Chemical Treatment of El Missikat Fluorite-Bearing ore Material, Egypt, for Recovery of Pure Yttrium Oxide

Khalil, M.M.H.(1), El Hussaini, O.M.(2), Abd El Wahab, G.M.(2), Swafy S.K(2) and Khalafalla M.S(2)
(1)Faculty of Science, Ain Shams University, Cairo, Egypt
(2)Nuclear Materials Authority, Cairo, Egypt

El Missikat fluorite-bearing ore material, Eastern Desert, Egypt, has been identified as containing a quantity of heavy rare earth elements (REE) such as yttrium fluorite mineral. Rare earth concentrate prepared from sulfate solution of El Missikat fluorite-bear ing ore material assays 0.615% of REE. The solution was prepared from ore material by two subsequent thermal steps namely; fluorine deactivation by MgO and roasting by (NH₄)₂SO₄. The deactivated roasted matrix was leached with hot distilled H₂O to prepare the solution which assays 0.846g/L of REE. This solution was treated with 20g of oxalic acid to produce impure RE oxalate cake which was ignited at 950°C. It was then dissolved in concentrated HNO₃ to prepare the RE nitrate solution for extracting yttrium using tri butyl phosphate (TBP) in kerosene. The produced yttrium oxide was of 97.2% purity.

Keywords: Yttrium, Extraction, TBP, Fluorite-bearing ore material

Introduction

Rare earths (lanthanides, yttrium and scandium) are important elements in photo-electronic and metallurgical industries as well as in nuclear energy programs. The demand for rare earth elements and their alloys as structural materials, fluxes, radiation detectors, diluent of plutonium etc. in nuclear technology is steadily increasing. High purity yttrium is often required, for a variety of uses in these specialized areas, [1]. Gabal El-Missikat occurs as a high mountain peak of younger granites (675 m above sea level) intruding the low lying older granites. Field relationships and observations confirm the presence of some different rock units such as: metavolcanics, older granitites, porphyry dyke swarms and younger granites. These granites are composed of quartz, alkali feldspar, plagioclase, biotite and muscovite with minor amounts of fluorite, zircon, rutile, apatite, monazite, epidote and topaz. Opaques, kaolinite and sericite, kaolinite and muscovite are found as secondary minerals [2].

According to Abd El Wraith et al. [3], El Missikat younger granite is rich in fluorine and high heavy REE (Y- group) enrichment in comparison with light REE (Ce - group). Fluorite (CaF₂) and yttrium fluorite (Ca,Y)F₂ which have a widespread occurrence as a common mineral ore deposits. The deep green fluorite contains Yb₂O₃ and Y₂O₃, [4]. Highly radioactive fluorite- bearing granite in EL Missikat occurs as blue to violet crystals due to the presence of U and Th in crystal lattice of the fluorite [5]. The conventional acidic or alkaline processing of El Missikat fluorite –bearing ore material for extracting REE, U and Cd have a serious environmental problems caused by HF and SO₂ evolution. Thus, the nonconventional processing might be preferred for such type of raw materials via two essential steps including thermal fluorine deactivation followed by roasting of the...
deactivated matrix. Fluorine deactivation process was carried out using MgO (to avoid the formation refractory insoluble REE fluorides) as the equation below, [6]:

\[ 2RE(FCO_3) + MgO \rightarrow RE_2O_3 + MgF_2 + 2CO_2 \]

A roasting step of the water insoluble fluorine deactivated matrix was carried out using (NH₄)₂SO₄ salt to convert it to water soluble matrix.

The separation of REE by solvent extraction (SX) technique depends on the slight difference in the distribution coefficient of an individual REE between two immiscible liquid phases (aqueous and organic phases). Different organic solvents have been used for investigating the separation of individual REE, one of these is tri-butyl phosphate (TBP) [8], di-2-ethylhexyl phosphoric acid (D2EHPA), [1] and carboxylic acids, [9]. Complete separation of yttrium from trivalent cations in aqueous HNO₃ can be achieved using tri-butyl phosphate (TBP), [7]. Finally, this work aims to treatment the refractory fluorite-bear granite for separation of yttrium using TBP in kerosene from El Missikate Fluorite-bearing ore material.

