Structural Characterization and Magnetic properties of Chromium substituted Nickel nano ferrites

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ABSTRACT

Mixed Ni-Cr nano ferrite system having the compositional formula NiCr_xFe_{2-x}O₄ (Where X = 0.1, 0.3, 0.5, 0.7, 0.9 and 1.0) were synthesized using Citrate-Gel auto combustion method. The structural characterization was investigated by X-ray diffraction (XRD). X-ray analysis shows that the samples are cubic spinel structure (single phase) without any impurity peak and average crystallite size was in the range 8.55-10.36nm. The X- ray density, bulk density and porosity were calculated. Magnetic properties were measured at room temperature in the applied field of ± 6 KOe. The obtained M-H loop area is very narrow, hence the synthesized nano ferrites become soft magnetic materials with less magnetization. Magnetic parameters such as saturation magnetization, magnetic moment and Y-K angle were measured and discussed with regard to Cr³⁺ ion concentration. The Curie temperature is measured from Loria-Sinha method. The observed results can be explained on the basis of composition.

Keywords: Ni-Cr nano ferrites, Citrate-Gel auto combustion method, XRD, magnetic properties.

1. INTRODUCTION

Among all the magnetic materials, ferrites are the important non-conductive ferromagnetic ceramic compounds which as composing of iron oxides (Fe_2O_3 or Fe_3O_4) and metal oxides (MO). These are show the magnetic and electrical transport properties with large number of application in various fields [1-2]. The ferrite composition is MFe₂O₄, where M is divalent metal ion and Fe is trivalent ion. Many physicists are focusing on the study of properties of nanostructured materials due to the nanostructured systems form a bridge between single molecule and bulk system and their increasing applications in several technological and scientific fields [3-4]. The nanostructure material shows evidence of novel properties over those exhibited by bulk material of same composition [5].

In the current years researchers are focus on synthesis and characterization of new class of soft nano size magnetic materials, which possess both large initial permeability and saturation magnetization [6-7]. In the field of science and technology nano size materials are important because same composition of different nano size particles can exhibit the changes in properties. When the size of the particle reduced into nano scale, the properties are change and may be useful for various applications [8].

In nanoscience and nanotechnology, a number of considers such as preparation method, preparative conditions and dopent are affected on ferrite properties. Structural parameters and magnetic properties of the material improved through changing microstructure, controlling the chemical composition and size [9-10]. Hence many researchers are paying attention on the preparation of nano size magnetic material with high purity, ultra-fine size, good dispersion and remarkable magnetism. In the previous period,

The Ceramic method [11] is commonly used to obtain nano particles, but they have disadvantages like high sintering temperature, time period is more and stoichiometry change with divalent volatilization results [12]. Hence, in recent years the wet chemical methods such as Solgel [13], Co-precipitation [14], hydrothermal [15], spray dying [16] and Citrate-gel technique [17] are used for preparation of nano scale materials because, easy preparation, require low temperature, less cost.

Among the ferrites, Ni ferrite and substituted nickel ferrites have extensive analysis due to having high frequency function of magnetic materials [18]. These materials scientifically interest because of its promising and interesting applications in microwave devices, colour imaging, magnetic refrigerators and high density recording devices [19]. Particularly trivalent such as AI^{+3} , Cr^{+3} are substituted in it, as for its fascinating effect on electromagnetic properties of nickel ferrite. In view of immense importance of nickel nano ferrite applications and their considerable changes in their properties by doping with trivalent ion decided to prepare chromium dopent nickel nano size ferrite with chemical formula NiCr_xFe_{2-x}O₄ (X=0.1, 0.3, 0.5, 0.7, 0.9 and 1.0) with using Citrate gel auto combustion technique, which has not been reported earlier and to study their structural and magnetic properties in systematic manner.

