Towards zero-waste valorisation of rare-earth-containing industrial process residues: a critical review

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Abstract
The supply risk for some critical rare-earth elements (REEs), which are instrumental in many cleantech applications, has sparked the development of innovative recycling schemes for End-of-Life fluorescent lamps, permanent magnets and nickel metal hydride batteries. These waste fractions represent relatively small volumes, albeit with relatively high rare-earth contents. Nevertheless, rare earths are also present in lower concentrations in a multitude of industrial process residues, such as phosphogypsum, bauxite residue (red mud), mine tailings, metallurgical slags, coal ash, incinerator ash and waste water streams. This review discusses the possibilities to recover rare earths from these “secondary resources”, which have in common that they contain only low concentrations of rare-earth elements, but are available in very large volumes and could provide significant amounts of rare earths. The success rate is set to increase if the rare-earth recovery from these industrial waste streams is part of a comprehensive, zero-waste, “product-centric” valorisation scheme, in which applications are found for the residual fractions that are obtained after removal of not only the rare earths but also other valuable (base) metals.

Keywords: bauxite residue; lanthanides; phosphogypsum; rare earths; red mud; zero-waste valorisation; metal recovery; mine tailings; slag and ash.
1. Introduction

For more than two decades, at least 95% of the annual global supply of the rare earths (REEs) has been provided by Chinese rare-earth producers. The tightening of rare-earth export quotas has caused supply risks outside China, as evidenced by the rare-earth crisis of 2011 with record high prices. This situation endangers the secure and sustainable supply of these critical (clean) technology metals, in particular the heavy rare earths, to many economies in the world. Paradoxically, on the positive side, this situation has also stimulated other countries to look for alternative rare-earth resources and to develop their own rare-earth industry. To tackle the rare-earth supply challenge, a threefold approach can be proposed. This strategy is particularly relevant for the heavy rare earths for which the supply risk is far greater than for light rare earths such as lanthanum and cerium. A first component of this strategy is to substitute critical rare earths by less critical metals. Secondly, the supply risk can be mitigated by investing in sustainable primary mining from old or new rare-earth deposits. Mining companies are now actively seeking for new exploitable rare-earth deposits, while old mines are being re-opened. A good example is the Mountain Pass Mine in California, which restarted production in 2012. Nevertheless, many non-Chinese mines, such as the Mountain Pass mine, are mostly light rare earth mines and, hence, do not deliver a meaningful stream of the most critical heavy rare earths. Furthermore, many countries such as Japan and most EU Member States do not possess any type of primary rare-earth deposits on their territory. Consequently, they will have to invest in technospheric mining (Johansson et al., 2013) of secondary rare-earth containing resources in order to obtain a source of both light rare earths and, particularly, heavy rare earths. This is the third component of the strategy.

Technospheric mining, however, can take many forms. With respect to (critical) metal containing streams, such mining incorporates (1) the direct (preconsumer) recycling of metal scrap and swarf generated during the production of metal based (intermediate) products (as
for instance NdFeB magnets) and (2) the (postconsumer) recycling and/or urban mining of, respectively, flows and stocks of complex, multi-material, metal-containing products (as for instance a hybrid electric vehicle). (3) landfill mining of historic urban solid waste (Jones et al., 2011), (4) metal recovery from flows of industrial process residues from primary and secondary metal production and, finally, (5) metal recovery from stocks of landfilled industrial process residues, as shown in Figure 1. When considering the potential for rare-earth recovery from these five secondary sources, it is clear that at present most attention is going out to the direct recycling of scrap and the recycling/urban mining of End-of-Life rare-earth containing products. The state-of-the-art in this domain has been described in detail in recent review papers (Binnemans et al., 2013a; Binnemans and Jones, 2014; Tanaka et al., 2013; Anderson et al., 2013; Innocenzi et al., 2014). The most interesting (heavy) rare earths sources intended for recycling/urban mining are permanent magnets and lamp phosphors (Binnemans et al., 2013a).

In contrast to recycling of rare earths from End-of-Life products, much less attention has been devoted to previously landfilled stocks and freshly produced flows of rare-earth-containing industrial process residues (Binnemans et al., 2013c) (top part in Figure 1). In general, these secondary resources contain much lower concentrations of rare earths than the End-of-Life consumer goods that are considered for direct recycling or postconsumer recycling/urban mining activities. Nevertheless, the volumes of these residues are enormous so that the total amounts of rare earths locked in these residues are also very large and may secure an independent source of rare earths as well as shield resource-poor countries from export quotas and price fluctuations. This review paper gives an overview of the possibilities to recover rare earths from the most important industrial process residues. These include residues from metal production (as shown in Figure 1) – phosphogypsum, bauxite residue (red mud), mine tailings, metallurgical slags – and industrial process residues from thermal
treatment facilities (coal ash, incinerator ash). Likewise, it is investigated if waste water streams may be a source of rare-earths as well. In all cases the zero-waste valorisation concept is promoted. For more background on the extractive metallurgy of rare earths, the reader is referred to general references (Gupta and Krishnamurthy, 2004; Gupta and Krishnamurthy, 1992; Habashi, 2013; Xie et al., 2014).

Figure 1: Closing the loop through technospheric mining, revealing the importance of metal recovery from both flows and stocks of industrial process residues from primary and secondary metal production.
2. Phosphogypsum

2.1. Formation and composition of phosphogypsum

Phosphogypsum is the main by-product of the production of phosphoric acid ($H_3PO_4$) by sulphuric acid ($H_2SO_4$) digestion of a concentrated slurry of pulverised phosphate ores (Koopman and Witkamp, 2000). Phosphoric acid is an important raw material for the manufacturing of phosphate fertilisers. Phosphate ores can be divided into two main types according to their origin: sedimentary and igneous phosphate rock (Habashi, 1998).

Sedimentary phosphate rock (also known as phosphorite) represents about 85–90% of the world reserves and is found in Florida, Morocco and the Middle East. Igneous phosphate rock represents the remaining 10–15% of the world reserves. It is found in the Kola Peninsula (Russia) and Brazil. Phosphate rock varies widely in composition (Becker, 1989). Apatite is the main phosphate mineral in most phosphate deposits (Rutherford et al., 1994). In sedimentary phosphate rock, apatite occurs in an amorphous form: francolite. Francolite has a complex chemical composition and can be represented by the formula $(Ca,Mg,Sr,Na)_{10}(PO_4,SO_4,CO_3)_6F_{2-3}$ (Benmore et al., 1983). In igneous phosphate rock, apatite occurs as the variety fluoroapatite, $Ca_{10}(PO_4)_6F_2$. Phosphate rock also contains trace amounts of many other elements, including thorium, uranium and rare earths. The rare-earth content depends on the type of phosphate rock (Habashi, 1998; Habashi, 1985) (Table 1).

Sedimentary phosphate rock contains 0.01–0.1 wt% of rare earths, but also about 0.01 wt% of uranium. Igneous phosphate rock is much richer in rare earths than sedimentary phosphate rock (1–2 wt%) and contains only very small amounts of uranium. The potential of phosphate rock as a source of rare earths has already been recognised in the 1960s (Anonymous, 1966). Analysis of phosphate rock from Florida gave a total rare-earth content of 0.059%, with the main elements being lanthanum (0.015 wt%), cerium (0.012 wt%), yttrium (0.011 wt%) and neodymium (0.007 wt%). The scandium concentration was very low: 0.0003 wt% (3 ppm). It
was estimated that the 6 million tonnes of phosphate rock that have been processed in the USA in 1964 could have yielded 3,500 tonnes of rare earths (cf. estimated worldwide REO production in 2013: 110,000 tonne (Gambogi, 2014). Cerium, lanthanum and neodymium account for 80% of the total rare-earth content of phosphate rock (Habashi, 1985).

Table 1. Rare-earth content of phosphate rock. Adapted from (Habashi, 1985).

<table>
<thead>
<tr>
<th>Locality</th>
<th>Total rare-earth oxides (wt%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kola (Russia)</td>
<td>0.8–1.0</td>
</tr>
<tr>
<td>Florida (USA)</td>
<td>0.06–0.029</td>
</tr>
<tr>
<td>Algeria</td>
<td>0.13–0.18</td>
</tr>
<tr>
<td>Morocco</td>
<td>0.14–0.16</td>
</tr>
<tr>
<td>Tunisia</td>
<td>0.14</td>
</tr>
<tr>
<td>Egypt</td>
<td>0.028</td>
</tr>
<tr>
<td>Vietnam</td>
<td>0.031</td>
</tr>
</tbody>
</table>

The \( \text{H}_2\text{SO}_4 \) digestion of fluoroapatite can be represented by the following chemical reaction scheme (Habashi, 1985):

\[
\text{Ca}_{10}(\text{PO}_4)_6\text{F}_2+10 \text{H}_2\text{SO}_4+20 \text{H}_2\text{O} \rightarrow 6 \text{H}_3\text{PO}_4+10 \text{CaSO}_4 \cdot 2\text{H}_2\text{O}+ 2 \text{HF} \tag{1}
\]

Depending on the production method, the phosphogypsum formed primarily consists of either calcium sulphate dihydrate, \( \text{CaSO}_4 \cdot 2\text{H}_2\text{O} \) (gypsum) or calcium sulphate hemihydrate, \( \text{CaSO}_4 \cdot \frac{1}{2}\text{H}_2\text{O} \), and it contains also small amounts of silica, fluoride compounds and unreacted phosphate rock. The amount of phosphogypsum produced during phosphoric acid production exceeds the mass of the product, i.e. 4.5 to 5.5 tonnes of phosphogypsum are generated per
tonne of phosphorus pentoxide (El-Didamony et al., 2012). Phosphogypsum is stored in stacks. The stacks form as the slurry containing the by-product phosphogypsum is pumped from the phosphoric acid plant onto a disposal site. The major component of the material in these waste piles is calcium sulphate dihydrate. In Table 2, the average composition of phosphogypsum produced via different processes is shown (Rutherford et al., 1994).

Table 2. Typical elemental composition (wt%) of phosphogypsum prepared via different processes. Adapted from (Rutherford et al., 1994).

<table>
<thead>
<tr>
<th>Component</th>
<th>Dihydrate</th>
<th>Hemihydrate</th>
<th>Hemi-dihydrate</th>
</tr>
</thead>
<tbody>
<tr>
<td>CaO</td>
<td>32.5</td>
<td>36.9</td>
<td>32.2</td>
</tr>
<tr>
<td>SO₃</td>
<td>44.0</td>
<td>50.3</td>
<td>46.5</td>
</tr>
<tr>
<td>P₂O₅</td>
<td>0.65</td>
<td>1.50</td>
<td>0.25</td>
</tr>
<tr>
<td>F</td>
<td>1.2</td>
<td>0.8</td>
<td>0.5</td>
</tr>
<tr>
<td>SiO₂</td>
<td>0.5</td>
<td>0.7</td>
<td>0.4</td>
</tr>
<tr>
<td>Fe₂O₃</td>
<td>0.1</td>
<td>0.1</td>
<td>0.05</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>0.1</td>
<td>0.3</td>
<td>0.3</td>
</tr>
<tr>
<td>MgO</td>
<td>0.1</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>H₂O (crystalline)</td>
<td>19.0</td>
<td>9.0</td>
<td>20.0</td>
</tr>
</tbody>
</table>

The rare earths are concentrated in phosphogypsum during the production of phosphoric acid. About 70 to 85% of the rare earths originally present in the phosphate rock end up in the phosphogypsum. The remainder stays dissolved in the leaching solution, which also contains the phosphoric acid. A smaller part of the rare earths is incorporated in the phosphogypsum if the leaching is performed at lower temperatures or with less concentrated H₂SO₄ (Habashi, 1985). The concentrations of the different rare earths in a phosphogypsum sample is shown in
The average concentration of rare earths in phosphogypsum is 0.4 wt% (Habashi, 1985). This seems to be a very low value compared to the rare-earth content of ore minerals such as bastnäsite, monazite or xenotime. Indeed, the rare-earth deposits that are currently mined have rare-earth oxide (REO) contents between 3 and 15 wt%. However, the estimated global production of phosphate rock in 2013 was 224 million tonnes (Jasinski, 2014), so that the total amount of rare earths in the mined phosphate rock is also large in absolute terms. This indicates that phosphate rock and the derived product phosphogypsum may become valuable sources of rare earths in the near future, provided the rare-earth prices are high enough. In comparison with red mud (see section 3), phosphogypsum contains lower concentrations of scandium.

