



# 2008201945 02 May 2008 2008201945 02 May 2008

# **ABSTRACT**

**A process for recovering rare earth elements from a composite ore containing rare earth elements that includes a monazite group mineral and an apatite mineral, includes pre-leaching the composite ore with an acid so as to substantially dissolve the 5 apatite mineral into the leach liquor and precipitating rare earth elements from the preleach liquor. The residue of the pre-leaching step is subjected to an acid bake treatment, followed by a water leach, to produce a water leach liquor rich in rare earth elements. Impurities including thorium and iron are separated from the water leach liquor by introducing a neutralising additive to the water leach liquor rich in rare earth 0 elements, and rare earth elements are precipitated from the post-neutralisation liquor.**

P/00/011 **Regulation 3 2**

**AUSTRALIA** 

*Patents Act 1990*

# **COMPLETE SPECIFICATION STANDARD PATENT**

Invention Title: **Recovery of rare earth elements**

**The following statement is a full description of this Invention, including the best method of performing it known to us:**

# **Recovery of Rare Earth Elements**

# **Field of the invention**

**This invention relates generally to the recovery of rare earth elements from ores containing the elements. More particularly, the invention is concerned with recovering 5 rare earth elements from a composite ore containing rare earth elements that includes a monazite group mineral and an apatite mineral.**

## **Background of the invention**

**Rare earth elements are of increasing economic importance as the highly specialised properties of individual elements find applications in a wide range of pervasive products.**

- **0 For example, lanthanum is a component of catalysts employed in oil refineries, neodymium is a component of solid state lasers, neodymium magnets are widely employed in headphones, hard disk drives, and DVD and CD-ROM drives, and europium and terbium are dopants in primary colour phosphors for plasma and LCD screens.**
- **5 At present, the principal source of most rare earth elements is China, and the principal deposits are in the Sichuan and Baotou regions.**

**With limited prospects for expanded production, and diversion of an increasing proportion to domestic customers, China is unlikely to be able to meet the expected increase in world demand for rare earth elements over the coming years. Since the gap**

**20 will not be met by traditional sources outside China, such as monazite deposits for certain of the elements, there is a growing requirement to tap non-traditional ores containing rare earth elements. One of these is a composite ore containing rare earth elements that includes a monazite group mineral and an apatite mineral. Apatite is a phosphate of calcium, while a "monazite group mineral" is a composite monoclinic 25 phosphate of rare earth elements and several other substituents.**

**When a composite ore of this structure has been subjected to conventional or other known processing regimes, it has proven difficult to convert the ore to an economically**

**useful deposit. High thorium content in both principal components of the ore have proven a hurdle to economic conversion of the phosphate base of the apatite to a "clean" fertiliser product, while the uranium content associated with the thorium cannot be economically extracted in sufficiently pure form.**

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**Primary treatments that were attempted included a nitric acid leach, a hydrochloric acid leach, a sulphuric acid bake/water leach and a caustic crack. It was found that direct treatment of the ore resulted in either low rare earth recovery, or poor leach selectivity for rare earths over calcium and phosphorous.**

**The present invention aims to provide a process for recovering rare earth elements from composite ores containing rare earth elements that include a monazite group mineral and an apatite mineral, preferably in a manner that also permits economic extraction of the phosphate values of the ore and removal of thorium to an extent that <sup>1</sup>5 can meet end-product specifications.**

# **Disclosure of the invention**

**The invention provides, in a first aspect, a process for recovering rare earth elements from a composite ore containing rare earth elements that includes a monazite group '0 mineral and an apatite mineral, including:**

**pre-leaching the composite ore with an acid so as to substantially dissolve the apatite mineral into a pre-leach liquor;**

**25 precipitating rare earth elements from the pre-leach liquor;**

**subjecting the residue of the pre-leaching step to an acid bake treatment, followed by a water leach, to produce a water leach liquor rich in rare earth elements;**

**30 separating impurities including thorium and iron from the water leach liquor by introducing a neutralising additive to said water leach liquor rich in rare earth elements; and**

**precipitating rare earth elements from a post-neutralisation liquor.**

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**It is found that the acid pre-leach is effective in achieving sufficient dissolution of the calcium phosphate lattice of the apatite to allow rare earth elements to report to both the leach liquor and the solid residue.**

**The acid employed for the pre-leach is preferably nitric or hydrochloric acid, more preferably the latter.**

**Preferably, rare earth elements are precipitated from the pre-leach liquor as <sup>1</sup>0 phosphates.**

**Precipitation of rare earth elements as phosphates from the pre-leach liquor is conveniently by addition of ammonia or hydrated lime. Advantageously, precipitation conditions are selected to recover, with the rare earth elements and in a form 15 separable subsequently, a major proportion of thorium present in the pre-leach liquor from the composite ore.**