2. EXPERIMENTAL PROCEDURE

The molar quantity of AR grade of metal nitrates such as Nickel Nitrate, Chromium Nitrate, Ferric Nitrate, Citric acid and Ammonia as raw materials. Calculated quantities of metal nitrates by sensitive digital balance and they were dissolved in deionized double distilled water and then mixed together in a beaker. Aqueous solution of calculated quantity of citric acid was added in 1:3 molar ratio of nitrate to citric acid as chelating agent to this nitrate mixture. This mixture was thoroughly stirred using magnetic stirrer to get a homogeneous solution. Ammonia solution was added to this nitrate-citrate mixture to adjust the p^{H} to 7. The mixed solution was then heated at about 80^{0} C with uniform stirring on a hot plate to obtain a highly viscous gel

denoted as citrate precursor. The resultant gel was further heated on the hot plate maintained at a temperature of 180°C to 200°C. Finally, water molecules were removed from the mixture, the viscous gel then began frothing. The gel gave a fast flameless auto combustion reaction with the evolution of large amounts of gaseous products. It started in the hottest zones of the beaker and propagated from the bottom to the top, like the eruption of a Volcano. The reaction was completed in a minute, giving rise to dark grey voluminous product with a structure similar to a branched tree. Finally the as-burnt ferrite powders were grained by using Agate Mortar and Pestle then calcined in a muffle furnace at a temperature at 700°C for 5hr. the calcined ferrite powders were again grained by using Agate Mortar and Pestle to obtain a better crystallization and homogeneous cation distribution in the spinel. By adopting this procedure, mixed Ni-Cr nano ferrites are prepared.

The structural characterization was determined by a Bruker D8 Advance X-ray diffractometer with Cu K α radiation ($\lambda = 0.15405$ nm) source between the Bragg angles 20^o to 80^o in steps of 0.04^o/Sec. The prepared samples crystallite sizes were measured from broadening peak (311), using the Scherer's equation [20].

$$D_{hkl} = \frac{0.91\lambda}{\beta Cos\theta} - - - - (1)$$

X-ray density (d_x) measured with the following relation [21]

$$d_x = \frac{8M}{Na^3} \left(\frac{g}{cm^3}\right) - \dots - \dots - (2)$$

The bulk density d_m was determined using formula [22]

$$d_m = \frac{m}{\pi r^2 h} - \dots - \dots - \dots - \dots - \dots - (3)$$

Porosity (P) of the ferrite was determined using formula [23]

$$p = 1 - \frac{d_m}{d_x} - \dots - \dots - \dots - (4)$$

The calcined powders morphology and average crystallite size were characterized by TEM (Tecnai-12, FEI, Netherlands).

The magnetic properties were carryout from obtained M-H loops at room temperature by using VSM (GMW Magnet System, model 3473). From M-H loops saturation magnetization (M_s) directly extracted and the magnetic moment and Y-K angle measurements are calculated as follows.

Magnetic moment (μ_B) calculated with the following relation [24]

$$\mu_B = \frac{M_W X M_S}{5585} - - - - - - - - (5)$$

Where M_W – Composition molecular weight M_S - Saturation magnetization

Yefet-Kittel (Y-K) angles are calculated with the following relation [25]

$$\mu_B = (6+X)coscos_{Y-K} - 5(1-X) - - - - - - - (6)$$

Where
$$X - Cu^{2+}$$
 concentration.

The Curie temperature calculated from Loria-Sinha method (gravity method).

3. RESULTS AND DISCUSSION

3.1 X-rd analysis

Figure 1 illustrates the X-RD pattern of mixed Ni-Cr nano ferrite system, it shows the crystalline phases were identified with standard data PDF# 862267 for Nickel ferrites (NiFe₂O₄) from the ICDD data. In XRD pattern the highest reflection comes from (311) peak that shows spinel structure and all samples represents formation of cubic spinel structure in single phase without other noticeable additional impurity phases for chromium substituted nickel nano ferrite [26].

The crystallite size was computed from the highest peak (311) FWHM value and is given in **table 1**. It shows that the prepared samples crystallite size is in the nanometer scale between 8.55nm-10.36nm. To my knowledge small size mixed Ni-Cr nano ferrites samples are possible only with the Citrate-gel auto combustion method no other method has resulted the such a small size nano ferrites.

It was found that **figure 2** shows increases the X-ray density (d_x) from 5.326gram/cc to 5.368gram/cc and the bulk density (d_m) decreases from 5.218gram/cc to 4.813gram/cc with increases Cr^{3+} ion concentration in Ni nano ferrite. It may be because of larger atomic weight and density of Fe (55.847gm/mole, 7.874gm/cm³) compare with atomic weight and density of Cr (51.996gm/mole, 7.14gm/cm³). The X-ray density is more than the apparent density due to the existence of pores which depends on the preparation state. **Figure 3** shows that porosity increase with increase Cr ion concentration in Ni nano ferrite and it shows similar behavior of X-ray density. Similar behavior observed for the Cr substitution nano ferrite system with other researcher reports [27-28].