Table 3. Concentrations of rare earths in a phosphogypsum sample of the Belgian company Prayon SA, dried at 250 °C (Germeau et al., 2013).

<table>
<thead>
<tr>
<th>Element</th>
<th>Concentration (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>La</td>
<td>1450</td>
</tr>
<tr>
<td>Ce</td>
<td>2310</td>
</tr>
<tr>
<td>Pr</td>
<td>235</td>
</tr>
<tr>
<td>Nd</td>
<td>899</td>
</tr>
<tr>
<td>Sm</td>
<td>163</td>
</tr>
<tr>
<td>Eu</td>
<td>34.9</td>
</tr>
<tr>
<td>Gd</td>
<td>98.7</td>
</tr>
<tr>
<td>Tb</td>
<td>7.45</td>
</tr>
<tr>
<td>Dy</td>
<td>45.5</td>
</tr>
<tr>
<td>Ho</td>
<td>7.37</td>
</tr>
<tr>
<td>Element</td>
<td>Concentration</td>
</tr>
<tr>
<td>---------</td>
<td>---------------</td>
</tr>
<tr>
<td>Er</td>
<td>15.7</td>
</tr>
<tr>
<td>Tm</td>
<td>1.36</td>
</tr>
<tr>
<td>Yb</td>
<td>5.6</td>
</tr>
<tr>
<td>Lu</td>
<td>0.568</td>
</tr>
<tr>
<td>Y</td>
<td>180</td>
</tr>
<tr>
<td>Sc</td>
<td>1.21</td>
</tr>
<tr>
<td>Total</td>
<td>5455</td>
</tr>
</tbody>
</table>

2.2. Recovery of rare earths from phosphogypsum

About 50% of the rare earths present in phosphogypsum can be recovered by leaching the phosphogypsum at ambient temperature with a 0.1 to 0.5 M H$_2$SO$_4$ solution in a solid-to-liquid ratio of 1:10 (Habashi, 1985). It is impossible to recover all the rare earths present in phosphogypsum without destruction of the phosphogypsum lattice. The leaching efficiencies can be increased by a gravity flow of the H$_2$SO$_4$ solution through a column packed with phosphogypsum (Lokshin et al., 2011). Another method for enhanced H$_2$SO$_4$ leaching is the mechanical activation of phosphogypsum by ball-milling (Todorovsky et al., 1997). The rare earths can be recovered from the leaching solution by precipitation, for instance as sodium rare-earth double sulphates (Lokshin and Tareeva, 2010) or rare-earth hydroxides (Habashi, 1985), or by solvent extraction (Habashi, 1985). In Figure 2, a flow sheet is shown for H$_2$SO$_4$ attack of phosphate rock, with precipitation of rare-earths as hydroxides (Habashi, 1985). Jarosinski and co-authors described a comprehensive process for the recovery of rare earths from phosphogypsum (Jarosinski et al., 1993) (Figure 3). The three basic steps are: (1) leaching of the rare earths from the phosphogypsum with dilute H$_2$SO$_4$; (2) concentration of the leachate by evaporation, followed by recovery of the rare earths by solvent extraction with nonyl–phenyl phosphoric acid (NPPA) or by selective precipitation with hydrofluoric acid;
production of anhydrite (CaSO$_4$) from the purified phosphogypsum by recrystallisation in concentrated H$_2$SO$_4$. Different process parameters had to be used, depending on whether the starting materials were calcium sulphate hemihydrate or dihydrate. The purified anhydrite can be used for production of plaster. Not all rare earths present in phosphate rock end up in phosphogypsum; a part dissolves in the aqueous phosphoric acid solution. The process was tested on a pilot scale, but showed to be too complicated and uneconomical to be industrialised (Wang et al., 2010).

In a two-stage hemihydrate–dihydrate process, calcium sulphate hemihydrate is precipitated and separated from phosphoric acid by filtration in the first step and the hemihydrate is hydrated to the dihydrate in the second step (Zielinski et al., 1993). The second step offers the best conditions for the rare earth recovery because the hydration proceeds via dissolution of hemihydrates salt. This dissolution step brings all rare earths into solution. On the other hand, rare-earth ions in solution inhibit the crystallisation of the dihydrate, so that the rare earths have to be removed from the solution. The rare earths can be recovered from the solution by solvent extraction with a dialkylphosphoric acid, followed by precipitation stripping to form sodium rare-earth double sulphates.

Leveque and Sabot developed a process for the comprehensive recovery of rare earths, uranium and thorium from phosphogypsum. This process consists of a leaching step with H$_2$SO$_4$, followed by solvent extraction of rare earths, uranium and thorium from the sulphate solution by a mixture of bis(octylphenyl)phosphoric acid and trioctylphosphine oxide (TOPO) (Leveque and Sabot, 1981). By using a mixture of extractants, a synergistic effect was achieved. H$_2$SO$_4$ was used as leachant, because (1) it showed selectivity towards the dissolution of the targeted metals, (2) it did not change the filterability of the phosphogypsum, (3) the process can be easily integrated in an existing phosphoric production plant, based on
H$_2$SO$_4$ attack of phosphate rock. Nevertheless, no evidence of commercialisation of this process was found in the literature.

The Belgian company Prayon SA developed a process for the recovery of rare earths from phosphogypsum (Germeau et al., 2013). Leaching is done by a dilute H$_2$SO$_4$ solution, while a suspension is stirred at high speeds (up to 6000 rpm). It is proposed to recover the rare earths from the leachate by solvent extraction or by precipitation with oxalic acid.
Figure 2. Flow sheet for the attack phosphate rock by H$_2$SO$_4$, with recovery of rare earths by precipitation as hydroxides. Adapted with permission from (Habashi, 1985). Copyright 1985 Society of Chemical Industry.
Figure 3. Process for recovery of rare earths from phosphogypsum, with production of purified anhydrite for use as plaster. Adapted with permission from (Jarosinski et al., 1993). Copyright 1993 Published by Elsevier B.V..

Preston and co-workers developed a process to recover rare earths from calcium sulphate sludge derived from the production of phosphoric acid from South-African phosphate rock (Preston and Du, 1998). Higher leaching efficiencies were obtained by leaching with HNO₃ instead of H₂SO₄, and maximum recovery of rare earths was observed for 3M HNO₃ (Preston et al., 1996a). The recovery markedly decreased with increasing temperatures. Interesting observations were made upon addition of Ca(NO₃)₂ to the HNO₃ leachant. Mixtures of Ca(NO₃)₂ and HNO₃ were found to be much more effective than solutions of only HNO₃. For instance, the treatment of a sludge containing 6.8% of rare-earth oxides by 1M HNO₃ resulted in 35% recovery of the rare earths after 72 h, whereas a mixture of 1M HNO₃ and 0.5M Ca(NO₃)₂ yielded a 76% recovery under similar conditions. The enhanced leaching was attributed to replacement of the rare-earth ion impurities in the calcium sulphate matrix by calcium ions. An alternative explanation is a lowering effect on the free fluoride ion concentration, which leads to a higher solubility of rare-earth ions. Addition of Ca(NO₃)₂ reduces the HNO₃ concentration required for maximum leaching from 3M to 1M. The dissolved rare-earth ions were recovered from the leach solution by solvent extraction with tributyl phosphate (TBP) or dibutylbutylphosphonate (DBBP) dissolved in xylene. The highest extraction percentages were observed for DBBP. For efficient extraction, the nitrate concentration in the leachate had to be increased by addition of 2.5M NH₄NO₃. By using a higher concentration of Ca(NO₃)₂ (3M) in the leachant, NH₄NO₃ could be omitted, and the DBBP could be replaced by the cheaper TBP. Stripping could be performed by water. Precipitation by oxalic acid and calcination gave a mixed rare-earth oxide. The process was
further tested by continuous counter-current tests on a lab scale, and also on a pilot-plant scale (Preston et al., 1996a). It was suggested to integrate the process in an existing plant that produces nitrate and phosphate fertilisers, in line with the previous suggestion to redesign the overall process, also upstream. Further research and development led to the invention of three additional processes: (1) a process for the precipitation of high-purity cerium dioxide and the recovery of a heavy rare-earth concentrate (Preston et al., 1996b), (2) a process for the separation of the light (La, Ce, Pr, Nd) and middle (Sm, Eu, Gd, Tb) rare-earth fractions by solvent extraction with bis(2-ethylhexyl)phosphoric acid (Preston et al., 1996c), and (3) a process for preparation of magnet-grade (95%) neodymium oxide from the light rare-earth fraction by extraction with the quaternary ammonium salt Aliquat 336 nitrate (Preston, 1996).

HNO₃ was found to be a more efficient leachant than H₂SO₄ for the recovery of rare earths from hemihydrates phosphogypsum, because of the high solubility of rare-earth phosphates and rare-earth alkali metal double sulphates in HNO₃ (Lokshin et al., 2002). Leaching with fresh portions of HNO₃ led to a 96% recovery of the rare earths, while only 7.8% of the calcium content went into solution. Based on these observations, the authors concluded that the rare earths do not isomorphously co-crystallise with CaSO₄·½H₂O, but rather exist as separate phases. Grinding of the hemihydrates phosphogypsum before leaching with HNO₃ led to a higher recovery of rare earths, because the grinding process removed the calcium sulphate crust that covers the surface of the grains of the rare-earth phases. In contrast to what was observed by Preston for leaching of calcium sulphate sludge (Preston et al., 1996a), Lokshin and co-workers did not observe enhanced leaching of hemihydrate phosphogypsum upon addition of Ca(NO₃)₂ to the HNO₃ leachant (Lokshin et al., 2002).

Although leaching of metal values from phosphogypsum is typically done by aqueous solutions of mineral acids, it is possible to largely suppress the aqueous phase and to make use of organic solvents. In that case one should use the term “solvometallurgical processing” (or
Solvometallurgical methods have been invented in the 1950s and early 1960s for the recovery of uranium from low grade complex ores (Grinstead, 1958; Grinstead, 1959b; Grinstead, 1959a; Kruse, 1963; Galvanek, 1959), but unfortunately these methods have been largely neglected ever since, notwithstanding their potential to recover metals from ores and industrial process residues. Two different approaches can be used: solvent leaching and slurry solvent extraction. In solvent leaching, the leaching process is performed with a complexing agent (acting as an extractant) dissolved in an organic solvent. The organic solvent can be miscible or immiscible with water, but it is important that both the extractant and the metal complex are soluble in the solvent. After leaching, the metals can be recovered from the organic phase. In slurry solvent extraction, the finely crushed ore is wetted by a small volume of acid solution, and this slurry is contacted with a water-immiscible organic phase, containing an extractant. This approach is similar to conventional solvent extraction, but the volume of the aqueous phase is significantly reduced. The solvometallurgical processing methods are generic in the sense that they can be applied to a host of different ore types. Besides the largely reduced consumption of acids and a major reduction in the volumes of the leaching solutions, the main advantage is the possibility to achieve a much higher selectivity for metal recovery compared to conventional hydrometallurgical leaching methods. El-Didamony and co-workers investigated the possibility to leach metals from phosphogypsum with organic extraction agents dissolved in kerosene (El-Didamony et al., 2012). Although the main aim of this study was the removal of radionuclides from phosphogypsum, the experimental results also indicated that this method was applicable to recover the rare earths. The best results were observed for leaching of phosphogypsum with tributylphosphate (TBP) dissolved in kerosene. This treatment removed 69.8% of the rare earths present in the solid material. The method was improved by using mixtures of TBP and TOPO (TOPO =
trioctylphosphine oxide) in kerosene instead of pure TBP mixtures. In this case the recovery of the rare earths equalled 80% (EI-Didamony et al., 2013).

