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Preparation and Evaluation of NiFe₂O₄ and CuFe₂O₄ Nanocatalysts by Combination of Sol- Gel Auto-Combustion Method and Irradiation Technique

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ABSTRACT

The dried nitrate-urea gcls exhibit the combination of auto-catalytic combustion behavior and ultrasonic, which can be used to synthesize the nanocatalyst ferrite powders. Cu and Ni ferrites nanocatalyst powders with composition of $CuFe_2D_4$ and $NiFe_2O_4$ were synthesized by a sol-gel auto catalytic combustion process. The molar ratio between metal inns and urea was 1:1.2. The sol- gel process was done at 80 °C. The nanoparticle crystallines have been calcined at 800 °C.

Comhustioa behavior and crystallite size of synthesized powders were iavestigated with the help of Scanning Electron Microscopy observation and X-ray diffraction technique. X- ray diffraction and Scanning Electron Microscopy were carried out for characterization of the powders. The grain size of the prepared ferrite powders is found to he in the range 30-35 nm.

Keywords: Auto-Combustion: Irradiation Technique: Sol- Gel, CuFe2O4; NiFe2O4; SEM; XRD

INTRODUCTION

The sol- gel method, in particular, is one of the most useful and attractive techniques for the synthesis of aanosized ferrite materials, because of its advantages such as; good stoichiometric control and the production of ultrafiae particles with a narrow size distribution in a relatively short processing time at very low temperature. Sol-gel methods generally refer to the hydrolysis and condensation of metal alkoxides or alkoxide precursors, leading to dispersions of oxide particles in a sol. The sol then dried or geled by solvent removal or by chemical reaction. In general, water is used as the solvent, hut the precursors can also be hydrolyzed hy an acid or hasic medium The eatalysis process induces the formation of colloidal as well as polymeric from of the gel [1, 2].

Soft and hard ferrites are a group of technologically

important magnetic materials. One of these ferrites is NiFe₂O₄. This material is largely used in electric and electronic devices, radarabsorbing coatings, ferro-fluids, and catalysts [3]. Nickel ferrite dominates the corrosion product oxide inventory in pressurized heavy water reactors and hence it plays a major role in the activity transport process [4].

Spinels of the type of $M^{2*}M^{3*}_{2}O_{4}$ attract the research interest because of their unique properties and multiple applications in various fields [5-8]. Copper ferrite, CuFe₂O₄, is one of these compounds. Several methods have been employed to prepare copper ferrite, such as, solgel method [9-14], eo- precipitation [15-19], solid state reaction [20] and auto - comhustion

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[21]. Combustion synthesis processes are characterized by high-temperatures, fast heating rates and shart reaction times. These features make CS an attractive method for the manufacture af technologically useful materials at lower costs eampared to canveational ceramic processes. In solid state comhustian (SSC), initial reactants, inter-mediates and final products are all in the solid state. Comhustion af solid reactants can occur in two modes: (i) Linear ar self-propagating, high temperature synthesis (SHS) and (ii) hulk or valume comhustion synthesis (VCS) [22].

For solid-state synthesis, it is difficult to achieve unifarmity of product and needs higher synthesis temperature and longer sintering time [23]. Compared with solid-state synthesis, the co- precipitation method makes the materials react uniformly at molecular level and has the advantages of lower polycrystalline-synthesized temperature and sharter sintering time [24 - 26]. The results show that the nanometer CuFe₂O₄ has high a catalytic activity. In this study, CuFe₂O₄ and NiFe₂O₄ nanocatalysts was synthesized by cambination auto- combustion and irradiation beam technique.

EXPERIMENTAL

CuFe₂O₄ and NiFe₂O₄ nana powders were prepared by soil-gel method. (Cu $(NO_3)_2$ -3H₂O, Merck), (N_1) $(NO_3)_2.6H_2O_1$ Merck), - íFe (NO₃)₃·9H₂O, Mcrck), (urae), Merck) and NH₄OH (Merck) were used as raw materials. All the reagents were used without further purification. X-ray diffraction (XRD) patterns af nanopowders were obtained with an X-ray diffractometer (Model: XPERT-MPO, Philips) using Cu Ka radiation (λ =1 5406 *A) with aperated at 40 kV and current of 40 mA. The shape and morphalogy af powder were analyzed by Seanning Electron Microscopy (SEM- Phillips XL 30). The HF-Frequenz 35 KHz, 240 w / Made In Germany was used as ultrasonic hath at 15 °C.

