International Journal of Bio-Inorganic Hybrid Nanomaterials

2Calcination Treated of Mixed Spinel ZnFe₂O₄ Synthesized by **Combustion Method**

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Received: 2 September 2014; Accepted: 6 November 2014

ABSTRACT

ZnFe $_2\rm O_4$ mixed spinel ferrite nanoparticles obtained through a citrate-nitrate combustion method. Ignited products calcined at different temperatures in 600-900°C range and effect of calcination treatment temperature investigated. Final Products characterized by X-ray Diffraction (XRD), Fourier-Transform Infrared spectroscopy (FTIR), and Field Emission Scanning Electron Microscopy (FESEM). The average particle size calculated from Scherrer equation. Results demonstrate good agreement with total rule of increase in particle size by increasing of calcination temperature. Average particle size from X-ray data sheets for 900°C was about 54 nm.

Keyword: Nano-powder; ZnFe₂O₄; Combustion; Surfactant; X-ray diffraction.

1. INTRODUCTION

Nanostructured materials are considered more than their bulk forms, due to quantum confinement that due to their advanced properties [1]. Cubic spinel ferrite nanoparticles such as $Fe₃O₄$, $MnFe₂O₄$, $NiFe₂O₄$ and $\text{ZnFe}_{2}\text{O}_{4}$ exhibit excellent magnetic properties, chemi cal stability and mechanical resistance $[2, 3]$. Among these, Zinc ferrite nanoparticles have wide applications such as $T1$ contrast agent [4], biomedicine applications [5], hyperthermia treatments [6], gas sensing [7-9], photo-catalysis [10], and photochemical water splitting [11]. This ferrite is of interest not only to technologi-
cal applications as denoted, but also has great attrac-

tion to basic researches in their magnetic and electrical

properties [1]. Based on cation distribution between A

(tetrahedral) and B (octahedral) sites in spinel struc-

ture, these divided in two categories: normal spi properties [1]. Based on cation distribution between A ture, these divided in two categories: normal spinel and $tetrahedral$) and B $(octahedral)$ sites in spinel strucreverse spinel. In normal spinel M^{2+} ions ($M = Fe^{2+}$, Mn^{2+} , Ni^{2+} , Zn^{2+}) occupied only tetrahedral sites and in reverse spinel where as it occupied only octahedral sites in an inverse one $[12, 13]$. Against bulk form, the nano-crystalline $\text{ZnFe}_{2}\text{O}_{4}$ system always shows up as a tween A and B sites $[14, 5]$. Various synthesis methods mixed spinel by distribution of Fe^{3+} and Zn^{2+} ions bedeveloped to produce zinc ferrite nanoparticles, such

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as simple and combine mechanical routes $[15, 16]$, Precipitation [17-20], different type of Combustion reaction [1, 21-24]. Among these methods, combustion routes are highly interesting for their low cost and simplicity.

In this work, fine ZnFe_2O_4 nanoparticles have been synthesized via a simple combustion technique. Results demonstrate that calcination temperature play an important role in the phase formation and size controlling in this process.

2. MATERIALS AND METHODS

Materials 2.1.

Zinc (II) nitrate tetra hydrate (98%) , Iron (III) nitrate Aldrich Co. and used without any purification. Deion-
ized-water applied as solvent. monia (25%) were purchased from Merck and Sigma-
Aldrich Co. and used without any purification. Deionnona-hydrate (99-101%), Citric acid (99%) and Am-
monia (25%) were purchased from Merck and Sigma-

2.2. Synthesis of ZnFe₂O₄

Zinc ferrite nanoparticles were prepared successfully propriate amount of metal nitrates in Zn:Fe, 1:2 molar by means of a citrate-nitrate combustion method. Apratio mixed together and solve in minimum amount tios with metal nitrates, solves in minimum amount of deionized water. Citric acid weighted in 1:3 ration. The pH of homogenized mixture adjusts at 7 by of deionized water, and added to metal nitrates soludropwise ammonia addition. After adequate heating

time (about 3 hours), viscous obtained gel expose at top of a burner flame. After some minutes gel was boiling and lose remain water. Next auto propagating combustion process start and gel transformed to loose powder. Obtained powders were calcined at 600, 700, 800, 900 $^{\circ}$ C for 2 h, washed by ethanol and deionized sonic bath for 15 minutes characterized after drying water then solve in chloroform and irradiated in ultraand denoted as S1, S2, S3, S4, respectively.