2.3. Leaching of phosphate rock by nitric acid or hydrochloric acid

Instead of using H\textsubscript{2}SO\textsubscript{4}, phosphate rock can also be treated by HNO\textsubscript{3} or HCl. In contrast to leaching with H\textsubscript{2}SO\textsubscript{4} no solid residues comparable to phosphogypsum are created by leaching with HNO\textsubscript{3} or HCl, although still some solid residues will be formed. When phosphate rock is leached with a HNO\textsubscript{3} solution, all rare earths are solubilised as nitrates (Habashi, 1985). Phosphate rock, with the exception of silicate gangue material, is readily soluble in 50–60 % HNO\textsubscript{3} at 60 °C:

\[
\text{Ca}_{10}(\text{PO}_4)_6\text{F}_2+20 \text{HNO}_3 \rightarrow 6 \text{H}_3\text{PO}_4+10 \text{Ca(NO}_3)_2+2 \text{HF}
\] (2)

After dissolution of the phosphates, the slurry has to be filtered to remove silica. The solution is cooled to -5 °C to induce crystallisation of Ca(NO\textsubscript{3})\textsubscript{2}·4H\textsubscript{2}O, which is removed from the solution by centrifugation. The solution can be defluorinated by addition of NaNO\textsubscript{3} to precipitate Na\textsubscript{2}SiF\textsubscript{6}. The rare earths can be recovered by adjusting the pH to 0.3–1.4 with ammonia, resulting in the formation of a phosphate precipitate containing 80 to 100% of the rare earths (Habashi, 1985). Another method to recover the rare earths is to extract them with tributylphosphate (TPB) (Werner et al., 1966a; Werner et al., 1966b), but this solvent extraction step should be done before crystallisation of Ca(NO\textsubscript{3})\textsubscript{2}·4H\textsubscript{2}O, since the calcium ions enhance the extraction. After extraction, the rare-earths can be removed from the organic phase by stripping with water, followed by precipitation as hydroxides by addition of ammonia to the water phase. Uranium is also extracted by TBP. Selectivity can be obtained by pH control: uranium is extracted at very low pH values (0.6–0.7) and the extraction efficiency drops at higher pH values, whereas maximum recovery of rare earths can be
achieved at a pH of 1.0 where a precipitate starts to form (Habashi et al., 1986). Uranium can be extracted from the solution together with H₃PO₄ by tertiary amyl alcohol and can be stripped by a 50% NH₄NO₃ solution (Habashi and Awadalla, 1986). The co-extracted H₃PO₄ may be stripped with NH₃. The rare earths, left behind in the leachate solution, can be extracted by TBP. A flow sheet for the treatment of phosphate rock with HNO₃ and the recovery of the rare earths from the chloride solution is shown in Figure 4.

Figure 4. Proposed flow sheet for the treatment of phosphate rock with HNO₃, with recovery of rare earths. Adapted with permission from (Habashi et al., 1986). Copyright 1986 Society of Chemical Industry.
The HCl route has received less attention than the HNO₃ route. Just as in the case of HNO₃, digestion of phosphate rock with HCl will bring all the solid material in solution, with the exception of silicate gangue material (Habashi et al., 1987):

\[
\text{Ca}_{10}(\text{PO}_4)_6\text{F}_2 + 20 \text{HCl} \rightarrow 6\text{H}_3\text{PO}_4 + 10\text{CaCl}_2 + 2\text{HF} \quad (3)
\]

A flow sheet for the treatment of phosphate rock by HCl and the recovery of the rare earths from the chloride solution can be found in Figure 5. The phosphate rock is leached with a stoichiometric amount of HCl at 40 °C. Addition of NaCl to the leachate will cause removal of fluorine by precipitation of Na₂SiF₆. Radium can be removed from the solution by co-precipitation with BaSO₄ upon consecutive addition of BaCl₂ and Na₂SO₄. Uranium can be removed by solvent extraction with a dilute solution (5%) of TBP in an aliphatic diluent. H₃PO₄ is separated from CaCl₂ by solvent extraction with TBP. Stripping of H₃PO₄ from the organic phase by ammonia gives ammonium phosphate fertiliser. The CaCl₂ in the aqueous solution can be precipitated as gypsum by addition of H₂SO₄. In this way, HCl is reformed. In contrast to phosphogypsum, the gypsum formed by precipitation with H₂SO₄ from chloride solution is not radioactive. The rare-earth elements can be recovered from the solution by precipitation as phosphates by partial neutralisation of the acidic solution by Ca(OH)₂. Alternatively, the rare earths can be recovered from the solution as hydroxides by precipitation with ammonia, before gypsum is precipitated. The recovery of the rare earths by this process is about 60%. Attack of phosphate rock by HCl was recommended because it facilitates the recovery of rare earths compared to attack by H₂SO₄ (Shlewit, 2011).
Figure 5. Proposed flow sheet for the treatment of phosphate rock with HCl, with recovery of rare earths. Adapted with permission from (Habashi et al., 1987). Copyright 1987 Society of Chemical Industry.
Habashi proposed *in situ*, dump or vat leaching of phosphate rocks with dilute HNO₃ (20%) or HCl (10%) to get a solution of monocalcium phosphate, and to allow recovering uranium and rare earths (Habashi and Awadalla, 1988; Habashi, 1989; Habashi, 1994). The rare earths can be recovered from the nitrate leach solution at the natural pH of the leach solution by TBP and from the chloride leach solution by bis(2-ethylhexyl)phosphoric acid. Evaporation of the leach solution leads to formation of the double salts CaCl₂H₂PO₄·H₂O or Ca(NO₃)₂H₂PO₄·H₂O. The double salts can be decomposed at 200–250 °C to CaHPO₄.

Advantages of this process include (1) low capital costs, (2) elimination of the phosphogypsum problem, (3) enhanced possibility to recover rare earths, uranium and fluorine, and (4) enhanced possibility to control the radium content of the phosphate rock.

Phosphogypsum can be decomposed by reaction with ammonium carbonate, so that ammonium sulphate fertiliser and calcium carbonate are formed (Habashi, 1985; Burnett et al., 1996):

\[
\text{CaSO}_4·2\text{H}_2\text{O} + (\text{NH}_4)_2\text{CO}_3 \rightarrow (\text{NH}_4)_2\text{SO}_4 + \text{CaCO}_3 + 2\text{H}_2\text{O}
\]  

(4)

The rare earths report to CaCO₃ and can be recovered by dissolution of CaCO₃ in HNO₃, followed by removal of the rare earths by solvent extraction (Habashi, 1985). Alternatively, CaCO₃ can be calcined to CaO and this compound can be dissolved by leaching with an NH₄Cl solution. This results in a rare-earth rich residue.

2.4. *Recovery of rare earths from phosphoric acid*

Phosphoric acid produced by H₂SO₄ leaching contains about 1 g L⁻¹ of rare earths. This is only 15 to 30% of the rare earths originally present in the phosphate rock, since most of the rare earths report to the phosphogypsum. This is in contrast to uranium that largely reports to
the phosphoric acid solution (Bunus, 2000). Many researchers have investigated the possibility to recover rare earths and uranium from these phosphoric acid leaching solutions, and most of these methods were based on solvent extraction or ion-exchange (Kumar et al., 2011; Radhika et al., 2011; Kumar et al., 2010; Reddy et al., 2009; Bunus and Dumitrescu, 1992; Bunus et al., 1994; Bunus, 2000; Koopman et al., 1999b; Koopman et al., 1999a; Habashi, 1985). Lower temperatures, higher concentrations of phosphoric acid and larger liquid-to-solid ratios increase the rare-earth concentration in the phosphoric acid leach solution (Wang et al., 2010). Addition of surfactants enhances the growth of gypsum crystals and improves the leaching of rare earths (Wang et al., 2010). Non-equilibrium extraction via centrifugal contacting was able to increase the separation factor between rare earths and iron more than 250 times (Wang et al., 2011a).

3. Bauxite residue (red mud)

3.1. Formation and composition of bauxite residue

Bauxite, a mixture of impure hydrated aluminium oxides, is the most important aluminium ore. The estimated global production of bauxite in 2013 was 259 million tonnes (Bray, 2014). There are different classification schemes for bauxites, depending on their origin (Patterson et al., 1986). An often used method of classification divides the bauxites into laterite and karst types. The laterite-type bauxite deposits are the very large blanket deposits formed mainly by surficial weathering in tropical regions, and these bauxites are mostly lying on top of aluminosilicate rocks. The karst-type deposits are found on top of carbonate rocks. Laterite bauxites (about 88% of the global reserves) are found in Suriname, Guyana, Venezuela, Brazil, Guinea, India, Indonesia, Vietnam, Australia and the USA. Karstic bauxites (about 12% of the global reserves) are mainly found in Europe (Greece, France,
Hungary, Romania), Jamaica, Russia and China. Karstic-type bauxites are richer in rare earths compared to lateritic bauxite. Depending on the type of deposit, the principal aluminium-bearing components are Al(OH)₃ (gibbsite), γ-AlO(OH) (boehmite) and α-AlO(OH) (diaspore). Lateritic bauxites contain mainly gibbsite, whereas the karstic bauxites are rich in boehmite and diaspore. The chemical composition of representative bauxites is given in Table 4. Aluminium metal is produced from pure Al₂O₃, which is obtained via the Bayer process, where bauxite is digested in a concentrated NaOH solution at temperatures and pressures. This digestion in NaOH converts the aluminium minerals to sodium aluminate, which is soluble in the NaOH solution due to the amphoteric character of aluminium. The other components of bauxite do not dissolve or are converted to sparsely soluble compounds. The solution, which is rich in dissolved Al₂O₃, is separated from the bauxite residue (BR), more commonly known as red mud (Jones and Haynes, 2011; Power et al., 2011; Klauber et al., 2011; Grafe et al., 2011; Grafe and Klauber, 2011). In the translated Russian literature, the term “red sludge” is used. The pH of the wet red mud slurry varies between 9.7 and 12.9, with an average of 11.3 (Grafe et al., 2011). The bauxite residue composition depends on the type of bauxite and the process parameters of the Bayer process (Table 5) (Grafe et al., 2011). Bauxite residue has a complex mineralogy (Table 6). In Table 7, the main chemical components of different bauxite residues are given. Worldwide there is an annual bauxite residue production of 1.2×10⁸ tonnes (dry matter) and a total inventory of 3×10⁹ tonnes, stored in huge tailing ponds (International Aluminium Institute, 2013). These tailing ponds carry risks, as revealed in 2010 by the dam failure of the Ajka refinery in Hungary and the resulting loss of human lives and environmental catastrophe (Ruyters et al., 2011; Gelencser et al., 2011).
### Table 4. Chemical compositions of typical bauxites (in wt%). Adapted from (Patterson et al., 1986).

<table>
<thead>
<tr>
<th>Origin</th>
<th>Al$_2$O$_3$</th>
<th>SiO$_2$</th>
<th>Fe$_2$O$_3$</th>
<th>TiO$_2$</th>
<th>Loss on ignition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Australia</td>
<td>54.8</td>
<td>3.4–4.2</td>
<td>17.1</td>
<td>3.4</td>
<td>26.4</td>
</tr>
<tr>
<td>Brazil</td>
<td>55.9</td>
<td>4.8</td>
<td>9.4</td>
<td>1.3</td>
<td>28.6</td>
</tr>
<tr>
<td>France</td>
<td>53.0</td>
<td>7.8</td>
<td>21.4</td>
<td>2.6</td>
<td>13.3</td>
</tr>
<tr>
<td>Ghana</td>
<td>51.6</td>
<td>1.3</td>
<td>17.4</td>
<td>1.9</td>
<td>13.3</td>
</tr>
<tr>
<td>Greece</td>
<td>57.6</td>
<td>3.0</td>
<td>22.8</td>
<td>2.75</td>
<td>12.17</td>
</tr>
<tr>
<td>Guyana</td>
<td>55–61</td>
<td>1–10</td>
<td>0.8–5</td>
<td>2–5</td>
<td>30–35</td>
</tr>
<tr>
<td>Jamaica</td>
<td>49.1–50.6</td>
<td>0.7–6.1</td>
<td>18.9–20.5</td>
<td>2.5–2.7</td>
<td>24.6–27.3</td>
</tr>
<tr>
<td>Suriname</td>
<td>58.5–60.0</td>
<td>3.4–4.3</td>
<td>2.7–4.4</td>
<td>2.4–2.7</td>
<td>30.7–31.4</td>
</tr>
<tr>
<td>Arkansas (USA)</td>
<td>45–50</td>
<td>13</td>
<td>8</td>
<td>2.5–3</td>
<td>25</td>
</tr>
</tbody>
</table>

### Table 5. Process parameters of the Bayer process (Grafe et al., 2011).

<table>
<thead>
<tr>
<th></th>
<th>Gibbsite-rich bauxite</th>
<th>Boehmite-rich bauxite</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature (°C)</td>
<td>104–145</td>
<td>200–232</td>
</tr>
<tr>
<td>Pressure (atm)</td>
<td>1.0–3.0</td>
<td>6.0</td>
</tr>
<tr>
<td>Concentration NaOH (M)</td>
<td>3.6–8.9</td>
<td>3.6–5.0</td>
</tr>
</tbody>
</table>
Table 6. Mineralogical composition of bauxite residues. Adapted from (Grafe et al., 2011)