TEM micrograph of nano ferrite is represents in **figure 4** and its point out the nanostructure environment with platelet form. The crystallite size is in nanometer and also be in agreement well with crystallite size estimated from XRD pattern analysis.

3.2 Magnetic Properties

The magnetic hysteresis loops are obtained for the entire investigated the Ni-Cr nano ferrite samples in the range $\pm 6kOe$ applied field at room temperature measured by VSM. The obtained hysteresis loops are illustrated in **figure 5.** It shows the loop area is very narrow therefore the samples present soft ferrite nature with less saturation magnetization Hence these materials are desirable for transformers [29].

Composition dependence magnetic parameters

It is observed that the Ni-Cr nano ferrites have less saturation magnetization due to the grain size is small. From table 2 the saturation magnetization (M_s) value has decreased from 4.49emu/gr to 2.97emu/gr and the magnetic moment values are decreases from $0.188\mu_B$ to $0.122\mu_{\rm B}$ with increases Cr³⁺ ion concentrations in Ni nano ferrite as evidence in figure 6 and figure 7. It is explained based on the non-collinear spin arrangement [30-31]. The B-O-B coupling interactions at the B sublattice become stronger than that of A–O–B coupling between magnetic ions at the A and B sublattice due to the presence of a small canting of the B site moment with respect to the direction of the A site moment. The B–O–B coupling leads to the random existence of the small canted structure at the B site and forms triangular configuration in the ferrite system. As a result, the magnetic moments of the Fe ions at the B site are shifted from the collinear parallel to nonparallel arrangements. Therefore, the saturated magnetization is being decreased corresponding to the magnetic moment which is also decreased. The decrease in saturation magnetization and magnetic moment is credited to greater tenancy of Cr³⁺ at B sites, therefore the materials are getting changed into soft ferrite materials. Similar report was observed in Cr substituted Co-Zn nano ferrites by S. Bhukal et al [32] and Al substituted Co-Zn nano ferrites by Sonal Singhal et al [33].

The decrease of magnetization has been proposed by Yafet and Kittel (Y–K) by triangular arrangement of spins [34]. The Y-K angles of Ni-Cr nano ferrite system are reported here in the **table 2** as evidence in the **figure 8**. It is clear that increases the Y-K angles with increase Cr^{3+} ion concentration in Ni nano ferrite. It indicates that the spin canting takes place significantly at higher concentration of Cr content. Therefore, increases the spin arrangement at B-site. As a result, decrease A-B exchange interaction consequent decreases in magnetization.

It was found a decrease from 775.39 Kelvin to 635.34 Kelvin on increasing the Cr^{3+} concentration, because the Fe³⁺ ions are replaced by paramagnetic Cr^{3+} ions [35]. The Curie temperature is determined by an overall strength of the exchange interaction between A and B sublattice. When the replacement of Fe³⁺ ion with Cr^{3+} ions increases, the magnetization decreases in B-sublattice without disturbing the A-sublattice, this leads to a decrease in the A-B interactions (Fe³⁺-O²⁻-Fe³⁺) which results in decreases of Curie temperature. Similar behavior was observed in the trivalent substitution in nano ferrite system investigated by other researchers [36-37].

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4. CONCLUSION

The Citrate-gel auto combustion technique is to be a convenient and versatile for obtaining homogeneous nanostructured mixed Ni-Cr ferrites. X-ray diffraction confirms the formation of single phased cubic spinel structure without any impurity peak. X-ray density increases, bulk density decreases and porosity increases with increase in Cr substitution in the mixed Ni-Cr nano ferrite system. It shows the densification of the material. The obtained hysteresis loops of Ni-Cr nano ferrites illustrate soft magnetic materials with less loop area and small saturation magnetization, hence these materials are may be desirable for transformers to minimize the energy dissipation.saturation magnetization, magnetic moment are decreases and the Y–K angles increases with increase Cr^{3+} ion concentrations in Ni nano ferrites. In Ni-Cr nano ferrites it was decreases from 775.39K to 635.34K with increase in Cr^{3+} concentration.