2.3. Characterization process

ips, XPERT- MPD, operates at 40 kV and 30 mA. Powder X-ray measurement was recorded with Phil-The morphology was investigated by using FESEM, Philips XL30 with 40 kV operating voltage. An FTIR spectrum was recorded as KBr pellets on Shimadzo, FTIR-300 spectrophotometer.

3. RESULTS AND DISCUSSION

X-ray analysis presented at Figure 1. As shown in these four XRD patterns, four weak unmatched peak around 2theta 31, 33, 36, and 54 degrees attributed to presence of some impurity in samples S1, S2, and S3. As seems increase in calcination temperature due to ray pattern, three peaks completely eliminated and the decrease of these impurity peaks intensity. In S4 Xone remains in 32 degrees has very low intensity. The well matched peaks of S4 confirmed with JCPDS card No. 01-079-1150, for Zinc iron oxide by ZnFe_2O_4 stoi

3O .(ZnO and

Sample name	Calcination temperature $(^{\circ}C)$	Phase composition	Average Crystallite $size$ (nm)	Crystallinity percent $(\%)$
S ₁	600	$\text{ZnFe}_{2}\text{O}_{4}$ Unmatched impurity	33	73.60
S ₂	700	$\text{ZnFe}_{2}\text{O}_{4}$ Unmatched impurity	40	73.33
S ₃	800	ZnFe_2O_4 Unmatched impurity	46	77.99
S ₄	900	$\text{ZnFe}_{2}\text{O}_{4}$	54	83.59

Table 1: Chemical composition of calcined samples at different temperatures.

chiometric ferrite chemical formula.

The average crystallite sizes for main phase, Zn-
Fe₂O₄ nano-particles, calculated using Scherrer's The average crystallite sizes for main phase, Znequation (Eq. 1).

$$
d = K\lambda / \beta \cos \theta \tag{Eq. 1}
$$

Where d is the grain size, β is half-intensity width of the relevant diffraction: λ is X-ray wavelength and θ the diffraction angle.

The crystallinity percentage calculated from net to ple compositions, Average crystallite size and Crystallinity percentage abstracted in Table 1. total area ratio obtained from X-ray data series. Sam-
ple compositions, Average crystallite size and Crystaltotal area ratio obtained from X-ray data series. Sam-

rer's equation for fourth sample presented at Table 1. The average crystallite size calculated from Scher-As calculated, the single phase S4 has 54 nm average crystallite size. Results demonstrate a slight increase

1.S FIGURE 1.S FIR spectrum of S1. FIGURE 1.S of S1. FIGURE 1.S of <i>S1. FIGURE PIGURE FIGURE PIGURE CS1. PIGURE CS1. CS1. CS1. CS1. CS1. **CS1.** *CS1.* **CS**

ture. This was in good agreement by previous reports. in crystallite size by increase in calcination tempera-[25]. Obviously, when crystallite lattice can't abide applied treated temperatures may be broken up to sub-
phase as occur in $MnFe₂O₄$ ferrites in temperatures up applied treated temperatures may be broken up to subto 600° C or composed the new phases.

The FESEM images of S1 presented at Figure 2. As shown in these Figure ZnFe_2O_4 nanoparticles has a fore characterization, hard aggregations seem in some spherical shape. Against of ultrasound treatment bearea, probably due to long period of calcination or presents of impurities.

trated at Figure 3. Characteristic vibrations around The FTIR spectrum of selected sample, S1, illus- 3434 and 1628 cm⁻¹ were indicated the asymmetric and symmetric H-O stretching bounds. The 675 and 547 cm⁻¹ peak demonstrates metal-oxygen stretching

bonds in tetrahedral situation. The intense peak in 419 cm⁻¹ attributed to metal-oxygen stretching band in octahedral site. This result was in good agreement with some previous reports such as Wan et al. [4] and Prithviraj et al. [21].

4. CONCLUSIONS

Zinc ferrite nano-particles were produced successfully with the simple combustion method. Citrate-nitrate tures applied in burnt powders. Results demonstrate precursor selected and different calcination temperathat particle size increase by increasing in calcination temperatures. Also, single phase composition obtained at 900° C and in lower temperatures some impurities observed. For single phase sample, S4, particle size of 54 nm calculated

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