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**CERAMICS** INTERNATIONAL

Ceramics International 39 (2013) 8767-8771

www.elsevier.com/locate/ceramint

# Low temperature synthesis of nano-crystalline gadolinium titanate by molten salt route

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Received 9 July 2012; received in revised form 16 April 2013; accepted 17 April 2013 Available online 30 April 2013

#### Abstract

Nano-crystalline Gadolinium Titanate (Gd<sub>2</sub>Ti<sub>2</sub>O<sub>7</sub>) powder was successfully synthesized by "**Single Step Molten Salt Technique**". LiCl–KCl eutectic mixture was used as a molten medium for the reaction. The duration of the synthesis was 10 h. Stoichiometric proportion of the reactants were mixed in an (LiCl–KCl) eutectic medium and treated at 750 °C in an electrical resistance furnace. Single phase Gadolinium Titanate compound was obtained by the thermal process. The synthesized powders were characterized using XRD, FT-IR, UV, EDAX and XPS analyses. The morphology of the powder was examined using SEM and TEM techniques. From the above studies, it has been concluded that pure crystalline Gadolinium Titanate powders can be synthesized via low temperature molten salt process. © 2013 Elsevier Ltd and Techna Group S.r.l. All rights reserved.

Keywords: B. Electron microscopy; E. Nuclear applications; Molten salt method

### 1. Introduction

Pyrochlore compounds are considered as potential multifunctional materials, which find various technological applications in the areas such as piezoelectricity, ferro and ferrimagnetism, luminescence, giant magneto-resistance and nuclear waste immobilization. Properties of pyrochlore oxides widely vary from insulators or semiconductors to compounds with high ionic, electronic or mixed conductivity making them possible for monolithic fuel cells and gas sensors [1–3].

Compounds exhibiting the pyrochlore structure have been extensively investigated, especially for their properties of anion and mixed conduction [4,5], magneto-resistance and superconductivity. Many applications of these materials have been envisaged as sensors and transistors [6], dielectrics and fast ion conductors [4,7], electrocatalysts [8] and nuclear waste encapsulation [9]. In particular, titanate and stannate pyrochlores have been synthesized for over 40 years [10], and hence a profound knowledge of their structural trends [11],

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thermodynamic properties [12], disorders [13,14] and nonstoichiometry [15] exist in the literature.

Titanate pyrochlore,  $A_2Ti_2O_7$ , materials are important, because of their potential use as solid electrolytes and mixed ionic/electronic conducting electrodes [16–20], catalysts and ferroelectric/dielectric device components [21].

Rare-earth pyrochlore (Gd<sub>2</sub>Ti<sub>2</sub>O<sub>7</sub>) has been synthesized by different methods such as sol–gel [22–24], solid state reaction [25,26], chemical-coprecipitation, calcinations method [27], single crystal growth [28–30], conventional mixed metal oxide method [31] and Pechini process. However, these methods (except in the case of single crystal growth method) involve multi-step complicated reaction routes with longer duration for synthesis. These reactions result in non-homogenous polycrystals with irregular morphology and broad particle size distribution. Furthermore, higher calcination temperature is needed to obtain a single phase compound. Single crystal growth method involves sophisticated instrumentation. To overcome all these disadvantages, we have made an attempt through a novel molten salt method (MSM) to synthesize high purity Gadolinium Titanate (Gd<sub>2</sub>Ti<sub>2</sub>O<sub>7</sub>) compound.

In this method, the molten salt is used as the reaction medium, where the reactants are dissolved, producing pure compounds of interest. Since, the rate of diffusion of reactants

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<sup>0272-8842/\$-</sup>see front matter © 2013 Elsevier Ltd and Techna Group S.r.l. All rights reserved. http://dx.doi.org/10.1016/j.ceramint.2013.04.063

in the molten melt is much higher than those in the solid-state reaction, the homogeneous oxide materials are formed at an accelerated rate. In this study, we report the preparation of nano-crystalline  $Gd_2Ti_2O_7$  compound using Gadolinium oxide  $(Gd_2O_3)$  and Metatitanic acid  $(H_2TiO_3)$  as the reactants. LiCl-KCl eutectic salt mixture was used as a flux during the synthesis process.