**Experimental**

**Materials characterization**

The representative studied raw material was collected from Gabal El Missikat granite, Eastern Desert, Egypt. It was ground to a mesh size of 60, washed with tap water to eliminate slimes. After drying, separation of heavy mineral with bromoform, the heavy fraction was separated iso-dynamically (1.5 amp nonmagnetic). The separated fractions were picked under the binocular microscope. To identify the mineralogical composition, the prepared sample was examined by X-ray diffraction (PhillipsPW3710) with a generator (PW1830) and Cu target tube (PW223/20) operated at 40 kV and 30 mA.

Magnesium oxide used in fluoride deactivation process was locally prepared at Nuclear Materials Authority, Egypt from the bittern solution of the Egyptian salt company El Max, Borg El Arab, Alexandria, [12]. The purity of MgO is not less than 99%. Solvent extraction experiments were carried out with AR grade of organic solvent, TBP of purity assays 99.5 %. Kerosene (boiling point 160–200°C) was used as diluent. All the other chemicals used in this work are also of high purity.

**Procedure**

The chemical composition of the raw material was identified by analyzing the major elements according to [13]. A portion of the fine ground ore material (0.5 g) was digested with acid mixture (H₂SO₄, HF, HNO₃, and HCl) until complete dissolution and diluted with distilled H₂O in 250 mL volumetric flask. Major elements were analyzed and their oxides were determined. For SiO₂ analysis, a 0.1 g weight of the ore material was mixed with 1.0 g of NaOH and fused using Lenton muffle furnace. The fused matrix was dissolved in 1:1 HCl solution and diluted with distilled H₂O in 250 mL volumetric flask then silicon content was estimated spectrophotometrically at $\lambda_{640}$ nm, using Shimadzu UV-vis160A [14].

To estimate the trace elements content in the studied ore material, 1.0 g was digested in acid mixture (as previously mentioned) until complete dissolution and dilution with distilled H₂O in 100 mL volumetric flask. The latter was directed to different analysis techniques for trace elements determination. The elements such as: Zn, Ni, Mn, Cd, V, Cu...etc. were measured using a Unicam Atomic Absorption Spectrophotometer (model-969 AAS) flame type at principal wavelengths. Uranium was analyzed using an oxidimetric titration method against ammonium metavanadate, [15]. The total REE referring to Y as heavy rare earth elements was quantitatively analyzed by 0.05 % Arsenazo (III) at $\lambda_{654}$ nm (using Shimadzu UV-vis160A spectrophotometer [14].

Determination of the fluoride content was conducted by dissolving 1.0 g of the ore using 5 ml of HCl and warming the solution to a lesser degree than the boiling temperature. 2.0 mL of the solution volume was treated with 0.5 mL of sodium acetate tri-hydrate to adjust the pH value to pH 5.0 followed by the addition of the total ionic strength adjustment buffer (TISAB) to adjust the ionic strength of the solution. The latter was then diluted up to 500 mL with double distilled H₂O in a plastic flask. A volume of 50 mL was taken and its fluoride ion concentration was measured by the ion selective electrode (Jenway, UK).
Preparation of leach liquor
Practically, a 300 g weight of El Missikat fluorite-bearing ground ore material underwent two roasting steps aiming to capture fluorine in closed system. The first step MgO was mixed well with ground ore at ratio (wt/wt) 1:10 and roasted at 350°C for 2h. After cooling the fluoride deactivation matrix was roasted again with (NH₄)₂SO₄ salt at(wt/wt) ratio of 1/2 at 200°C for 2h. The produced water soluble fluorine deactivation matrix was leached with distilled hot H₂O at (80-90°C) at S/L ratio of 1/2 and stirring for 2h. After cooling, washing and filtration the water insoluble MgF₂ residue was collected on filter paper. The filtrate contained the elements: REE, U and Cd was collected in 2L. This leach liquor's pH was found to be 8.5; It was used for yttrium extraction.