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Chemical formula</th>
<th>Range (wt%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hematite</td>
<td>$\alpha$-Fe$_2$O$_3$</td>
<td>7–29</td>
</tr>
<tr>
<td>Goethite</td>
<td>$\alpha$-FeOOH</td>
<td>7.3–24.3</td>
</tr>
<tr>
<td>Magnetite</td>
<td>Fe$_3$O$_4$</td>
<td>0–8</td>
</tr>
<tr>
<td>Diaspore</td>
<td>$\alpha$-AIOOH</td>
<td>0.5–0.6</td>
</tr>
<tr>
<td>Boehmite</td>
<td>$\gamma$-AIOOH</td>
<td>1–9.6</td>
</tr>
<tr>
<td>Gibbsite</td>
<td>Al(OH)$_3$</td>
<td>1–5</td>
</tr>
<tr>
<td>Quartz</td>
<td>SiO$_2$</td>
<td>1.2–4.9</td>
</tr>
<tr>
<td>Rutile</td>
<td>TiO$_2$</td>
<td>1.1–5.4</td>
</tr>
<tr>
<td>Anatase</td>
<td>TiO$_2$</td>
<td>0.3–5</td>
</tr>
<tr>
<td>Sodalite</td>
<td>Na$_8$[Al$_6$Si$<em>6$O$</em>{24}$][(OH)$_2$]</td>
<td>2.7–24</td>
</tr>
<tr>
<td>Cancrinite</td>
<td>Na$_6$[Al$_6$Si$<em>6$O$</em>{24}$]·2CaCO$_3$</td>
<td>0–51</td>
</tr>
<tr>
<td>Calcite</td>
<td>CaCO$_3$</td>
<td>1–11.2</td>
</tr>
<tr>
<td>Perovskite</td>
<td>CaTiO$_3$</td>
<td>0–11.5</td>
</tr>
</tbody>
</table>
Table 7. Elemental compositions of bauxites residues (wt%) Adapted from reference (Grafe et al., 2011).

<table>
<thead>
<tr>
<th>Bauxite Origin</th>
<th>Refinery</th>
<th>$\text{Al}_2\text{O}_3$</th>
<th>$\text{Fe}_2\text{O}_3$</th>
<th>$\text{SiO}_2$</th>
<th>$\text{TiO}_2$</th>
<th>$\text{CaO}$</th>
<th>$\text{Na}_2\text{O}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Greece</td>
<td>Aluminium de Greece</td>
<td>15.6</td>
<td>42.5</td>
<td>9.2</td>
<td>5.9</td>
<td>19.7</td>
<td>2.4</td>
</tr>
<tr>
<td>Hungary</td>
<td>Ajka</td>
<td>14.8</td>
<td>42.1</td>
<td>13.5</td>
<td>5.2</td>
<td>6.1</td>
<td>8.2</td>
</tr>
<tr>
<td>Turkey</td>
<td>Seydisehir</td>
<td>20.24</td>
<td>39.84</td>
<td>15.27</td>
<td>4.15</td>
<td>1.8</td>
<td>9.43</td>
</tr>
<tr>
<td>China</td>
<td>Shandong</td>
<td>6.9</td>
<td>12.8</td>
<td>19.1</td>
<td>3.43</td>
<td>46.0</td>
<td>2.37</td>
</tr>
<tr>
<td>China</td>
<td>Shanxi</td>
<td>7.3</td>
<td>6.8</td>
<td>13.9</td>
<td>2.5</td>
<td>33.9</td>
<td>2.7</td>
</tr>
<tr>
<td>India</td>
<td>Renukoot</td>
<td>21.9</td>
<td>28.1</td>
<td>7.5</td>
<td>15.6</td>
<td>10.2</td>
<td>4.5</td>
</tr>
<tr>
<td>India</td>
<td>Korba</td>
<td>19.4</td>
<td>27.9</td>
<td>7.3</td>
<td>16.4</td>
<td>11.8</td>
<td>3.3</td>
</tr>
<tr>
<td>India</td>
<td>Damanjodi</td>
<td>14.5</td>
<td>54.8</td>
<td>6.4</td>
<td>3.7</td>
<td>2.5</td>
<td>4.8</td>
</tr>
<tr>
<td>Australia</td>
<td>Pinjarra</td>
<td>17.1</td>
<td>36.2</td>
<td>23.8</td>
<td>3.9</td>
<td>3.9</td>
<td>1.6</td>
</tr>
<tr>
<td>Jamaica</td>
<td>Kirkvine</td>
<td>13.2</td>
<td>49.4</td>
<td>3.0</td>
<td>7.3</td>
<td>9.4</td>
<td>4.0</td>
</tr>
<tr>
<td>Guinea</td>
<td>Auginish</td>
<td>23.6</td>
<td>30.4</td>
<td>9.65</td>
<td>17.85</td>
<td>6.4</td>
<td>5.3</td>
</tr>
<tr>
<td>Ghana</td>
<td>Burntisland</td>
<td>23.43</td>
<td>36.31</td>
<td>18.25</td>
<td>5.97</td>
<td>4.38</td>
<td>12.36</td>
</tr>
</tbody>
</table>

3.2. Recovery of rare earths from red mud

All rare earths end up in the bauxite residue during the processing of bauxite by the Bayer process, because these elements are associated with iron and titanium minerals that remain unchanged (Derevyankin et al., 1981c). The enrichment factor of the rare earths in bauxite residue compared to bauxite is about a factor of two (Table 8) (Ochsenkühn-Petropoulou et al., 1994). The average concentration of rare earths in bauxite from the

26
Parnassos-Ghiona mountains in Greece is 506 ppm, whereas the average concentration of the resulting bauxite residue is 1040 ppm (Ochsenkühn-Petropoulou et al., 1994). It must be mentioned that Greek bauxite is often mixed with tropical bauxites during its processing to alumina. The rare-earth concentration in Jamaican red mud varies between 1500 and 2500 ppm (Wagh and Pinnock, 1987). Bauxite residue is rich in scandium: bauxite residue from Greece contains about 130 ppm of scandium, whereas bauxite residue produced from Jamaican bauxite contains up to 390 ppm (Wagh and Pinnock, 1987). However, the highest rare-earth concentrations can be found in the bauxite residue produced from bauxites from Moengo in Suriname, with scandium concentrations as high as 1700 ppm (Logomerac, 1971), while Russian bauxite residue was reported to contain 135 ppm of Sc$_2$O$_3$ (Fomin et al., 2004). These scandium concentrations are much higher than the average abundance of 22 ppm of scandium in the Earth’s crust, and point to a significant enrichment of scandium in bauxite and bauxite residue. Although minerals with a high scandium content do exist (e.g. thortveitite and kolbeckite), they only form very small deposits (Wang et al., 2011b). The lack of rich scandium deposits hampers the widespread use of scandium, although scandium could find useful applications as an alloying metal for aluminium (Royset and Ryum, 2005) and as a recyclable Lewis acid catalyst for the production of fine chemicals (Kobayashi, 1999). It can be expected that the demand of scandium will be increasing in the near future, due to the increasing need of scandium for preparing scandium-stabilised zirconia used in solid oxide fuel cells (Badwal et al., 2000). For these reasons, the recovery of scandium from red mud could be of high economic interest (Petrakova et al., 2014). Scandium represents more than 95% of the economic value of rare earths in red mud. Every alumina plant with a capacity of more than 1 million tonnes of Al$_2$O$_3$ has the potential of producing 150 tonnes of scandium or more (Yatsenko and Pyagai, 2010). However, very little information is available on the mineralogy of the rare-earth-rich phases in red mud. Allanite-(La) and dissakisite-(Ce) were
identified as the rare-earth phases in an Indian bauxite residue sample, containing 70 ppm of lanthanum and 100 ppm of cerium (Abhilash et al., 2014).

Table 8. Average concentration of rare earths in Greek bauxites and bauxite residues, and the enrichment factor of rare earths in bauxite residue. Adapted from reference (Ochsenkühn-Petropoulou et al., 1994). The values are expressed in ppm (g/tonne).

<table>
<thead>
<tr>
<th>Element</th>
<th>Bauxite</th>
<th>Bauxite residue</th>
<th>Enrichment factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>La</td>
<td>87.2</td>
<td>149.0</td>
<td>1.78</td>
</tr>
<tr>
<td>Ce</td>
<td>224.3</td>
<td>418.0</td>
<td>1.87</td>
</tr>
<tr>
<td>Pr</td>
<td>13.9</td>
<td>25.8</td>
<td>2.13</td>
</tr>
<tr>
<td>Nd</td>
<td>62.4</td>
<td>115.0</td>
<td>1.99</td>
</tr>
<tr>
<td>Sm</td>
<td>13.2</td>
<td>28.9</td>
<td>2.30</td>
</tr>
<tr>
<td>Eu</td>
<td>2.5</td>
<td>5.0</td>
<td>1.89</td>
</tr>
<tr>
<td>Gd</td>
<td>12.8</td>
<td>23.3</td>
<td>1.86</td>
</tr>
<tr>
<td>Tb</td>
<td>n.d.</td>
<td>n.d</td>
<td>n.d.</td>
</tr>
<tr>
<td>Dy</td>
<td>7.0</td>
<td>12.8</td>
<td>2.15</td>
</tr>
<tr>
<td>Ho</td>
<td>2.0</td>
<td>4.3</td>
<td>2.15</td>
</tr>
<tr>
<td>Er</td>
<td>8.1</td>
<td>17.2</td>
<td>2.15</td>
</tr>
<tr>
<td>Tm</td>
<td>n.d.</td>
<td>n.d</td>
<td>n.d.</td>
</tr>
<tr>
<td>Yb</td>
<td>8.0</td>
<td>15.6</td>
<td>1.99</td>
</tr>
<tr>
<td>Lu</td>
<td>1.4</td>
<td>2.4</td>
<td>1.76</td>
</tr>
<tr>
<td>Y</td>
<td>55.9</td>
<td>93.9</td>
<td>1.68</td>
</tr>
<tr>
<td>Sc</td>
<td>59.0</td>
<td>127.9</td>
<td>2.17</td>
</tr>
</tbody>
</table>
There are numerous studies on the extraction of rare earths, and of scandium in particular, from bauxite residue. The two main approaches are either purely hydrometallurgical flow sheets or a combination of pyrometallurgical and hydrometallurgical processes (Wang et al., 2011b). The purpose of the hydrometallurgical processes is to selectively leach the minor metals from bauxite residue. It has been an important discovery that the rare earths are readily leachable from bauxite residue by diluted mineral acids, whereas majority elements such as iron cannot be leached from bauxite under the same experimental conditions (Fulford et al., 1991a). A comparative study of leaching with different acids (HCl, HNO₃ or H₂SO₄), has shown that 0.5 M HNO₃ is the best leachant and that the leaching process can be performed at ambient temperatures and pressures (Ochsenkühn-Petropoulou et al., 1996). 80% of scandium and 96% of yttrium could be recovered, but the leaching procedure was less efficient for the light lanthanides (30 to 50% recovery). The leaching process with dilute HNO₃ has been performed at a pilot scale and optimised for the recovery of scandium from bauxite residue (Ochsenkühn-Petropoulou et al., 2002) (Figure 6). An issue related to leaching with HNO₃ is the difficulty to recover nitrate ions adsorbed to bauxite residue (Petrakova et al., 2014). Flushing with an excess of water create large volumes of waste water contaminated by nitrate ions. Removal of nitrate ions from waste water is difficult since most nitrate salts are water soluble, so that nitrates cannot be precipitated. An analytical method based on reverse-phase HPLC was developed for the determination and quantification of rare earths from red mud (Tsakanika et al., 2004). Dilute H₂SO₄ is the most efficient leachant for leaching of scandium from Australian bauxite residue (Wang et al., 2013), in contrast to leaching of scandium from Greek bauxite residue. This difference in leaching behaviour was attributed to the differences in mineralogical composition of these two types of bauxite residue (Wang et al., 2013). Subsequent tests to recover scandium from the leachate were not done on real samples but rather on synthetic
ones prepared by dissolution of rare-earth sulphate salts. Scandium could be extracted from the leachate by a mixture of bis(2-ethylhexyl)phosphoric acid (D2EPHA) and tributylphosphate (TBP) at pH 0.4, with hardly any co-extraction of iron. The authors proposed a conceptual flow sheet for the recovery of Sc$_2$O$_3$ from bauxite residue (Figure 7). Leaching with HCl is very efficient for recovery of rare earths, but a main disadvantage of this method is that large amounts of iron are codissolved (Borra et al., 2015).