**Preferably, especially where the apatite is a fluorapatite, a fluoride component of the pre-leach liquor is removed, e.g. by addition of a sodium salt to cause precipitation of !0 Na2SiF6. Silicon is typically sufficiently present in the composite ore to allow this precipitation. The sodium salt conveniently matches the acid employed for the preleach,** for **example** NaCI in the case of a hydrochloric acid pre-leach or NaNO<sub>3</sub> with a **nitric acid pre-leach. The sodium salt is typically required to stoichiometric excess, e.g. 7 to 8 times. Recovery of the fluoride component is substantially more effective 25 when the pre-leach acid was hydrochloric.**

**The acid bake is preferably carried out in a rotary kiln, and is preferably effected for between <sup>1</sup> and 3 hours, most preferably 1.5 to 2.5 hours.**

**30 The acid bake treatment is preferably a sulphuric acid bake, and typically at a temperature in the range 200-280°C, more preferably 210-250°C, most preferably about 230°C.**

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**Preferably, the thorium and iron separation is effected by neutralisation with a suitable alkali or alkaline earth additive selected to minimise rare earth co-precipitation or absorption. A calcium compound is thought to be unsuitable for the latter purpose. A magnesium compound such as magnesium oxide or magnesium carbonate is 5 satisfactory and may be preferred.**

**The pH of the liquor following addition of the neutralisation additive is preferably in the range 2 to 3.5, most preferably about 2.5. This pH is thought to be optimum for maximum thorium and iron removal whilst minimising rare earth loss to the solid residue of the neutralisation step.**

**0 It is found that a minimum Fe.P molar ratio of 1, preferably an Fe:P molar ratio substantially equal to 1, in the water leach liquor feeding the neutralisation is desirable in order to minimise rare earth loss to the solid residue by premature precipitation of rare earth phosphates. Hence, it is preferred if necessary to add Fe upstream of the neutralisation step, e.g. to the acid bake, in order to optimise thorium and iron removal 5 with minimal rare earth element loss to the solid residue of the neutralisation step. Preferably, thorium removal is optimised to ensure an acceptable thorium level in the downstream rare earth end-product.**

**The precipitation of the rare earth elements from the post-neutralisation liquor is preferably by carbonate or double salt precipitation. The resultant composite rare earth 20 carbonates or hydroxides may be leached to produce a composite rare earth chloride as an end-product with acceptable impurity levels. The leach may be particularly desirable to remove aluminium from the composite rare earth carbonate.**

**With carbonate precipitation, 110% stoichiometric addition of a carbonate is preferred.**

**In an embodiment, the rare earth phosphates precipitated from the pre-leach liquor are 25 subjected to said acid bake together with the pre-leach residue. Alternatively, the rare earth phosphates precipitated from the pre-leach liquor are separately treated by an acid bake or by a caustic conversion process to permit recovery of rare earth elements therefrom.**

**Preferably, the pre-leach acid addition is about 110% stoichiometric with respect to calcium. As mentioned, apatite is a phosphate of calcium and calcium may also be a significant substituent in the monazite group mineral, for example where it is cheralite. This level of acid addition with respect to calcium optimises calcium removal by 5 subsequent precipitation from the liquor, for example by crystallisation as calcium nitrate.**

**The initial concentration of the nitric acid pre-leach is preferably in the range 30 to 60wt%, more preferably in the range 40 to 50wt%. The initial concentration of the hydrochloric acid pre-leach is preferably in the range 10 to 40wt%, more preferably in 0 the range 20 to 30wt%.**

# **Examples**

**A composite ore from Nolans Bore in the Northern Territory of Australia was subjected to a series of processing tests according to the invention. Rare earths in the ore were present in cheralite, (LREE, Ca)(P,Si)O4, which is a monazite group mineral, in** 15 **association** with fluorapatite,  $Ca_5(PO_4)_3$  (F,OH). The main ore resource consists of **about 82% w/w fluorapatite and 13% w/w cheralite. The remainder of the ore is about 3 to 4% clay with some calcite, dolomite, quartz and other minor minerals. Rare earths substitute for the calcium in the apatite structure. However, the bulk of the rare earths and thorium in the ore is contained in the cheralite, which is isomorphous with monazite. 20 SEM examination suggested that approximately 67% of the rare earths are associated with the cheralite and 33% are associated with the apatite. Two assays of the composition of the ore, determined by XRF, are set out in Table 1.**

