3</td>
<td>20.2</td>
</tr>
</tbody>
</table>
The REE extractions in the tests for Sample A increased with increasing acid addition. Highest REE recoveries were obtained at pH 1 with extraction efficiencies of 97%. The REE recoveries were reduced slightly at a pH value of 2 where 80% total REE (TREE) extraction was achieved. During the controlled acid concentration leaching tests, the REE leaching efficiency also increased with increasing acid concentration between 1M and 3M. There was not much difference between the extraction of light REE (LREE), middle REE (MREE), and heavy REE (HREE) per test. Mn, Fe, Ca, and Al were the major impurities in the pregnant leach solution (PLS). Similar to the REE trend, impurity dissolution increases with increasing acid strength.

For Sample B, REE leaching was generally poor in all the tests, with the highest extraction of 46% TREE obtained with 10M HCl. MREE were the most resistant of the REEs, while HREE were easier to dissolve compared to the rest of the REE. The poor dissolution of REE in Sample B was linked to its association of the REE with the stable perovskite phase (CaTi mineral) which hosted a significant portion of the REE. Similarly to Sample A; Fe, Mn, Al, and Ca were the dominant impurities in Sample B. The concentrations of Fe (14.3 g/L), Mn (16.6 g/L), Al (17.7 g/L), and Ca (20.2 g/L) in the 10 M HCl test were much higher than the TREE (11.4 g/L). This indicates that a fair portion of the acid was consumed by impurities rather than by the dissolution of REE.

Mineralogy of washed residues

The mineralogical analysis of the residue of Sample A and Sample B was conducted to understand how the phases were affected during the leach step. SEM imaging with EDS using element maps provided confirmation of the presence of REE minerals in the perovskite mineral phase in both samples. However due to the higher concentration of the perovskite phase in Sample B than Sample A, the REE detected were proportionally higher. This result suggests that the REE that were associated with the MnCaSi and needle phases were solubilized, and limited or completely no dissolution of REE occurred in the perovskite phase. This correlates well with the leach results which showed poor REE dissolution in sample B.

Overall the results for the HCl leach tests have shown that REE could be recovered from the slag material. The results showed that it was much easier to recover REE from the high-Mn slag (Sample A) compared to the low-Mn slag (Sample B). It is critical that perovskite is minimized. Alternatively, other methods of cracking this phase must be investigated.

**FLOWSHEET OPTIONS**

The envisaged flowsheet for the PyEarth™ process comprises two sections: (1) smelting of the iron-rich ore (2) leaching of the resulting slag and separation and purification of the rare earth oxides. Figure 4 presents the generic flowsheet of the PyEarth process. This flowsheet includes a DC arc furnace where no flux is added to produce a high-MnO slag with < 10% REE, and pig iron. The slag is subjected to direct HCl leaching where the REE are leached at efficiencies of up to 95%. The impurities are precipitated out of the REE leach solution, followed by precipitation of REE and the recovery of HCl and CaSO₄ by addition of H₂SO₄.

Other possible flowsheets for this process could include prior extraction of the Mn into a manganese salt from the MnO-rich slag, followed by direct HCl leaching of the REE residue. Alternatively, a fluxed smelting process where the MnO is reduced to the alloy phase, leaving REE richer slag, is currently being optimized at Mintek.
CONCLUSIONS AND RECOMMENDATIONS

The critical steps of the PyEarth™ process developed at Mintek – namely the concentration through direct smelting, and the leaching of the resulting rare earth rich slag – were demonstrated successfully.

Direct smelting, under various conditions, of an iron-rich rare earth ore sample containing 2.3% REE and 33% Fe in Mintek’s 200 kW DC pilot furnace resulted in the production of slags of various MnO and REE contents, as well as pig iron of various compositions. High-manganese slags containing up to 8% REE were generated in the unfluxed smelting conditions, while the lime fluxing recipes produced low-manganese slags containing slightly more than 10% REE. More than 98% of the iron in the ore was recovered into a saleable iron-manganese alloy, while rare earth oxides remained fully unreduced in the slag phase. The production of rare earth rich slag through direct smelting was found to be efficient, flexible, and controllable.

Leaching of a high-manganese slag sample with HCl resulted in over 95% TRE extraction efficiencies, while poor efficiencies of up to 46% REE were achieved when leaching a low-manganese slag sample due to its high perovskite content.

The PyEarth process was found to be suitable for the extraction of rare earths from dormant southern African iron-rich rare earth deposits. The current pilot work was also able to identify research areas for the improvement and optimization of this process.

REFERENCES


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