H$_2$SO$_4$ leaching studies on Indian bauxite residue sample containing 70 ppm of lanthanum and 110 ppm of cerium showed that maximum recovery was obtained with a 3M H$_2$SO$_4$ solution (Abhilash et al., 2014). However, while maximum recovery of lanthanum was achieved at 35 °C, the maximum recovery of cerium occurred at 75 °C. It must be realised that upon leaching of bauxite residue with acids, part of the acid is consumed for neutralisation of the highly alkaline bauxite residue. After leaching, the rare earths can be recovered from the leachate by selective precipitation as oxalate, or by solvent extraction. Different ion-exchange resins have been tested to selectively take up scandium from the leachate after leaching bauxite residue with H$_2$SO$_4$ (Smirnov and Molchanova, 1997). Scandium could be eluted from the loaded resin by a Na$_2$CO$_3$ solution and scandium was precipitated as Sc(OH)$_3$ from the concentrated eluate by addition of NaOH solution. The literature data show that the acid leaching results strongly depend on the type of bauxite residue, because the mineralogical composition of the bauxite residue determines which acid is best suited for leaching. Therefore, a recovery process for rare earths from bauxite residue needs to be tailored for a given type of bauxite residue. For instance, results obtained for experiments on Greek bauxite residue samples are not necessarily applicable to Australian bauxite residue. The close association of scandium with iron minerals is evident from the correlation between the dissolution of iron and the recovery of scandium (Borra et al., 2015). About 50% of the scandium in bauxite residue samples can be recovered without bringing too
much iron into solution. However, trying to recover more than 50% of the scandium does inevitably result in dissolution of a major part of the iron. It is not be possible to recover 100% of the scandium present in the bauxite residue without dissolving all of the iron.
Figure 6. Conceptual flow sheet for the recovery of scandium from bauxite residue by a HNO₃ leaching method. Adapted with permission from (Ochsenkühn-Petropoulou et al., 2002). Copyright 2002 American Chemical Society.
The rare earths can be selectively dissolved by digesting bauxite residue with a dilute acid solution made by saturating water with SO₂ (forming sulphurous acid), leaving most of the iron undissolved (Fulford et al., 1991a; Fulford et al., 1991b; Kuwabara et al., 1967). Also sodium, aluminium and silicon present in the form of sodalite are brought into solution. The rare earths can be recovered by solvent extraction. In the Canadian Orbite process, bauxite residue is leached with a concentrated HCl solution in an autoclave at temperatures between 125 and 150 °C (Boudreault et al., 2013b) (Figure 8). This brings aluminium, iron and the rare earths into solution. By saturating the leachate with HCl, it is possible to induce crystallisation of AlCl₃·6H₂O. Solid AlCl₃·6H₂O is separated from the solution and converted by calcinations to Al₂O₃, with recovery of HCl gas. The remaining solution is concentrated and FeCl₃ is hydrolysed to Fe₂O₃ (hematite) between 155 to 170 °C. After removal of Fe₂O₃ a solution of rare earths is obtained, from which the rare earths can be recovered by solvent extraction. A main issue of a process based on HCl is the highly corrosive character of HCl gas and HCl solutions. In 2013 the Japanese company Nippon Light Metals has set up pilot plant studies on the grounds of the Jamaica Bauxite Institute (JBI) in Hope Gardens, St. Andrew (Jamaica) to extract rare earths and especially scandium from bauxite residue (Richardson, 2013). Technical information on this Japanese process is not available. In a recent review, leaching with H₂SO₄ is considered to the economically most efficient method for recovery of rare earths from bauxite residue (Petrakova et al., 2014).
Figure 8. Flow sheet of the Canadian Orbite process for zero-waste valorisation of bauxite residue, including recovery of rare earths. Adapted from (Boudreault et al., 2013b).

Meanwhile, Russian researchers have reported that it is possible to efficiently leach scandium from bauxite residue in alkaline conditions (Yatsenko and Pyagai, 2010; Fomin et al., 2004). The development of the process was based on the observation that scandium forms a soluble anionic complex compound with carbonate ions (Pasechnik et al., 2004). Whereas only 0.43 g L\(^{-1}\) Sc\(_2\)O\(_3\) can be dissolved at 25 °C in a solution containing 100 g L\(^{-1}\) Na\(_2\)CO\(_3\), scandium is much better soluble in a NaHCO\(_3\) solution: 16.7 g L\(^{-1}\) Sc\(_2\)O\(_3\) can be dissolved in a solution containing 100 g L\(^{-1}\) NaHCO\(_3\) (Fomin et al., 2004). In one version of the process, bauxite residue is leached with solutions of Na\(_2\)CO\(_3\) and/or NaHCO\(_3\) (5–12%) in three steps at 50 °C for 2 h at a solid-to-liquid ratio of 1:2.5 to 1:5. The scandium is precipitated from the leachate by adding a solution of sodium aluminate or sodium zincate (prepared by dissolving...
Al₂O₃ or ZnO in an aqueous NaOH solution) while heating the solution for 2 h at 80 °C. The resulting precipitate is separated from the solution, washed with an aqueous NaOH solution (10–25%) with simultaneous heating until boiling, followed by filtration and washing with an aqueous NaOH solution (1–5%). In a next step, the solid is dissolved in an aqueous HCl solution (1–5%) and the remaining solid is filtered off. Scandium is precipitated as Sc(OH)₃ by addition of an excess of an aqueous NH₃ solution (10–25%) to the filtrate (Diev et al., 2003). In a modified version of the process, leaching is carried out by passing a gas mixture containing air with CO₂ (10–17 vol.%) through the mixed Na₂CO₃/NaHCO₃ solutions (Yatsenko et al., 2005). The leachate is first treated with an organic flocculant to precipitate titania, is subsequently electrolysed to remove impurities, and finally scandium is precipitated by sodium zincate in combination with an organic flocculant. In a further improved version of the process, multiple leaching steps with mixtures of sodium carbonate and sodium hydrogen carbonate are carried out while a flow of flue gases from the bauxite sintering or calcination furnaces (containing CO₂) is passed through the solution (Yatsenko and Pyagai, 2010; Pyagai et al., 2012). Scandium is co-precipitated with zinc oxide. The advantage of alkaline leaching is that no acid is consumed to acidify the highly alkaline red mud sludge. The disadvantage is that only scandium can be recovered and not the other rare earths, because the other rare earths have a much weaker tendency to form soluble carbonato complexes.

The imidazolium ionic liquid 1-ethyl-3-methylimidazolium hydrogensulphate, [EMIM][HSO₄], has been used for high-temperature leaching of bauxite residue (Davris et al., 2014). At a temperature of 190 °C, recovery was between 60 and 70% for most of the rare-earth elements, although 100% recovery of lanthanum was possible under some conditions. All of the iron and titanium present was solubilised, as well as about 35% of aluminium. Substitution of the ethyl group by a longer alkyl chain had little effect on the leaching
efficiencies. By leaching with the carboxyl-functionalised ionic liquid betainium bis(trifluoromethylsulphonyl)imide, [Hbet][Tf₂N], a much higher selectivity for dissolution of the rare earths was achieved, even though the recovery percentages are lower than in [EMIM][HSO₄]. [Hbet][Tf₂N] is able to selectively dissolve rare-earth oxides, leaving behind aluminium and iron oxides (Nockemann et al., 2006).

Bioleaching experiments on bauxite residue have been carried out with the acid-producing fungus *Penicillium tricolor* to remove the rare earths and radionuclides (Qu and Lian, 2013). The authors directly isolated from bauxite residue this strain of fungi, which can excrete the maximum amounts of organic acids (oxalic acid and citric acid). They tested the bioleaching efficiency under various bioleaching processes and pulp densities. The highest leaching efficiencies were found for yttrium and the heavy rare earths, followed by scandium. The lowest efficiencies were found for the light rare earths. This leaching behaviour is similar to what is observed for mineral acids.

Most combined pyro/hydrometallurgical treatments of bauxite residue use a pyrometallurgical step to first recover iron from bauxite residue and to subsequently concentrate the rare earths in an oxide slag (Liu et al., 2009b; Logomerac, 1979a; Logomerac, 1979b; Loginova et al., 2013). The efforts made to extract iron from bauxite residue have been described above. The hydrometallurgical step consists of leaching the rare earths from the slag with a diluted mineral acid (Sargic and Logomerac, 1974). During reductive smelting of bauxite residue, 98–99% of the scandium content reports to the slag phase (Tanutrov et al., 2013). Scandium has been recovered from slag obtained by smelting of Chinese bauxite residue, by leaching with 30% H₂SO₄, followed by solvent extraction with bis(2-ethylhexyl)phosphoric acid in sec-octanol/kerosene diluent, and stripping with NaF solution from the loaded organic phase to obtain Na₃ScF₆ (Xu and Li, 1996) The combined use of pyrometallurgical and hydrometallurgical methods for recovery of metal values of bauxite
residue is called complex processing (Zazubin et al., 1967; Sargic and Logomerac, 1974; Loginova et al., 2013).

4. Mine tailings

The older beneficiation methods to separate rare-earth ores from gangue were not very efficient. Large losses occurred during the flotation process of ground rare-earth ores, which has led to the cumulative generation of large volumes of solid waste still containing significant concentrations of rare earths (Jordens et al., 2013). This powdery solid waste was often dumped in tailing ponds close to the mine or the beneficiation plant. The concentrations in these residues are so high that these tailings can be considered as genuine rare-earth deposits. For instance, the tailings of the Mountain Pass mine in California are considered as the second largest rare-earth deposit in the United States (the largest being the Mountain Pass mine itself). The Mountain Pass mine tailings still contain between 3 to 5% of rare-earth oxides. The recovery of rare earths from the huge polymetallic iron-niobium-REE deposit of Bayan Obo in Inner-Mongolia (China) is very inefficient. The deposit is mainly mined for its iron ore and only about 10% of the rare-earth content is recovered (Zhang et al., 2014). The situation is even worse for niobium, with virtually zero recovery of this metal. Concurrently, large volumes of mine tailings rich in rare earths are found near the Chinese rare-earth deposits or processing units (Xu and Peng, 2009). Also these “alternative” deposits represent a vast untapped resource of rare earths. The tailings in Baotou are receiving wide attention because they are polluting the environment with radioactive thorium-containing dust. More than 150 million of tonnes of tailings are being stored in a pond with an area of circa 12 km² close to the Yellow River in Baotou city (Yu et al., 2012). Rare-earth-tailings from Bayan Obo ore in China have been used for the production of glass ceramics (Zhao et al., 2010; Luo
Research is ongoing towards the concentration of the rare earths in the slag phase upon reductive smelting of the iron-rich Bayan Obo tailings. This work is discussed in section 6 on metallurgical slags. Carbochlorination is a promising method for the recovery of rare earths, thorium and niobium from the tailings of Bayan Obo ore (Yu et al., 2012). The tailings powders are mixed and heated in a stream of chlorine gas at 550 to 600 °C, transforming the metal compounds into chlorides. Some of these metal chlorides are volatile and can be removed from the reaction mixture by sublimation. Chloride salts of the rare earths and thorium are left behind in the solid residue, and can be subsequently recovered. Another approach to solubilise the rare earths and niobium from the Bayan Obo tailings is by sulphating roasting at 250 °C, followed by leaching with water at 60 °C (Zhang et al., 2014). Not only during the beneficiation process, but also during the cracking of rare-earth ores by acids or bases, significant amounts of rare earths are lost. Likewise, the historical processing of monazite has generated large volumes of rare-earth-rich tailings at Rhodia Solvay’s rare-earth processing plant at La Rochelle (France).

More efficient extraction of rare earths from old mine tailings would not only give an easy access to significant amounts of rare earths, but is also beneficial from an environmental point of view as this could be integrated in remediating mine sites through either in situ landfill mining or ex situ landfill mining (Jones et al., 2013; Jones et al., 2012). In the former case the metals are extracted in situ; in the latter case the residues are excavated and further processed to recover any critical metals and to, ideally, valorise the remaining mineral phases in other applications, in line with the zero-waste principle (see further).