Separation of total REE
The prepared leach liquor was adjusted to a pH of 1.1. It was treated with oxalic acid aiming at precipitate the total REE. About 20 g / L of the solution was added with continuous stirring for 1h. After filtration and washing, the obtained cake was ignited at 850°C for 2h to produce the RE₂O₃ concentrate. The latter was washed, dried and redissolved in 31.5%HNO₃ acid at S/L ratio of 1/3. The REE nitrate solution was directed to solvent extraction for selective recovering of pure yttrium.

Individual separation of yttrium using TBP
Yttrium extraction from the prepared RE (NO₃)₃ solution was performed after pH adjustment to 1.0 with ammonia solution. It was then mixed with tributyl phosphate, (TBP) (sp.gr.0.979 g/mL) in kerosene to study the relevant extraction parameters such as TBP concentration, pH value, contact time and volume ratios (A/O) to optimize yttrium extraction. On the other hand, the concentration of the stripping solution, phase’s ratios (O/A) was done for optimizing yttrium stripping.

Concentrated yttrium strip solution was adjusted to a pH of 1.1, then 10% oxalic acid was added with continuous stirring for 1h. The obtained precipitate was filtered and washed with distilled H₂O to get rid of any adsorbed impurities. After dryness, the precipitate was ignited at 950°C for 3h and was identified by means of XRD.

Results and Discussion
Mineralogical composition
The present ore material underwent heavy liquid separation using bromoform where the heavy fraction was Iso-dynamically separated. The binuclear microscope examination of the separated fractions revealed that the non-magnetic fractions containing yttrium fluorite (Ca,Y)F₂ mineral distributed at 1 and 1.5 Amp. The separated mineral grains were investigated by XRD as shown in Figure(1): yttrium fluorite (Ca,Y)F₂, quartz,(SiO₂), and Musecovite, (K,Na) (Al, Mg, Fe,₃ (Si₃ Al)O₁₀ (OH)₂.

![Figure (1): XRD pattern of the mineralogical composition](image_url)
**Chemical composition**

The chemical analysis of ore material is given in Table (1) for major contents and some trace elements. The chemical composition reflects the previously mentioned mineralogical composition.

**Leach liquor preparation**

The ground fluorite bearing ore material was mixed with MgO at (wt/wt) ratio of 10% and roasted at 350°C for 2h in a closed system for fluoride deactivation. The fluoride matrix underwent the second roasting step with \((\text{NH}_4)_2\text{SO}_4\) salt at (wt/wt) ratio of 1/2 and 200°C for 2h. The obtained roasted cake was leached with hot water where the residue was mainly composed of MgF₂. On the other hand; the collected filtrates of 2L volume had pH of 8.5. The content of this leach liquor was given in Table (2).

**Separation of total REE**

The total REE (0.845 g/l) in the prepared leach solution were selectively precipitated by the gradual addition of oxalic acid 20 g/L with continuous stirring. Almost complete precipitation of REE as strong stable insoluble RE oxalate cake was obtained at pH 1.1 and stirring for 60 min. with precipitation efficiency of 98%. These precipitation conditions of RE oxalate from solutions are quite close agreement with, [16]; [17], [18] and [19]. The precipitation proceeds according to the following equation:

\[
2\text{RE}^{3+} + 3\text{H}_2\text{C}_2\text{O}_4 \rightarrow \text{RE}_2(\text{C}_2\text{O}_4)_3 \text{ (solid)} + 6\text{H}^+
\]