# 2. Experimental procedure

# 2.1. Synthesis of Gd<sub>2</sub>Ti<sub>2</sub>O<sub>7</sub> powders

Gadolinium oxide (98% purity, Aldrich), and Metatitanic acid (Travancore Titanium Products Limited, Kerala, India) were used as the starting materials for the synthesis. The reactants were mixed in stoichiometric proportions and thoroughly ground using mortar and pestle. An equimolar mixture of lithium chloride and potassium chloride salts were used as the flux. The reactants and the flux were placed in an high density alumina crucible and fired at 750 °C for 10 h. At this temperature, the salt medium forms a molten flux, and the precursor oxides disperse, dissociate, rearrange, and diffuse rapidly throughout the melt forming the final product. Finally, the resultant product was washed with dilute hydrochloric acid (10%) followed by hot distilled water for several times. Then the residue was dried at 50 °C for 15 min to remove the surface moisture, yielding a free flowing fine crystalline white powder.

 $Gd_2O_3+2H_2TiO_3 \rightarrow Gd_2Ti_2O_7+2H_2O$ 

### 2.2. Characterization

XRD analysis was carried out using an X-ray powder diffractometer (Philips 8030 X-ray Diffractor) with Cu Ka radiation ( $\lambda = 1.5406$  Å). The XRD patterns were compared with the standard JCPDS: (23-0259) files. FTIR measurements were performed with a Perklin Elmer UK paragon-500 spectroscope in the mid infrared region  $(4000-400 \text{ cm}^{-1})$ . For each sample preparation, approximately 2 mg of sample was dispersed in 300 mg of KBr powder and pressed into a pellet and used for the analysis. The elemental analysis of the powders was done using the energy dispersive X-ray analysis (EDAX) technique. This analytical tool was an attachment to the SEM (scanning electron microscope) unit. The surface morphology was analyzed by using a scanning electron microscopy (JEOL-JSM-3.5 CF-JAPAN). The nano-size of the powder was assessed by a transmission electron microscope. TEM images were recorded using a JEOL-3010 transmission electron microscope using 300 kV accelerating voltage with a 400-mesh ultrathin carbon type A copper grid.

#### 3. Results and discussion

#### 3.1. X-ray diffraction

Fig. 1 shows the X-ray diffraction patterns of synthesized  $Gd_2Ti_2O_7$  powders obtained by the molten salt route. The

diffraction peaks of the samples can be indexed as spinel Gadolinium Titanate. From the XRD data, the lattice constant 'a' was calculated using the formula

$$a = d\sqrt{h^2 + k^2 + l^2}$$

where *d* is the inter-planar spacing (Å) and *h*, *k*, *l* are the Miller indices. XRD patterns of the synthesized polycrystalline powders show characteristic reflections of  $Gd_2Ti_2O_7$  pyrochlore type crystal strcture. The main reflections observed correspond to (222), (440) and (622) planes with the  $2\theta \sim 30^{\circ}$  (100% relative intensity), and weak peaks at 50.12° (50%) and 60.198 (60%) are confirming the presence of  $Gd_2Ti_2O_7$  pyrochlore compound. Few peaks appearing in the spectrum can be assigned to the presence of rutile TiO<sub>2</sub>. The presence of the pyrochlore phase does not have any influence on the pyrochlore phase [26].

The unit cell parameters have been determined from the XRD data and found to be 10.1687 Å. This calculated lattice constant value is in good agreement with the standard JCPDS data (a = 10.1860 Å) [4]. The line broadening of the diffraction peaks has been ascribed to the fact that the synthesized powders have nano-structured morphology. The average crystalline size of the powder was determined by using Debye–Scherrer's formula

## $D = 0.9\lambda/(\beta \cos\theta)$

where  $\beta = \Delta/2$  ( $\Delta$  is the full width at half maximum for a given diffraction peak),  $\lambda =$  wavelength of the Cu K $\alpha$  radiation and (wave length  $\lambda_{Cu} = 1.5406$  Å), and  $\theta =$  Bragg's angle.

The average crystalline size of the particles is found to be in the range of 31–39 nm.

#### 3.2. FT-IR spectroscopy

The FT-IR spectrum of the single phase  $Gd_2Ti_2O_7$  compound is shown in Fig. 2. FT-IR bands have been analyzed for identifying the functional groups present in the compound. From the FT-IR spectrum, it is perceived that the absorption bands appearing between 1700 and 2400 cm<sup>-1</sup> correspond to





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