Economically interesting concentrations of rare earths are found not only near rare-earth mines, but at tailing deposits of other mines as well. For instance, apatite associated with iron ores often contains significant amounts of rare-earth elements. The apatite-rich rocks on the tailings of the iron mines of the Mineville district, New York (USA), are a typical example
Reddish-brown apatite of this deposit contains between 5.8 and 20.6 wt% total rare earths, the green variety between 0.5 and 2.0 wt%, and the white and transparent varieties only trace amounts. Monazite and bastnäsite fill microfractures in apatite crystals. The tailings of the Pea Ridge iron ore mine, Washington County, Missouri (USA) contain considerable amounts of rare earths (Grauch et al., 2010; Long et al., 2010). The rare earths are present in the material from breccia pipes that cross-cut the magnetite-hematite ore body and its altered rhyolite host rock. The main rare-earth-bearing ore minerals are monazite and xenotime, that are present as fine grains within apatite. The apatite does not occur only in the breccia, but is also dispersed throughout the whole iron ore body. The apatite itself contains minor amounts of rare earths in its crystal structure. Besides the rare earths, the breccia contains significant concentrations of gold, silver and tin. The Pea Ridge deposit is a high-grade rare-earth deposit (on average 12 wt% rare-earth oxides, estimated total: 72,000 metric tonnes) and is relatively rich in heavy rare earths and yttrium. In Europa, apatite-iron ores are found in the Kiruna area in northern Sweden (Palsson et al., 2014). The main rare-earth minerals are apatite, monazite and allanite. Flotation tests have been carried out on the solid materials from the tailing pond in order to concentrate the rare-earth-containing minerals. Haine and Filippov explored the potential to recover rare earths from the waste generated by kaolin (China clay) extraction plants in Cornwall, UK (Dehaine and Filippov, 2014). The kaolin deposits are formed by weathering of granite rocks. In the micaceous waste fraction, the rare earths are concentrated in monazite grains, and the main rare earths are cerium, lanthanum and neodymium. Likewise, tailings of uranium mines often contain large concentrations of rare earths. For instance, tailings of uranium mines in Queensland (Australia), Kazakhstan and Kyrgyzstan are presently under investigation as possible new rare-earth “alternative deposits” (Golev et al., 2014). Furthermore, tailings of titanium mines could also offer possibilities for the extraction of rare earths (Jha et al., 2008). Ilmenite is
transformed into water-insoluble alkali titanate and water-soluble ferrite by alkali roasting (Sanchez-Segado et al., 2015; Jha et al., 2011; Lahiri and Jha, 2009). After roasting, the insoluble alkali titanate is separated from the rare-earth oxide mixture in colloidal form and water-soluble ferrite. Further leaching of alkali titanate is carried out with oxalic acid and ascorbic acid solutions, to remove the remaining Fe$^{2+}$ ions into the leachate and to precipitate a high-purity synthetic rutile containing more than 95% TiO$_2$. The mining tailings of tin placer deposits often contain considerable amounts of monazite and xenotime (Szamalek et al., 2013). Cassiterite is the main tin ore in these deposits.

On the other hand, an interesting source of scandium can be found in the residues of the processing of tungsten ores (Xu and Li, 1996; Natansohn et al., 1992; Gokhale and Bhat, 1967; Guo et al., 1988). If pressurised leaching with aqueous alkali is used to yield sodium wolframate from wolframite, scandium is left behind in the tungsten residue. Scandium can be recovered from this residue by leaching with HCl solution, followed by solvent extraction with bis(2-ethylhexyl)phosphoric acid (D2EPHA). Scandium can also be recovered from the residues generated during the processing of tin, nickel, tantalum and niobium ores (Wang et al., 2011b).

5. Coal ash and incinerator ash

Coal contains trace concentrations of many elements, including rare earths. These elements, with the exception of volatile elements such as mercury and cadmium, end up in the bottom and fly ashes of power plants after burning the coal for electricity production (Querol et al., 1995). The burning of coal leads to an enrichment of some metals in the ashes so that these ashes be considered as low-grade ores for different metals (Seredin and Finkelman, 2008; Seredin et al., 2013; Franus et al., 2015). Concentrations of more than 0.1% of rare-
Earth oxides are often found in coal ashes (Seredin and Dai, 2012). Ashes from coal of the Russian Far East contain even more than 1% of rare-earth oxides (Seredin, 1996). Given the fact that still huge amounts of coal are burnt worldwide – despite the negative environmental and climate impact of this energy source – coal ashes could be an important source of rare earths (Calus-Moszko and Bialecka, 2013). It has been recommended to use method similar to those used for the extraction of rare earths from weathered crust deposits (ion adsorption clays) (Blissett et al., 2014). Experiments to leach rare earths with dilute H\textsubscript{2}SO\textsubscript{4} from coal fly ash have been carried out (Kashiwakura et al., 2014). First, a fast dissolution of the rare earths on the surface of the ash particles was observed, followed by slow dissolution of the rare earths inside the particles. In a patented process, it has been proposed to leach the rare earths from coal ashes by dilute HNO\textsubscript{3}, followed by solvent extraction with TBP (Joshi et al., 2013). In another process, it is proposed to leach the rare earths with HCl from coal fly ash (Boudreault et al., 2013a).

Ashes from oil shales are another potential resource of rare earths (Yang et al., 2010). Extraction of rare earths from these ashes is a challenge due to the high iron and aluminium contents relative to the rare-earth concentrations. A conceptual flow sheet for the recovery of rare earths was proposed, consisting of four steps: (1) leaching with HCl; (2) removal of iron by solvent extraction with the trialkylamine N235; (3) selective precipitation of aluminium and rare earths as hydroxide and separation from impurities; (4) dissolution of the hydroxides in HNO\textsubscript{3}, followed by extraction of the rare earths by TBP to remove them from other metals dissolved in the solution.

The ashes of incineration plants contain a wide variety of metals. A study of the ashes of a Swiss incineration plant burning municipal solid waste showed that the concentrations of rare earths are very low and the extraction of these elements from the ashes would not be economically feasible (Morf et al., 2013). Analysis of the metal content of a Danish
incineration plant gave similar low concentrations of rare earths (Allegrini et al., 2014). The concentration of rare earths in Chinese medical waste incinerator ashes was found to be lower than the crustal abundance of the rare earths (Zhao et al., 2008). Research on the ashes of incineration plants show that landfills storing municipal waste are unlikely to become a resource of rare earths, although these landfills often contain high concentrations of base metals (Jones et al., 2013; Jones et al., 2012). This is not a surprise, given the fact that widespread use of rare-earth products (e.g. neodymium permanent magnets) in consumer goods is only a quite recent phenomenon. The situation could be different in the case of some specialised industrial landfills, for instance landfills storing lamp phosphor waste from End-of-Life fluorescent lamps.

6. Metallurgical slags

Efficient pyrometallurgical processes have been developed for the recovery of metal values from electronic scrap, spent automobile exhaust catalysts and spent industrial catalysts. In its Hoboken site near Antwerp (Belgium) Umicore operates an integrated smelter and refining plant for recovering metals from Waste Electrical and Electronic Equipment (WEEE) (Felix and Vanriet, 1994; Meskers et al., 2009; Vanbellen and Chintinne, 2006). Recovered metals include: silver, gold, indium, bismuth, tin, selenium, tellurium, antimony, arsenic, the platinum-group metals (platinum, palladium, ruthenium, rhodium, iridium) and the base metals copper, nickel and lead. However, the employed metallurgical flow sheets have not been developed for the recovery of rare earths. Due to the high affinity of the rare-earth metals for oxygen, these elements end up in diluted (oxidic) form in the slag of the so-called “copper smelter” and finally in the slag of the lead blast furnace. Moreover, the rare-earth content of these slags consists largely of cerium, which is not a critical metal. In fact, there is
an oversupply of cerium, due to the large natural abundance of cerium and its relatively low demand (Binnemans et al., 2013b). Other companies such as Aurubis and Boliden use pyrometallurgical processes comparable to that of Umicore for the multi-metal recycling from electronic scrap, but none of cases the rare earths are recovered from the slag phases (Boliden, 2014; Aurubis, 2014).

Concurrently, pyrometallurgical flow sheets for the recycling of valuable metals from batteries can produce a slag relatively rich in rare earths. In 2011, Rhodia (Solvay) and Umicore announced that they had jointly developed a process for recycling of rare earths from nickel metal hydride rechargeable batteries (Anonymous, 2011; Rhodia, 2011). Although the details of the process have not been disclosed, it is based on Umicore’s patented Ultra High Temperature (UHT) smelting technology (Cheret and Santen, 2007). An industrial-scale pilot plant is operational in Hoboken since September 2011 and this pilot plant has an initial annual capacity of 7000 tonnes, corresponding to approximately 150000 (hybrid) electric vehicle batteries or 250 million mobile phone batteries (Umicore, 2011; Umicore, 2009). Not only nickel metal hydride batteries, but also lithium-ion batteries could be recycled in this facility. At this moment, the process has been optimised for End-of-Life portable nickel metal hydride batteries. The batteries are fed in a vertical shaft furnace, together with a small amount of coke and a slag former (Cheret and Santen, 2007). At the bottom of the shaft furnace, oxygen-enriched air is injected into the furnace. The process requires relatively little external energy input, because the combustion of both the plastic casing of the batteries and the organic electrolytes releases large amounts of energy. The metals are converted into a Ni-Co-Cu-Fe alloy and a slag. The slag consists mainly of oxides of calcium, aluminium, silicon, and iron, and also contains lithium and rare earths (Verhaeghe et al., 2011). These oxide slags can be processed to recover lithium and to produce rare-earth concentrates that are subsequently used
as a feed in the rare-earth separation plant of Solvay (formerly Rhodia) in La Rochelle, France.

In comparison to natural ores, slags have the advantage that the formation of mineral phases during cooling can be influenced by use of different slag formers, cooling rates and furnace conditions. This processing of slags is defined as *hot stage slag engineering* (Durinck et al., 2008). The slag composition and microstructure can be changed by adjusting the primary pyrometallurgical process directly or immediately after separation of the slag from the molten metal, while the slag is still at high temperature. If a metal can be enriched in a certain mineralogical slag phase embedded in a matrix of other minerals by choosing a suitable slag composition and cooling trajectory, then a separation of this mineral phase might be possible by conventional ore processing methods. The metal could be much more effectively extracted from the resulting mineral concentrate than by processing the complete slag (Elwert et al., 2014). To present, interest from academia for research on the recovery of rare earths from metallurgical slags has been limited. Müller and Friedrich investigated the recycling of nickel metal hydride batteries and measured the rare-earth content in CaO–SiO$_2$–CaF$_2$ based slag both in laboratory tests and pilot scale trials (Muller and Friedrich, 2006; Muller and Friedrich, 2004). As expected, the rare-earth metals collected in the slag phase and a rare-earth-rich slag (50–60 wt% rare earths) were obtained. In a pyrometallurgical process for recycling of NiMH batteries, the batteries were first dismantled (Tang et al., 2013). In a next step, the electrodes and the polymer were heated to 600 °C to combust the organic components. After separation of nickel metal from the black powder that contained NiO, CoO and the rare earths, the powder was further treated by a slagging process using a calcium silicate slag. Nickel and cobalt were recovered as an iron-copper-cobalt-nickel alloy and the rare earths reported to the slag phase. The affinity of neodymium and dysprosium to phosphate-containing phases was investigated for the Al$_2$O$_3$–CaO–MgO–P$_2$O$_5$–SiO$_2$ slag
In the phosphate-free $\text{Al}_2\text{O}_3$–$\text{CaO}$–$\text{MgO}$–$\text{SiO}_2$ slag system, dysprosium and neodymium were more or less randomly distributed throughout the different silicate phases, although local enrichments of up to 15 wt% of rare-earth oxides were found. Addition of $\text{P}_2\text{O}_5$ led to the formation of a britholite-like silico-phosphate, with up to 57 wt% of rare-earth oxides. This phase formed at an early stage of the cooling process and it is assumed that it scavenged the rare earths before the silicate phases were crystallising. Even at a content of $\text{P}_2\text{O}_5$ as low as 4.57%, the concentration of rare-earth oxides in the silicate matrix phase was already very strongly reduced. This shows that the rare earths in slags have a strong affinity for phosphate-containing phases. A rare-earth-rich slag has been obtained by a pyrometallurgical process based on carbon-bearing pellet reduction and melting technology to recover iron from complex ores of Bayan Obo in China (Ding et al., 2012; Ding et al., 2013). The main mineral phases of the ore were iron oxides, bastnäsite-(Ce) and fluorite. During the smelting process, nearly all the rare earths reported to the slag phase and the resulting slag contained 14% of rare-earth oxides. The slag was remelted at 1400 °C and the melt was cooled using three types of cooling conditions (water quenching, air cooling, and furnace cooling) to investigate the influence of cooling on the microstructure and phase characteristics of the slag. The rare-earth elements were concentrated in a cerium calcium silicate phase and could be leached from the slag by 4 M HCl, leaving behind $\text{CaF}_2$ and $\text{ThO}_2$. The recovery of the rare earths was more than 97% in all cases; the lowest recovery was observed for the water-quenched slag, while the highest recoveries (> 99%) were observed for the air-cooled and the furnace-cooled slag. The recovery of the rare earths from the original slag was in between the two extremes. One could wonder whether remelting of the slag is justified from an economic point of view, since it resulted in only 1% extra leaching of the rare earths. Although the rare earths report to a cerium calcium silicate slag phase, europium and
scandium do not. These rare earths are concentrated in rare-earth-containing barium fluorophlogopite phase (Zheng and Lin, 1994). This rare-earth-rich phase could be collected from crushed and powdered slag by a flotation process. Another type of rare-earth-rich slag was obtained by the reductive smelting of a magnetite ore containing monazite (Kim et al., 2014). The authors investigated the kinetics of lanthanum leaching from this slag by 0.3 M H$_2$SO$_4$. Scandium reports to the slag phase during the reduction of tungsten residue in an electric arc furnace (Xu and Li, 1996).