Samples of the ore, riffled and ground to  $P_{80}$ =180  $\mu$ m, were subjected respectively to **nitric acid and hydrochloric acid pre-leach treatments. The details and results of these 25 treatments are set out respectively in Table 2 and Table 3. In the nitric acid pre-leach (Table 2), Ca and P extractions of 97% and 94% respectively were obtained. Rare earth dissolution of 40% can be expected. A rare earth residue concentrate containing <3% Ca and P, and 11% rare earths was produced. Similar results were obtained by hydrochloric acid pre-leaching (Table 3).**

**In the example reported in Table 2, the initial HNO<sup>3</sup> concentration is 54wt%. Further calculations and experiments have indicated that the broadly preferred initial HNO<sup>3</sup> concentration is 30 to 60wt%, but it is more preferably in the range 40 to 50wt%. The corresponding figures for the hydrochloric acid pre-leach are 10 to 40wt% and 20 to 5 30wt%.**

**The pre-leach liquors were treated by addition of ammonia to successfully precipitate rare earth phosphates. The details of the treatments and the results are set out in Tables 4 and 5. Rare earth recoveries of 89% and 98% were obtained from the nitrate and chloride liquors respectively.**

- **0 Four samples of solid residue from the hydrochloric acid pre-leach were subjected to a sulphuric acid bake in a rotary kiln, under standard conditions of 750 kg/lt H2SO4 addition and a temperature of 230°C for 2 hours. Rare earth recovery was consistently about 96%. It had previously been ascertained, in smaller sample tests in a static furnace that rare earth recovery fell from 90-92% at 230°C to less than 80% at 200°C, 5 and was not improved by increasing the temperature from 250°C to 300°C. As earlier**
- **foreshadowed, an Fe:P ratio of about <sup>1</sup> is thought to be required for an optimum neutralisation step, so bakes of pre-leach residue at temperatures above 250°C are preferably avoided since they were found to reduce Fe dissolution.**

**In the smaller sample tests, a bake time of 2h was found to be optimum; at a bake time 20 of 0.5h, rare earth recovery was significantly lower.**

**The solid residue of each acid bake test was water leached to produce a water leach liquor rich in rare earth elements. Waste solids were removed: these were found by SEM and XRD analysis to contain very little rare earth remnant and to consist mainly of quartz, alkali feldspar and remnant silicates, surrounded by clays and micas. The liquor 25 was subjected to a standard solvent extraction procedure to remove uranium values, e.g. employing a tertiary amine as solvent.**

**Neutralisation of the water leach bake liquor was then carried out as a purification step to selectively precipitate thorium, iron and aluminium from the rare earths. This was**

**effected at room temperature using 20wt% slurries of the neutralising additive, magnesia or magnesium carbonate. It was found that calcium carbonate and lime were less effective neutralising agents due to much higher rare earth losses, possibly due to co-precipitation/adsorption with gypsum due to the added calcium.**

By testing, it was further found that a minimum Fe:P molar ratio of above 1, preferably<br>about 1, was desirable in the water leach liquor feeding the neutralisation in order to<br>prevent rare earth loss to the solid residue b **about 1, was desirable in the water leach liquor feeding the neutralisation in order to prevent rare earth loss to the solid residue by premature precipitation of rare earth phosphates. It is thought that as the pH increases, iron phosphate precipitates. If there is not enough iron to react with the phosphate in the first instance, then the phosphate 0 will combine with rare earths and precipitate as rare earth phosphate. The Nolans Bore feed to the acid bake has a molar ratio Fe:P = 0.6, and therefore some iron addition to the acid bake or water leach liquor would be desirable in order to minimise rare earth losses in the neutralisation step.**

**The results of a variety of neutralisation tests are set out in Table 6.**

**5 Post-neutralisation liquor samples were now treated with ammonium carbonate to precipitate a composite rare earth carbonate, a commercial end product suitable for further processing by known techniques to separate out the rare earth components.**

**The composition of the composite rare earth carbonate is set out in Table 7. One optional further treatment, especially for the purpose of further reducing the aluminium 20 and thorium content, is to leach this carbonate with hydrochloric acid to produce a rare earth chloride end-product.**

**The earlier mentioned rare earth phosphates precipitated from the pre-leach liquor were split into samples. Some were successfully treated in a process stream similar to that outlined above for the pre-leach residue, but other processing options may be more 25 useful in extracting the rare earths from the phosphates.**



**Table <sup>1</sup>**

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Leach temp range  $(^{\circ}C)$  = 22-60 $^{\circ}C$  (uncontrolled)

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**Table 2**



# **Table 3 HCI Bulk Pre-Leach of Nolans Bore Ore at 45°C 510 kg/t HCI (-110% stoich. for Ca), 28 wt% initial solids**



CaSO4(head contains 0.2% S)