Appropriate amounts of ferric, nickel and copper nitrates and throurea, were first dissolved in a minimum amount of deionized water. The molar ratio of nitrates to thiourea was 1:2. A small amount of ammonia was added to the solution to adjust the pH value at about 10. During this procedure, the solution was continuously stirred using a mechanical agitator.

Then, the mixed solution was poured in to a dish and heated and stirred constantly to transform it into a xerogel. When ignited points were observed, the dried gel burnt in a selfprapagating cambustion manner until all the gel was burnt aut completely to form a loose powder. The powder was then calcined at 800 °C for 4 h. In orther hand, to prepare manadisperse CuFe₂O₄ and NiFe₂O₄ nanoparticles, the powder was dispersed by ultrasonic technique for 15 minutes.

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RESULTS AND DISCUSSION

The XRD patterns af the Cu and Ni ferrites nanot crystalliae powders are shown in Fig.1. The particle size af the samples has been determined emplaying the Scherrer equation: $D = k\lambda / \beta \cos \theta$ [1]

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Where β is the full width half maximum (rad), λ the wavelength of the X-ray, θ the angle between the incident and diffracted heams (degree) and D the particle size of the sample (nm). The results of XRD shaw at lower temperature the diffraction lines have confirmed the farmatian af single phase of spinel ferrite NiFe₂O₄ nanoparticles and for nanopawders af CuFe₂O₄ were observed composition of phases CuFe₂O₄, CuO and Fe₂O₃.

The nanoparticles structural αf CuFe₂O₄ are Tetragonal, Moaoclinic, Rhamhuhedral, and for NiFe₂O₄ is Cuhie. The grain sizes af the prepared ferrites are found to he in the range 30-35 nm.

The structural morphology of the nanoparticles was investigated using Scanning I Electroa Microscopy (SEM). Fig. 2, shows the SEM images af CuFe₂O₄ and NiFe₂O₄ nanocatalysts.

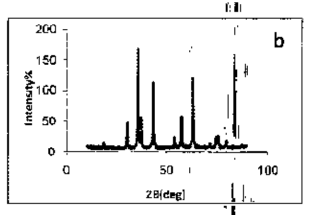


Fig. 1. The XRD patterns for CuFe₂O₄ (a) and NiFe₂O₄ (b) ferrite nanocatalysts powder

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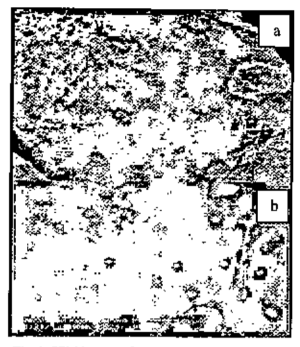


Fig. 2. SEM images of nanoparticles for a) CuFc₂O₄ and b) NiFe₂O₄.

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CONCLUSION

A nitrate- urea gels were prepared from metal nitrates and fuel by a sol- gel auto- combustion process in order to synthesize CuFe₂O₄ and NiFe₂O₄ ferrites. The well- crystalline copper ferrite and nickel ferrite was produced when pH = 10. So, it is necessary to adjust appropriate pH to produce pure copper ferrite and nickel fernite. The grain sizes of the prepared ferrites are found to he in the range 30- 35 nm. The particles have been calcined at 800 °C for 4 h. Then the products were placed in ultrasonic hath of n-butanol for 15 minutes. SEM results showed that the grains were regular sphere-shaped nanoparticles. The XRD data sbnw nanocatalyst pnwders have phases fnr CuFc2O4: CuFe2O4, CuO and Fe₂O₃ and for NiFe₂O₄: NiFe₂O₄.

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