Additionally, some information on the presence of rare-earth oxides in flux powders used in the continuous casting of steels is available, as well as on slag resulting from the smelting of rare-earth-containing iron ores (Zheng and Lin, 1994; Anacleto et al., 1993; Derevyankin et al., 1981a; Derevyankin et al., 1981b; Fang et al., 2001; Jia et al., 1995; Lin and Zheng, 1997; Yang et al., 2011). A wealth of information on the formation of rare-earth-containing minerals in magmas is available in the geological community, although geological conditions and time scales may not be very relevant for rare-earth-recovery operations. The available thermodynamic data for rare earths elements in FactSage, the most widely used thermodynamic calculation system for high temperature oxide systems, is fairly limited (Factsage, 2013). In order to make the recovery of rare earths from metallurgical slags more efficient, the problem of dilution of the rare earths in oxide slags has to be solved. Once the rare earths have been concentrated into solid rare-earth-rich phases, they can subsequently be removed much more easily from these phases by acid leaching.

7. Waste water

Waste water could also be a source for rare earths, but the potential of this waste stream for the recovery of rare earths is largely unexplored. In the first instance, rare earths could be
recovered from waste water produced during the extraction and separation of rare earths. Acid mine drainage (AMD) often contains considerable concentrations of rare earths (Merten and Buchel, 2004; Protano and Riccobono, 2002; Delgado et al., 2012). AMD is the outflow of acidic water from (old) metal mines or coal mines, and it is often heavily contaminated by metals (Akcil and Koldas, 2006). Some studies report on the possibility to recover uranium and other metals from AMD by ion-exchange resins or by biosorption, but limited information is available on the possibility to recover rare earths (Roig et al., 1997; Geremias et al., 2003; Fu and Wang, 2011). Whereas solvent extraction is the preferred method for recovery of rare-earth ions from concentrated aqueous waste streams and pregnant leach solutions, this method is not recommended for removal of rare-earth ions from diluted aqueous waste streams, because of the unavoidable contamination of the aqueous phase by organic solvents. Therefore, ion-exchange resins and chelating resins are to be preferred.

Yantasee and co-workers have investigated the possibility to bind rare-earth ions by functional groups on the surface of a mesoporous silica support (Yantasee et al., 2009). It has been proposed to use these materials for recovery of rare earths from acidic industrial waste water streams and mining effluents. Nanoporous silica surface modified with 3,4-hydroxypyridinone was tested for the recovery of cerium and europium from aqueous solutions (Johnson et al., 2012). The mechanism of the uptake of rare earths by chitosan functionalised with EDTA or DTPA has been studied (Roosen and Binnemans, 2014; Inoue, 2000; Roosen et al., 2014). Functionalised magnetic nanoparticles are efficient for the capture of rare-earth elements from water, and such nanoparticles can be easily retrieved from the solution by a permanent magnet (Dupont et al., 2014a; Dupont et al., 2014b; Legaria et al., 2015). Also functionalized non-magnetic nanoparticles such as SiO₂ have been used for uptake of rare-earth ions from aqueous solutions, but recovery of the metals is less straightforward than with magnetic nanoparticles (Dupont et al., 2014a; Topel et al., 2014).
Although the biosorption of precious metals and uranium by algae, fungi, bacteria and yeast has been widely explored, up to now relatively little attention has been paid to the biosorption of rare earths (Das and Das, 2013; Philip et al., 2000; Anagnostopoulos and Symeopoulos, 2008; Tsuruta, 2007; Moriwaki and Yamamoto, 2013; Andres et al., 2003). The fungus *Ganoderma lucidum* has been tested for the uptake of rare earths and thorium from waste water from the processing of monazite (Muraleedharan et al., 1994). By-pass cement dust (BCD) has been tested as a cheap adsorbent for rare-earth ions from aqueous solutions (Ali et al., 2011). BCD is a by-product of the manufacture of Portland cement and is generated during the calcination process in the kiln. This fine-grained material mainly consists of calcium oxide. Supported flowerlike nano-Mg(OH)\(_2\) exhibits a large specific surface area and high extraction ability for rare earths (Li et al., 2013). Pilot-scale experiments showed that this material is promising to recycle rare earths from industrial waste water.

8. **Towards zero-waste valorisation**

8.1. **Concept of zero-waste valorisation**

In this paper we have used the term “valorisation” to refer to the economically viable utilisation of a material stream which was hitherto considered as a non-valuable waste stream. The term “valorisation” differs from the official meaning of the word in UK English where it is described as the increase in the value of capital assets through the application of value-forming labour in production. However, as the term “valorisation” has been used for many years by a multitude researchers in the context of “slag valorisation” (Jones et al., 2009) or “waste valorisation”, we also employ it in this meaning in the present review paper. Hence, valorisation of industrial process residues implies that a certain value is created in a cost-effective way.
When considering the valorisation of rare-earth containing secondary resources, one can take a straightforward metal-centric approach in which the most valuable rare-earths are extracted from the residue. As these metals form only a minor fraction of the overall residue the chosen metal-centric processing method is only delivering a suboptimal solution, in which the largest fraction of the residue is not catered for, often implying that more costly and environmentally-unfavourable stockpiling or landfilling is required. It is clear that this metal-centric approach is not in line with the whole-system approach that is conveyed by the zero-waste philosophy. The latter strives to eliminate waste at source and at all points down the supply chain, thereby rejecting one-way linear resource-use in favour of a “closed-loop” circular system (Curran and Williams, 2012). In the zero-waste vision, “waste” is considered as the symbol of an inefficient society where resources are badly allocated (Zaman and Lehmann, 2013; Zaman, 2014). Zero-waste, therefore, envisions that all industrial inputs are somehow being used in final products or converted into value-added feedstocks for other industries or processes. This vision is completely in-line with the industrial symbiosis approach, in which distinct industries are re-organised into clusters in such a way that each company’s by-products match the input requirements of another industry, whereby the integrated whole produces (near) zero wastes (Curran and Williams, 2012). As pointed out by ZWIA, products and processes need to be designed and managed to avoid and eliminate waste and to recover all resources from a given waste stream (ZWIA, 2004). For industry greater competitiveness can be obtained, representing a continuation of the inevitable drive towards (resource) efficiency (Curran and Williams, 2012). Translated to the domain of zero-waste valorisation of metal-containing industrial process residues, this implies that a rare-earth metal-centric recovery approach needs to be replaced by a product-centric, whole system approach, in which (almost) all components of the residue need to be turned into value, i.e. “valorised”. The rejection of a metal-centric in favour of a product-centric approach has
received ample attention in the most recent UNEP report *Metal Recycling – Opportunities, Limits, Infrastructure* (Reuter et al., 2013). Although the concept was mainly developed for the integrated valorisation of critical and technology metal containing End-of-Life products (cf. Waste Electrical and Electronic Waste, WEEE), it is perfectly fit for secondary resources as well. In the context of End-of-Life products, it has been pointed out that the product-centric view considers the complex metallurgy of all elements at the same time, optimising both the metallurgy and recycling infrastructure in order to minimise losses, but also addressing the related issues, such as liberation, sorting and consumer recycling (Reuter and van Schaik, 2013). The work of Reuter and colleagues demonstrates that zero-waste valorisation will require more than the development of technological solutions, as also pointed out by Klauber et al. (Klauber et al., 2011). In the case of industrial process residues, the “secondary resource” should thus be regarded as a polymetallic raw material product, also hosting a number of potentially valuable minerals. Zero-waste, product-centric valorisation, therefore, means that tailored, integrated flow sheets need to be designed to recover both critical and base metals, while simultaneously finding solutions for the residual mineral matrix as well. For instance, in the case of stocks of previously produced bauxite residue, this zero-waste vision implies that all valuable metals (including the lanthanides and scandium) etc. need to be recovered, while valorising the mineral matrices into, for example, innovative construction materials. In the case of future bauxite residues or phosphogypsum, an even more ideal, whole-system scenario is to redesign the overall flow sheet upstream so that different residues are generated, which can find better valorisation options further downstream the valorisation chain. The zero-waste valorisation concept is now discussed in more detail for phosphogypsum (section 8.2) and bauxite residue (section 8.3).
8.2. Towards zero-waste valorisation of phosphogypsum

Phosphogypsum could be used as a resource for the manufacturing of cement, plasters and gypsum construction materials, but the high contents of fluorides, phosphates and water-soluble sodium compounds turn phosphogypsum into a raw material that is of an inferior quality compared to natural gypsum or anhydrite for these applications. Also organic matter can cause problems. After or during the recovery of the metal values, phosphogypsum needs to be treated before it can be used in the construction industry. Lokshin and co-workers described a zero-waste valorisation approach for the processing of phosphogypsum, which includes purification of phosphogypsum on dumps by percolation leaching with dilute H$_2$SO$_4$ solutions to remove these fluorides, phosphates and water-soluble sodium compounds of sodium, with simultaneous dissolution of rare-earth sulphates (Lokshin et al., 2013). Indian phosphogypsum was purified by treating it with a 10–20 % aqueous ammonia solution, followed by washing with water (Singh et al., 1993). The purified phosphogypsum was tested on a lab scale for cement manufacturing and the produced Portland cement was found to have properties similar to cement produced from the mineral gypsum.

Another approach is to purify phosphogypsum by treating it with a 3–4 wt% citric acid solution, followed by washing with water (Singh, 2002). This treatment transforms phosphate and fluoride impurities in water-soluble citrates, aluminates and iron compounds. The residue can be used for the production of Portland cement or gypsum plaster. The Belgian company Prayon SA has recently developed an integrated process to recover rare earths from phosphogypsum by leaching with H$_2$SO$_4$, yielding a pure, white residue that can be valorised as a construction material (Germeau et al., 2013).

One particular point of attention for the zero-waste valorisation of phosphogypsum is the fact that phosphate rocks are slightly radioactive due to the presence of uranium and its decay
products. The contribution of thorium to the radioactivity of phosphate rock is low, due to the very low thorium concentrations in phosphate rock. Upon the acid attack of phosphate rock, the radioactive equilibrium existing in the phosphate rock is disturbed and a redistribution of the radionuclides occurs. Most of the uranium and thorium report to the phosphoric acid and subsequently finish up in the phosphate fertiliser, while radium will be incorporated in the phosphogypsum since it shows chemical similarities to calcium (Papastefanou et al., 2006; Hull and Burnett, 1996). Also polonium will end up in the phosphogypsum. Four radionuclides in phosphogypsum are of concern: $^{226}\text{Ra}$, $^{231}\text{Pa}$, $^{210}\text{Pb}$ and $^{210}\text{Po}$ (Vanderheijde et al., 1990). Most attention has been paid to the radium content of phosphogypsum.