 $\ddot{\phantom{1}}$ 



afloc 333) but 25 no flocs formed Green precipitate

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# **Table 5**

# **Rare Earth Precipitation with NH<sup>3</sup> from Hydrochloric Acid Leach Liquor**

# *Precipitation conditions*



Volume Primary Filtrate  $(mL)$  = 143  $15$  Wash Volume (mL)  $=305$ 

### *Precipitate*



Comments: High concentration of RE's in the wash indicates that the wash is redissolving some precipitated RE's ie Acid concentration in wash is too strong % Precipitation based on liquor assumes no solution volume change





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\*MgCO<sup>3</sup> added instead of MgO, \*\*O<sup>2</sup> sparged

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The discussion of documents, acts, materials, devices, articles and the like is included in this specification solely for the purpose of providing a context for the present invention. It is not suggested or represented that any or all of these matters formed part of the prior art base or were common general knowledge in the field relevant to the present invention as it existed before the priority date of each claim of this application.

# THE CLAIMS DEFINING THE INVENTION ARE AS FOLLOWS:

1. A process for recovering rare earth elements from a composite ore containing rare earth elements that includes a monazite group mineral and an apatite mineral, 5 including:

pre-leaching the composite ore with an acid so as to substantially dissolve the apatite mineral into a pre-leach liquor;

precipitating rare earth elements from the pre-leach liquor;

subjecting the residue of the pre-leaching step to an acid bake treatment, followed by a water leach, to produce a water leach liquor rich in rare earth elements;

15 separating impurities including thorium and iron from the water leach liquor by introducing a neutralising additive to said water leach liquor rich in rare earth elements; and

precipitating rare earth elements from a post-neutralisation liquor.

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2. A process according to claim <sup>1</sup> wherein the acid employed for the pre-leaching is nitric acid.

3. A process according to claim 2 wherein the initial concentration of the nitric acid 25 is in the range 30 to 60wt%.

4. A process according to claim 2 wherein the initial concentration of the nitric acid is in the range 40 to 50wt%.

30 5. A process according to claim <sup>1</sup> wherein the acid employed for the pre-leaching is hydrochloric acid.

6. A process according to claim 5 wherein the concentration of the hydrochloric acid is in the range 10 to 40wt%.

7. A process according to claim 5 wherein the initial concentration of the hydrochloric acid is in the range 20 to 30wt%.

5 8. A process according to any one of claims <sup>1</sup> to 7 wherein rare earth elements are precipitated from the pre-leaching liquor as phosphates.

9. A process according to claim 8 wherein said precipitation of rare earth elements as phosphates from the pre-leaching liquor is by addition of ammonia or hydrated lime.

10. A process according to any one of claims <sup>1</sup> to 9 wherein the conditions of said precipitation of the rare earth elements from the pre-leaching liquor are selected to recover, with the rare earth elements and in a form separable subsequently, a major proportion of thorium present in the pre-leaching liquor from the composite ore.

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11. A process according to any one of claims <sup>1</sup> to 10 wherein the apatite is a fluorapatite, and a fluoride component of the pre-leaching liquor is removed by addition of a sodium salt to cause precipitation of the fluoride component.

20 12. A process according to any one of claims 1 to 11 wherein said acid bake treatment is carried out in a rotary kiln.

13. A process according to any one of claims 1 to 12 wherein said acid bake treatment is effected for between <sup>1</sup> and 3 hours.

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14. A process according to any one of claims 1 to 13 wherein said acid bake treatment is preferably a sulphuric acid bake.

15. A process according to any one of claims 1 to 14 wherein said acid bake 30 treatment is effected at a temperature in the range  $210-250^{\circ}$ C.

16. A process according to any one of claims <sup>1</sup> to 15 wherein said thorium and iron separation is effected by neutralisation with an alkali or alkaline earth additive selected to minimise rare earth co-precipitation or absorption.

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17. A process according to claim 16 wherein said neutralising additive is a magnesium compound.

18. A process according to any one of claims <sup>1</sup> to 17 wherein a minimum Fe:P molar ratio of <sup>1</sup> is maintained in the water leach liquor feeding the neutralisation in order to prevent rare earth loss to a solid residue by premature precipitation of rare 5 earth phosphates.

19. A process according to any one of claims <sup>1</sup> to 18 wherein said precipitation of the rare earth elements from the post-neutralisation liquor is by carbonate or double salt precipitation.

20. A process according to claim 19 wherein rare earth carbonates or hydroxides resulting from said carbonate or double salt precipitation are leached to produce a composite rare earth chloride as an end-product.

15 21. A process according to any one of claims <sup>1</sup> to 20 wherein a pre-leach acid addition is about 110% stoichiometric with respect to calcium.