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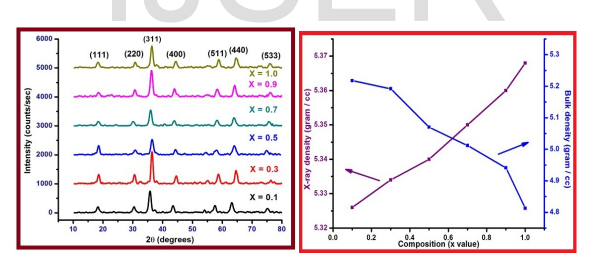
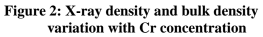


Figure 1: X-ray diffraction pattren of mixed NiCr_XFe_{2-X}O₄ nano ferrites



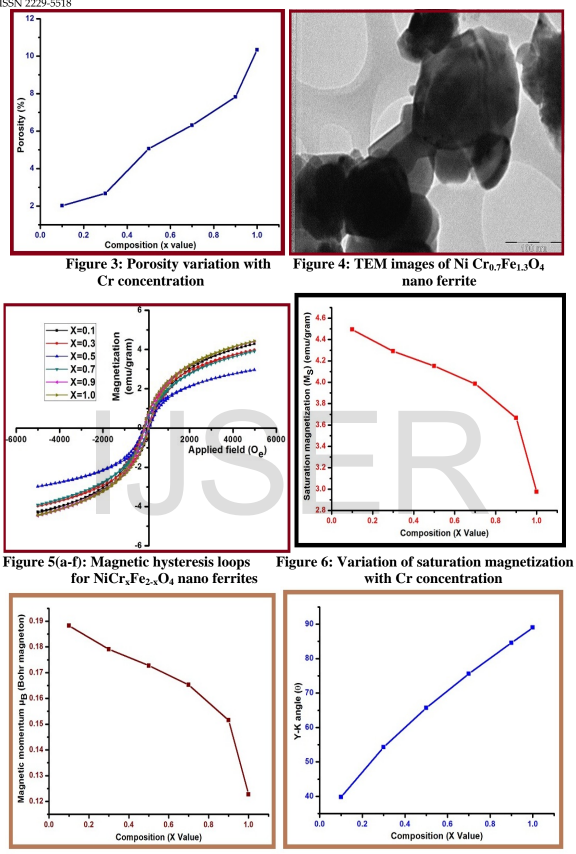


Figure 7: Variation of magnetic moment with Cr concentration

Figure 8: Variation of Y-K angle with Cr concentration

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Sl.No.	Composition	Molecular	Crystallite	X-ray	Bulk	Porosity
		weight	size (nm)	density	Density	(P%)
		(M)(gr)		(gram/cc)	(gram/cc)	
1	$NiCr_{0.1}Fe_{1.9}O_4$	233.945	8.96	5.326	5.218	2.01937
2	NiCr _{0.3} Fe _{1.7} O ₄	233.175	10.36	5.334	5.192	2.67538
3	NiCr _{0.5} Fe _{1.5} O ₄	232.340	7.95	5.34	5.07	5.0628
4	NiCr _{0.7} Fe _{1.3} O ₄	231.635	8.55	5.35	5.012	6.31649
5	$NiCr_{0.9}Fe_{1.1}O_4$	230.865	8.84	5.36	4.941	7.82673
6	NiCrFeO ₄	230.480	9.26	5.368	4.813	10.33693

Table - 1: Structural parameters of mixed NiCr_xFe_{2-x}O₄ nano ferrites

Table - 2: Magnetic parameters of mixed Ni-Cr nano ferrites

Sl.No.	Composition	Saturation magnetization (M _S) (emu/gr)	Magnetic moment (Bohr magneton)	Y-K angle (θ)	Curie Temperature
1	$NiCr_{0.1}Fe_{1.9}O_4$	4.49	0.18823	39.7751	808
2	NiCr _{0.3} Fe _{1.7} O ₄	4.29	0.17911	54.2687	736
3	$NiCr_{0.5}Fe_{1.5}O_4$	4.15	0.17273	65.7204	726
4	$NiCr_{0.7}Fe_{1.3}O_4$	3.99	0.16529	75.6083	686
5	$NiCr_{0.9}Fe_{1.1}O_4$	3.67	0.15154	84.5817	674
6	NiCrFeO ₄	2.97	0.12272	88.9955	668