Phosphogypsum may contain specific activities of $^{226}\text{Ra}$ between one and two orders of magnitude higher than those of natural gypsum, depending on the uranium content of the ore (Menzel, 1968). Due to the enrichment of radionuclides, phosphogypsum is classified as a technologically enhanced naturally occurring radioactive material (TENORM) (Shakhashiro et al., 2010; Tayibi et al., 2009; Gezer et al., 2012). The decay of $^{226}\text{Ra}$ in the phosphogypsum will result in the formation and reappearance of daughter nuclides and radioactive equilibrium will be re-established. Radiological measurements on phosphogypsum of Thessaloniki (Greece) showed that the activity of $^{226}\text{Ra}$ varied between 261 to 688 Bq kg$^{-1}$, with a mean value of 508 Bq kg$^{-1}$ (Papastefanou et al., 2006). It must be mentioned that also part of the uranium originally present in the phosphate rock ends up in the phosphogypsum (with the concentration depending on the type of phosphate rock and the chemical process used), so that also daughter elements of $^{238}\text{U}$ can be found in the phosphogypsum (Rutherford et al., 1994). The possible inhalation of the radon ($^{222}\text{Rn}$) formed by the decay of $^{226}\text{Ra}$ is considered as a health concern (Haridasan et al., 2002). Several authors have investigated the activities of radionuclides in phosphogypsum at different geographic locations. Hence, more information is available on radioactivity issues of phosphogypsum compared to bauxite residue (red mud).
(see section 4) (Rutherford et al., 1994; Mazzilli et al., 2000; Saueia et al., 2005; Azouazi et al., 2001; Hull and Burnett, 1996; Zielinski et al., 2011; Papastefanou et al., 2006; Paridaens and Vanmarcke, 2008; Paridaens and Vanmarcke, 2001; Vanderheijde et al., 1990). The presence of radionuclides in phosphogypsum is of concern when phosphogypsum is valorised, both at the stage of rare earths recovery and during the transformation of phosphogypsum into cement, plaster or other building materials. This implies that safety measures are required when working with phosphogypsum in order to avoid uncontrolled release of the radionuclides and to avoid the risk of exposure of the workers to those radionuclides. In the conversion of phosphogypsum to ammonium sulphate by reaction with ammonium carbonate, all the radionuclides that were originally present in the phosphogypsum ($^{226}$Ra, $^{210}$Pb and $^{210}$Po) will report to the calcium carbonate by-product (Burnett et al., 1996). As a result, the ammonium sulphate fertiliser will have a very low radioactivity. However, it must be taken into account that the concentration of radionuclides in the calcium carbonate has consequences if this calcium carbonate is further treated for recovery of rare earths.

Recovery of rare earths from phosphogypsum could be combined with removal of radionuclides, as part of remediation or recultivation of old phosphogypsum stacks to leave the landfills in a safer condition. If the phosphogypsum is not of a high enough quality for use in plaster or similar applications, excavation and ex situ leaching of the radionuclides and rare earths is not a feasible option. In such cases, in situ leaching could be option. This approach implies injecting lixiviants in the phosphogypsum stack, followed by collection of the leachates and recovery of the dissolved metals. No studies on in situ leaching of phosphogypsum stacks have been done yet, although in situ leaching is commonly used for recovery of metals from low grades ores, especially from low grade uranium ores (Gupta and Singh, 2003). However, in situ leaching can only be applied if precautions are taken to avoid contamination of ground water.
8.3. Towards zero-waste valorisation of bauxite residue

Zero-waste valorisation of bauxite residue implies not only removal of the rare earths and other metal values, but also finding applications for the metal-lean residues. Unlike some other high volume industrial wastes such as fly ash and metallurgical slag, bauxite residue currently finds no major industrial applications, besides minor use in cement and ceramic production (Pontikes and Angelopoulos, 2013; Liu and Zhang, 2011). Many researchers have already looked at the valorisation of bauxite residue, apart from its valorisation in the construction industry (Paramguru et al., 2005; Sushil and Batra, 2008; Liu et al., 2009b; Bhatnagar et al., 2011; Klauber et al., 2011; Liu and Wu, 2012). The removal of rare earths can be done in a preprocessing step consisting of leaching the rare earths from bauxite residue, followed by further processing of the remaining residue for recovery of the other metal metals (Piga et al., 1993). However, other processing schemes are possible. For instance, the rare earths could be removed from the slag that are formed during the removal of iron as pig iron.

Bauxite residue has a high (base) metal content and extraction of metals from bauxite residue might be economically feasible. Iron oxide is a main constituent of bauxite residue (up to 60% of the mass) and causes its red colour. In view of the rather large content of iron oxide, attempts have been made in the past to use bauxite residue as a source of iron. Iron recovery studies were initiated as early as in the 1950s and may be classified into two major approaches, i.e. smelting and solid state reduction (Hammond et al., 2013). In the former, bauxite residue is treated in a blast furnace in the presence of a reducing agent where the iron oxides are reduced, generating pig iron and a titanium-rich slag (also containing the rare earths) (Ercag and Apak, 1997; Kumar et al., 2006; Xenidis et al., 2011; Logomerac, 1979a;
Raspopov et al., 2013). In the solid-state reduction, bauxite residue is mixed with a reducing agent or contacted with a reducing gas at certain temperature, resulting in the formation of metallic iron or the conversion of ferric iron to magnetite, which can be used in a next step as feed for a conventional blast furnace (Xenidis et al., 2009; Samouhos et al., 2013; Chun et al., 2014; Liu et al., 2009a). So far, these iron recovery processes have not been commercially successful. The large sodium content of bauxite residue prohibits the use of blast furnaces. The high temperatures lead to the evaporation of sodium oxide in the lower (hotter) regions of the blast furnace and its re-deposition in the colder regions, where it attacks the ceramic refractory bricks and also leads to the formation of so-called sodium nests. These sodium nests are hard outgrowths that obstruct the free flow of the charge inside the furnace. Moreover, bauxite residue contains a lot of water, which would have to be removed first, thus representing a high cost if fossil fuels are used for drying. The solid-state reduction process, resulting into a beneficiated stream of magnetite, appears feasible but most likely economically not viable.

A concurrent valorisation of the iron and iron-lean residue, designed for bauxite residue, would possibly address the above. This implies a mentality shift from the concept of introducing bauxite residue into an existing process, to the concept where a process is developed specifically for bauxite residue. Obviously, the commitment of resources in the second trajectory is much higher, although it could be the way forward. An example of that approach is the ENEXAL bauxite residue treatment process (Balomenos et al., 2014; Balomenos et al., 2013). Through electric arc furnace carbothermic smelting, bauxite residue is fully converted into two marketable products: pig iron and mineral wool. Pig iron is used in the secondary steel industry as a steel scrap substitute, while the mineral wool can applied for the production of thermo-acoustic insulating products for the construction industry. No solid or liquid waste products are generated in this process, and thus in conjunction with the
alumina refinery plant, zero-waste valorisation of the bauxite ore can be achieved. This novel process has been applied for more than a year in Aluminium of Greece’s industrial scale pilot plant. So recovery of rare earths is not done in the ENEXAL process, but new process flow sheets with recovery of rare earths could be developed.

Besides iron, bauxite residue also contains economically relevant concentrations of aluminium and titanium. High silica contents in bauxites result in reduced Al$_2$O$_3$ recovery in the Bayer process since a part of the dissolved Al$_2$O$_3$ will precipitate in the form of complex aluminosilicates, e.g. cancrinite Na$_6$Ca$_2$[Al$_6$Si$_6$O$_{24}$](CO$_3$)$_2$ (Smith, 2009). Re-feeding these residues in the Bayer process does not result in recovering significant amounts of alumina. In conjunction with losses during the digestion and settling in the Bayer process, bauxite residue contains as a result up to 20 wt% Al$_2$O$_3$ which is a serious loss for the industry. To achieve Al$_2$O$_3$ recovery from high-silica bauxite residues, several processes have been developed in the past. The “MgO–Na$_2$CO$_3$ sinter process” involves adding MgO to bauxite residue and sintering at 900–1100 °C (Meher et al., 2011a). A variant of this method is the “BaO-Na$_2$CO$_3$ sinter process”, which also achieves high alumina recovery as dissolved sodium aluminates and low silicon dissolution (Meher et al., 2011b). An alternative to the sintering process is the Orbite process which uses high temperature HCl leaching to dissolve aluminium and iron as chlorides (Boudreault et al., 2013b). Efforts have also been made to recover titanium from bauxite residue by leaching with H$_2$SO$_4$ (Gatzini-Leonardou et al., 2008; Sayan and Bayramoglu, 2004). The recovery of metal values from bauxite residue has been reviewed recently (Liu and Naidu, 2014).

In fact, the work on alumina extraction should be seen in a wider scope and be coupled also with the iron removal as a first step. In this approach, a great amount of work already done in the early 1960s by the US Bureau of Mines on clays, could be now extrapolated both on bauxite residue and iron-lean slags (e.g. from the ENEXAL process) (Balomenos et al.,
The processes investigated on clays were based on HNO₃, sulphurous acid, HCl–isopropyl ether extraction, lime-soda with wet or dry grind option, H₂SO₄–electrolytic iron removal, H₂SO₄–chemical iron removal and potassium alum (Peters et al., 1967). The most promising acid processes were those based on HCl and HNO₃, with the separation of dissolved iron and aluminium salts, recovery of acid and crystallisation of an aluminium salt of sufficient purity to yield reduction-grade alumina identified as the major challenges to overcome (Goldberg, 1970).

Considering that bauxite residue contains small quantities of radionuclides (mainly ²³²-Th, ²³⁸-U, ²²⁶-Ra), bauxite residue can be classified as Technologically Enhanced Naturally Occurring Radioactive Material (TENORM) and falls under the legislation for Naturally Occurring Radioactive Materials (NORM) (Klauber et al., 2011; O'Connor et al., 2013; Qin and Wu, 2011; Gu et al., 2012; Somlai et al., 2008). These legal constraints have to be taken into account when bauxite residues are used in building materials. One could think of developing a process to leach out the radionuclides together with the rare earths, so the resulting bauxite residue would have very low radiation levels. However, this implies that downstream processing of the rare-earth-containing leachates should also include a step to safely dispose of the radionuclides present in these solutions.

9. Conclusions and outlook

In general, the concentrations of rare earths in industrial waste streams are low, typically less than 1% of rare-earth oxides. However, since the available waste volumes are huge, the total amount of rare earths in these waste fractions is very substantial, so that they can represent an interesting alternative source of rare earths, complementing rare earths obtained through primary mining and recycling of End-of-Life product flows. The recovery of rare
earths and other valuable elements can be part of a remediation process of historic landfills of industrial wastes. Recovery of valuable metals from industrial waste streams must be integrated in larger recycling schemes that recover most (base) metals and also find applications for the residues left after removal of the metals. The recovery of metals must thus be part of zero-waste valorisation schemes of industrial waste streams. Whereas radioactivity is not an issue in case of recycling of rare earths from End-of-Life consumer goods, it can be a serious issue in the case of processing industrial waste streams such as bauxite residue (red mud) or phosphogypsum. From an economic point of view, the most interesting industrial waste stream is bauxite residue because of its high concentrations of scandium. Extraction of scandium from bauxite residue can be an economically feasible industrial process, especially if bauxite residues with high scandium content can be processed, for instance Greek or Jamaican bauxite residues. Phosphogypsum can be a potential resource of light rare earths, but it is recommended to integrate the production of phosphoric acid and the recovery of the rare earths into one process, so that the rare earths are not incorporated in the phosphogypsum. It is a challenge to develop economically feasible and environmentally-friendly “new metallurgical systems” for the recovery of rare earths from industrial waste streams containing low concentrations of rare earths. However, as (heavy) rare earth prices tend to increase, recovery flow sheets for industrial process residues will become more attractive. By also finding medium to high value outlets for the other residues arising during these flow sheets, the overall business case of such flow sheets can be further improved. To provide more resilience to the rare-earth and raw materials industry, innovative fundamental and applied research in this field should, therefore, be of strategic interest, targeting comprehensive “new metallurgical systems”. These research activities can provide medium to long term answers for a likely situation in the future when the demand for (heavy) rare earths exceeds the available supply. Concerted efforts will be required as much of the reported
research data have been obtained on a trial-and-error basis, rather than on the basis of a rational design. Presently, only limited information is available with respect to the mineralogy of the different rare-earth rich phases in bauxite residue, phosphogypsum and metallurgical slag. Knowledge of these phases could aid the development of new more efficient leaching processes. There is also a need for methods that allow efficient recovery of rare-earth ions from dilute aqueous solutions, not only from waste water streams, but especially from the dilute leachates. Due to the fact that the concentrations of rare earths in industrial waste residues are low compared to primary rare-earth ores and reclaimed End-of-Life consumer goods (WEEE), tailored, zero-waste processes dedicated to the recovery of rare earths from these dilute waste streams must be developed.

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References


Factsage, 2013.


International Aluminium Institute, 2013. Bauxite Residue Management: Best Practice.


Richardson, J., 2013. Up to 2,500 per cent higher concentration levels in Ja's red mud: "God-blessed dirt". Jamaica Observer 13 February 2013.