After filtration, washing and drying, the obtained RE oxalate cake (3.72g), was ignited at 850°C for 2h. It was washed, dried to produce a weight of 2.1g of RE₂O₃ concentrated cake. A weighed portion of the latter was analyzed by UV-VIS spectrophotometer to determine the REE concentration and the associated impurities. The purity of RE₂O₃ cake achieved 80.3% with impurities e.g.18% of Ca, 0.3% of Fe and 0.03% of Mg.

| Table (1): Chemical composition of El Missikat ore material |
|---------------------------------|-----------------|-----------------|
| **Major Contents**              | Conc., (%)      | Trace elements  |
| SiO₂                            | 65              | U               |
| TiO₂                            | 0.21            | Ni              |
| Al₂O₃                           | 3.40            | Cd              |
| Fe₂O₃                           | 1.1             | Cu              |
| CaO                             | 5.20            | V               |
| MgO                             | 1.20            | Th              |
| Na₂O                            | 0.2             | REE             |
| K₂O                             | 0.13            | Cr              |
| P₂O₅                            | 0.2             | Cu              |
| F                                | 20.2            | Co              |
| **L.O.I.**                      | 2.06            | Zn              |
| Total                           | 98.9            |                 |

*REE: measured as Y (Marchizinco, 2000) [14]

**L.O.I.** loss of ignition

| Table (2): Chemical constituents of the leach liquor |
|--------------------------------|-----------------|
| **Elements’ Ions**            | Conc. (g/L)     |
| REE'³                          | 0.845           |
| U’⁶                           | 0.225           |
| Fe’³                          | 1.13            |
| Cd’²                          | 0.064           |
| F’                            | 6               |
| Si’⁴                          | 34              |

Individual separation of yttrium using TBP

Due to the great importance of individual REE, the separation of pure \( Y_2O_3 \) was done via solvent extraction method. So, the \( RE_2O_3 \) cake was completely dissolved in 31.5% HNO\(_3\) acid at S/L ratio of 1/3 and stirring for 2h at 70\( ^\circ \)C [19]. The prepared nitrate solution assaying 17g/L of \( Y_2O_3 \) was used for yttrium extraction by TBP in kerosene in the presence of 0.5M of EDTA as a complexing agent for masking different impurities. The following parameters were studied to optimize yttrium extraction step.

Effect of TBP concentration

The effect of varying TBP concentration in the range 0.5 – 2.0 M in kerosene upon yttrium extraction efficiency (\%) was studied. The pH of aqueous solution was 0.5 at A/O ratio of 1/1 and a contact time of 10 min. After separation, yttrium was analyzed in the aqueous solution. The obtained data shown in Figure(2) revealed that the maximum yttrium extraction efficiency of 64\% was attained at TBP concentration of 1.5 M. However, further increase in TBP concentration to 2.0 M has no influence due to the interfering of other REE metal ions in the aqueous solution.

Effect of pH value

The effect of pH value of the prepared nitrate solution upon yttrium extraction efficiency was studied in the pH range from 0.5 to 2.0. The other parameters were kept constant at TBP concentration of 1.5 M in kerosene, A/O ratio of 1/1 and contact time of 10 min. The obtained data in Figure (3) indicated that the percentage of yttrium extraction efficiency increased from 64 to 81.2\% with increasing the pH value of the nitrate solution from 0.5 to 1.0. Further increase in pH values to 1.5 and 2.0 have a reverse effect, where yttrium extraction efficiency decreased to 70.3 \%

Effect of contact time

Figure (4) shows the effect of contact time (shaking time) upon yttrium extraction efficiency from the nitrate solution of pH 1.0 at A/O ratio of 1/1 and shaking at different time ranging from 5 to 20 min. The data indicates that, the extraction efficiency of yttrium achieved its maximum value of 92.0\% after 15 min. Further increase in contact time to 20 min has negligible effect.

Figure (2): Effect of TBP concentration upon yttrium extraction efficiency at pH of 0.5, A/O of 1/1 and contact time of 10 min
Figure (3): Effect of pH value upon yttrium extraction efficiency at A/O ratio of 1/1, TBP concentration of 1.5M and contact time of 10 min.

