Growth and Structural Characterization of Gadolinium Neodymium Oxalate Crystals Grown in Hydro-Silica Gel

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ABSTRACT

Gadolinium Neodymium Oxalate (GNO) single crystals were grown in hydrosilica gel by the diffusion of a mixture of aqueous solutions of the respective rare earth nitrates into a gel, impregnated with oxalic acid, in a test tube. Pink, transparent, hexagonal GNO crystals were obtained on optimization of growth parameters. The crystals are found to be monoclinic by the XRD analysis. The presence of water of hydration and carboxylic group in the grown crystals was confirmed by IR analysis. EDAX analysis confirmed the presence of Gd and Nd in the grown samples. The thermal decomposition behavior of the grown crystals were studied by thermogravimetric analysis (TGA) and differential thermal analysis (DTA). The thermal analysis results concur with the proposed crystal structure.

Key words : Rare earths, Oxalate Crystals, Gel growth

1. INTRODUCTION

The compounds of rare earths, especially oxalates are technologically important on account of their luminescent, ferroelectric and ferroelastic behavior [1,2]. Superconducting compounds have been synthesized by the controlled precipitation of rare earth oxalates followed by calcination [3]. Czochralski technique has been employed for the growth of rare earth oxalates at elevated temperature [4]. These crystals are found to be defective due to thermal stresses introduced during growth. To avoid these defects, methods of growth at room temperature are preferred. The gel technique [5] has been successfully employed for the growth of several rare earth oxalate crystals [6-9]. The authors report here the growth and structural characterization of gadolinium neodymium oxalate crystals in hydro-silica gel, and analysis of the thermal decomposition behaviour of the grown crystals.
2. EXPERIMENTAL

Aqueous solution of sodium meta silicate of density 1.03 g/cm\(^3\) is mixed with 1M oxalic acid to get desired pH. The resulting solution is allowed to set in test tubes of length 150mm and internal diameter 15mm. At a pH of 6 gelling is completed within two days. Over the fully set gel, supernatant solution comprising of a mixture of equal volumes of 0.5M gadolinium nitrate and 0.5M neodymium nitrate solutions acidified with nitric acid was slowly added. The rare earth ions diffuse slowly through the gel and react with the oxalic acid already incorporated in the medium in a controlled manner. Well defined pink coloured crystals of gadolinium neodymium oxalate of size 3.5 x 2.5 x 1mm\(^3\) are obtained in a period of three weeks.

The proposed reaction is

\[
\text{Gd(NO}_3\text{)}_3 + \text{Nd(NO}_3\text{)}_3 + 3\text{H}_2\text{C}_2\text{O}_4 \rightarrow \text{GdNd(C}_2\text{O}_4\text{)}_3.\text{nH}_2\text{O} + 6\text{HNO}_3
\]

The gel medium prevents turbulence and being chemically inert, it provides a three dimensional structure which permits controlled diffusion of reagents. The morphology and size of the crystals are found to depend on various growth parameters such as pH of the gel, density of the gel, concentration of the feed solution and acidity of the feed solution.

![Figure 1. Growth system of GNO crystals](image)

The lattice parameters of the grown crystals were determined by single crystal X-ray diffraction analysis using an ENRAF (Bruker) Nonius CAD4 single crystal X-ray diffractometer. IR absorption spectrum was recorded in the range 400 cm\(^{-1}\) to 4000 cm\(^{-1}\) using a Shimadzu IR 470 spectrophotometer. The thermal characteristics of the grown crystals were studied using a Shimadzu thermal analyser DT-40 in the temperature range 30-800°C at a scan rate of 10°C per minute.
3. RESULTS AND DISCUSSIONS

3.1. X-ray Diffraction Studies

X-ray diffraction studies confirmed that gadolinium neodymium oxalate crystal belongs to the monoclinic system with space group $P 2_1/C$ with $Z = 2$. The unit cell dimensions are $a=11.19$ Å, $b=9.89$ Å and $c=10.24$ Å. The crystal structure is expected to be identical with that proposed by Sheng Hua Huang et al.[10] for lanthanum oxalate, as evidenced by the similarity of the lattice parameters of the double rare earth oxalate crystal as that of the single component ones. It may be concluded that GNO crystals are formed by the substitutional exchange of the two rare earth ions and has the same structure as that of lanthanum oxalate crystal, $La_2(C_2O_4)_3 \cdot 10H_2O$.

3.2. Infrared Absorption Studies

FTIR studies reveals the presence of water of crystallization in the grown samples which is evidenced by a broad envelope extending from 3000 cm$^{-1}$ to 3600 cm$^{-1}$, due to the stretching modes of water molecules. The broad band appearing in the IR spectrum around 1620 cm$^{-1}$ may be identified as due to the asymmetric stretching mode of an oxalate ion [11]. The CO$_2$ group may be identified by a strong peak around 1316 cm$^{-1}$.