Figure (4): Effect of contact time upon yttrium extraction efficiency at pH of 1.0, A/O ratio of 1/1 and TBP concentration of 1.5M.

Table (3): Effect of A/O ratio upon Y extraction efficiency (%)

<table>
<thead>
<tr>
<th>A/O ratio</th>
<th>A Conc., ppm</th>
<th>O Conc., ppm</th>
<th>$D_O^A$</th>
<th>Extraction eff., %</th>
</tr>
</thead>
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<td>$\infty$</td>
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</tr>
<tr>
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<td>425</td>
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<td>13</td>
<td>97.5</td>
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<td>37740</td>
<td>8.53</td>
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</tr>
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<td>4/1</td>
<td>6290</td>
<td>42840</td>
<td>6.81</td>
<td>63.0</td>
</tr>
</tbody>
</table>

As shown from McCabe Thiele diagram in Figure(5), the extraction would be completed in three stages. From the obtained results, the optimum conditions for 92% yttrium extraction (of 15.64g/LY₂O₃) from the concentrated RE nitrate solution were done at TBP concentration of 1.5M, pH aqueous solution of 1.0 and 15 min. shaking time at an A/O ratio of 1/1. The optimum conditions were applied using 80 mL of Y(NO₃)₃ solution. The loaded TBP was then directed to the stripping stage. Different affecting factors were studied for optimizing the stripping step as follows:

**Effect of stripping agent type**

The yttrium loaded TBP was stripped with different stripping agents e.g. hot H₂O and 1.0M of HCl, HNO₃ and H₂SO₄, at O/A ratio of 1/1 and 5 min. contact time. After separation, the strip solution was analyzed for yttrium stripping efficiency. The obtained data revealed that the maximum Y stripping efficiency (35%) was attained using 1.0M H₂SO₄, Fig (6).

![Figure (5): Effect of A/O ratios up on yttrium extraction efficiency at TBP concentration of 1.5M, contact time of 10min and pH1.0, (McCabe Thiele diagram)](image)

![Figure (6): Effect of stripping agent’s type upon yttrium stripping efficiency at O/A ratio of 1/1 and contact time of 5 min.](image)
Effect of $\text{H}_2\text{SO}_4$ concentrations

The yttrium loaded on TBP was contacted with $\text{H}_2\text{SO}_4$ strip solution of different concentrations ranging from 1.0 to 4.0M. The other stripping parameters were kept constant at O/A ratio of 1/1 and 5.0 min. contact time. The obtained data in Figure(7) emphasized that yttrium stripping efficiency shows an improvement from 35.0 to 59.2% as the strip solution concentration increased from 1.0 to 3.0M. Further increasing in $\text{H}_2\text{SO}_4$ concentration to 4.0M has limited influence.

Effect of contact time

The effect of changing the stripping time from 5.0 to 30.0 upon yttrium stripping percentage using 3.0M $\text{H}_2\text{SO}_4$ solution at O/A ratio of 1/1 Fig (8). It was noticed that yttrium stripping efficiency is strongly influenced by increasing the stripping time. It increased from 59.2 to 92.0% by increasing the contact time from 5 to 25 min. almost no improvement in stripping after 25 min.

Effect of O/A ratio

This factor not only represents the equilibrium state between both aqueous and organic phases, but also shows the suitable applied O/A ratio. The studied O/A ratios ranged between 1/4 …1/1…. 4/1 using 3.0M $\text{H}_2\text{SO}_4$ and 25 min. contact time. The obtained data in Table 4 revealed that yttrium stripping efficiencies 100, 99.0 and 97.0% were attained at O/A ratios of 1/4, 1/3 and 1/2, respectively. Practically, these ratios were not applied because the yttrium contents were diluted in strip solution which is not preferred to recover yttrium. On the other hand, yttrium stripping efficiency of 78.0, 60.0 and 46.5% were attained at O/A ratios of 4/1, 1/3 and 1/2, respectively. It was found that the yttrium concentration was of highest value at O/A ratio of 1/4, Table (4). Drawing McCabe Thiele diagram shows that stripping requires two stages to reach equilibrium Figure(9). The O/A ratio was selected to be 1/1 for applying the precipitation step.