3.3. Energy Dispersive X-ray Analysis

![EDAX spectrum of GNO crystals](image)

Figure 2. EDAX spectrum of GNO crystals

The elemental incorporation of Gd and Nd in GNO crystal is confirmed using the EDAX analysis. The presence of Gd and Nd is established from their characteristic peaks in the EDAX spectrum. The peak heights or areas are a measure of the respective elements incorporated in the specimen.
3.4. Thermal Analysis

![Figure 3. TGA and DTA curves of GNO crystal](image)

Extensive research on the mechanism and chemical intermediates associated with the thermal decomposition of oxalates have been conducted by many researchers [12-15]. Various reports are available in literature regarding the thermal studies of lanthanide oxalates prepared either by mechanical mixing or by co-precipitation methods [16, 17]. Based on thermogravimetric and differential thermal analysis, Glasner et al. [18, 19] suggested a decomposition mechanism for anhydrous rare earth oxalate via a divalent intermediate. Gallagher and co-workers have reported thermal studies on the decomposition of europium oxalate [20].

The thermal decomposition stages of gadolinium neodymium oxalate decahydrate crystal are studied from the TGA and DTA curves shown in Figure 3. These stages are analyzed from the proportionate mass at each stage taking the initial weight as standard. The three steps involved in the decomposition process of GdNd(C₂O₄)₃.10H₂O are dehydration, decomposition of oxalate to carbonate and decomposition of carbonate to oxide. The liberation of ten molecules of water of crystallization in GNO crystal takes place in two stages in the temperature ranges 60-160°C and 160-350°C. In the first stage, seven water molecules which are distributed randomly in the intervening lattice space are eliminated. In the second stage, the remaining three water molecules are liberated. These two different stages in the dehydration process appear as two distinct endothermic peaks in the DTA curve.

The next stage of decomposition occurs as a continuous process of dehydration. The oxalate decomposition of GNO crystals can be assigned to two stages. During the first stage in the temperature range 350-520°C, two CO₂ molecules and three CO molecules are lost resulting in the formation of an unstable intermediate, dioxygen carbonate. The reduction of this intermediate compound into the oxide form occurs in the range 520-650°C with the release of one CO₂ molecule. These decomposition reactions are represented by the two exotherms in the DTA
The thermal studies on GNO crystals are in good agreement with the proposed chemical formula and hydration number for these crystals.

The mechanism of thermal decomposition of GNO crystals may be described as follows.

\[
\begin{align*}
\text{GdNd(C}_2\text{O}_4)_3.10\text{H}_2\text{O} & \xrightarrow{\text{60-160°C}} \text{GdNd(C}_2\text{O}_4)_3.3\text{H}_2\text{O} \\
\text{GdNd(C}_2\text{O}_4)_3.3\text{H}_2\text{O} & \xrightarrow{\text{160-350°C}} \text{GdNd(C}_2\text{O}_4)_3 \\
\text{GdNd(C}_2\text{O}_4)_3 & \xrightarrow{\text{350-520°C}} \text{GdNdO}_2\text{CO}_3 \\
\text{GdNdO}_2\text{CO}_3 & \xrightarrow{\text{520-650°C}} \text{GdNdO}_3
\end{align*}
\]

The thermo analytical data for GNO crystals is given in Table 1.

<table>
<thead>
<tr>
<th>Stage</th>
<th>Decomposition Temperature range (°C)</th>
<th>Loss of material</th>
<th>Observed mass loss (%)</th>
<th>Calculated mass loss (%)</th>
<th>Nature of reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>60 – 160</td>
<td>7H₂O</td>
<td>16.08</td>
<td>16.91</td>
<td>Endo dehydration</td>
</tr>
<tr>
<td>2</td>
<td>160 –350</td>
<td>3H₂O</td>
<td>7.39</td>
<td>7.25</td>
<td>Endo dehydration</td>
</tr>
<tr>
<td>3</td>
<td>350 – 520</td>
<td>2CO₂ + 3CO</td>
<td>23.95</td>
<td>23.07</td>
<td>Exo decomposition</td>
</tr>
<tr>
<td>4</td>
<td>520 – 650</td>
<td>CO₂</td>
<td>6.14</td>
<td>5.90</td>
<td>Exo decomposition</td>
</tr>
</tbody>
</table>

Table-1. Thermo analytical data of GdNd(C₂O₄)₃.10H₂O

4. CONCLUSION

The gel grown gadolinium neodymium sample is crystalline as evidenced by XRD analysis and has identical structure as that of single rare earth oxalate crystals. Presence of water molecules in the sample is confirmed by the broad envelope in the IR spectrum extending from 300cm⁻¹ to 3600cm⁻¹ and that of carboxylic groups is evidenced by the peak at 1620 cm⁻¹. The presence of Gd and Nd in the grown sample is established from their characteristic peaks in the EDAX spectrum. The proposed chemical formula and crystal structure for the grown sample are in good
agreement with the results obtained from TGA and DTA studies. The observed mass loss and calculated mass loss agrees in all stages of dehydration and decomposition.

REFERENCES