![Figure (7): Effect of $\text{H}_2\text{SO}_4$ concentration upon Y stripping efficiency at O/A ratio of 1/1 and contact time of 5 min.](image)

![Figure (8): Effect of contact time of stripping upon yttrium efficiency at 3.0M $\text{H}_2\text{SO}_4$strip solution and O/A ratio of 1/1](image)
The optimum conditions for regeneration 95.0% of yttrium from the loaded TBP are: 3.0 MH₂SO₄ strip solution and shaking for 25 min. at an O/A ratio of 1/1.

Preparation of Y₂O₃ final product

Yttrium in the strip solution assaying 14.8 g/L was precipitated by the gradual addition of 10% oxalic acid with continuous stirring for 60 min. Almost complete precipitation of yttrium as a strong stable insoluble oxalate cake was attained at pH of 1.1 with 99% precipitation efficiency. After ignition at 950°C for 2h, Y₂O₃ was produced. A small portion of the latter was analyzed by means of XRD and SEM-EDAX to define its chemical composition as shown in Figs. (10 & 11). On the other hand, another 0.1g weight was completely dissolved in 25 mL volumetric flask and analyzed by both of UV-VIS spectrophotometer and ICP instruments for its purity. The obtained result showed that Y₂O₃ purity was not less than 97% with very low impurities of Si and Ca. Their lower concentrations were not detected by both XRD and SEM-EDAX analysis of the yttrium oxide product.

Table (4): Effect of (O/A) ratio upon Y stripping efficiency, %

<table>
<thead>
<tr>
<th>O/A ratio</th>
<th>A Conc., (ppm)</th>
<th>O Conc., (ppm)</th>
<th>Dₒ²</th>
<th>Stripping Eff., (%)</th>
</tr>
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<td>1/4</td>
<td>3190</td>
<td>0</td>
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<tr>
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<td>29090</td>
<td>2094</td>
<td>46.5</td>
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</table>

Figure (9): Effect of O/A ratio upon Y stripping efficiency at contact time 25 min and 3.0MH₂SO₄ stripping solution, (McCabe Thiele diagram)
Conclusions

El Missikat fluorite-bearing ore material contains fluorides and SiO₂, Al₂O₃ and CaO as major oxides in addition to low concentrations of REE, U and Cd. Two successive roasting processes of El Missikate fluorite-bearing ore material were performed using 10% (wt/wt) MgO followed by 1/2 (wt/wt) (NH₄)₂SO₄ salt to deactivate the fluoride. Fluorine deactivated matrix was leached with distilled hot H₂O at (80-90°C) at S/L ratio of 1/2 and stirring for 2h to prepare the solution. Precipitation of REE was attained by addition of 20g/l oxalic acid at pH of 1.1 after 60 min stirring with 98% precipitation efficiency.

The ignited REE cake was dissolved in 31.5% nitric acid with S/L ratio 1/3 for 2h at 70°C for yttrium extraction by tri-butyl phosphate (TBP) from the nitrate solution.

The optimum extraction conditions of yttrium were: 15 min. contact time, pH of 1.0 and 1.5 M solvent concentration at A/O ratio 1/1. Yttrium strip from the loaded organic phase was achieved by 3.0M of H₂SO₄ at O/A ratio 1/1 for 25 min contact time. Yttrium was precipitated from the strip solution by addition of 10% oxalic acid.
gradually to the solution with stirring until almost complete precipitation. The obtained yttrium oxalate precipitate was filtered, washed, dried and ignited at 950°C to yield an Y₂O₃ with 97.2% purity.

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