Original Research Article

Effect of temperature on Structural, Magnetic and Dielectric Properties of Cobalt ferrite Nanoparticles Prepared via Co-precipitation Method

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7 Abstract

Nano-crystalline cobalt ferrite powders have been synthesized by coprecipitation method and 8 9 characterized to investigate the effect of thermal treament on the structural, dielectric, and magnetic properties of the prepared sample. The structural/morphological, dielectric, and magnetic properties of 10 the products were determined by X-raydiffraction (XRD), transmission electron microscopy (TEM), 11 open ended coaxial probe and vibrating sample magnetometer (VSM). Results from analysis confirms 12 that thermal heat influences the magnitude of magnetization, dielectric and surface morhpology of 13 samples. The variation of dielectric properties with frequency reveals that the variation in magnitude is 14 due to Maxwell–Wagner type of interfacial polarization as well as hopping of charge from Fe²⁺ to Fe³⁺ 15 and from Co^{2+} to Co^{3+} ions at B-sites. 16

17 Keywords: nanoparticles, ferrites, spinel, cobalt ferrite, permittivity

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19 Introduction

Magnetic nanostructured materials have gained great interest of the scientists due to their unusual 20 physical properties compared to their bulk form. They have very large areas of applications in 21 technology, such as electronic devices, microwave devices, transformer cores, magnetic devices, 22 switching devices, recording tapes, permanent magnets, hard disc recording media, flexible recording 23 media, read-write heads, active components of ferrofluids, magnetic drug delivery, catalysis, color 24 imaging, magnetic refrigeration, detoxification of biological fluids, magneti- cally controlled transport 25 TDR.E.MOVE SPAG of anti-cancer drugs, magnetic resonance imaging (MRI) contrast enhancement and mag- netic cell 26

27 separation [1].

The ferrite nanoparticles have a spinel structure and the general spinel structure is in the form as AB₂O₄. Cobalt ferrite is a partially inverted spinel structure with cobalt atoms predominantly in the octahedral sites [2] and it is a well-known hard magnetic material, which has been studied in detail due to its high coercivity (5400 Oe), high chemical stability, good electrical insulation, significant mechanical hardness and moderate saturation magnetization (80 emu/g) at room temperature.

33 Nanoferrites are simultaneously good magnetic and dielectric materials. These properties of the ferrites are governed by the choice of the cations and their distribution between tetrahedral and octahedral sites 34 of the spinel lattice. The properties of the nanoferrites are also affected by the preparation conditions, 35 chemical composition, sintering temperature, doping additives, and the method of preparation [3]. 36 Several chemical and physical methods such as spray pyrolysis, sol-gel, coprecipitation, combustion 37 technique, high energy milling, and so forth have been used for the fabrication of stoichiometric and 38 chemically pure nanoferrite materials [4]. Among the available synthesis methods, solgel method has 39 attracted much attention due to its inherent advantages of low processing temperature and homogenous 40 reactant distribution. The products obtained by this method exhibit high crystalline quality, narrow size 41 LADINSERT SPACE distribution, and uniform shape [5]. The effect of rare-earth ions inclusion into the ferrite spinel 42 structure has been reported in many literatures, however little has been reported on the effect of 43 calcination temperature on dielectric and magnetic properties. Rashad et al, [6] indicated the change in 44 themagnetic properties of samariumsubstitutedCoFe2O4 synthesized by citrate precursor method and 45 9 COFE, OG-DSUB WIZX LOSPACE the results revealed that the saturation magnetization and coercivity are decreased with the addition of 46 Sm³⁺ ions. Peng et al. [7] have reported an increase in crystallite size of cobalt ferrite nanoparticles by 47 the doping of gadolinium. Guo et al, [8] have reported that the substitution of Sm3+ in NiFe₂O₄ 48 increases the lattice parameter and reduces the crystallite size of the materials. Tahar et al, [9] have 49 investigated the effect of Sm³⁺ and Gd³⁺ substitution on the magnetic properties of cobalt ferrite 50

synthesized by forced hydrolysis in polyol and reported that particle size increased slightly with rare
 earth substitution.

In this work we investigated the effect of temperature on structural, magnetic, and dielectric properties
 of CoFe₂O₄ nanoparticles synthetized via coprecipitation method.

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56 **Experimental**

57 Synthesis

Required amount of iron nitrate and cobalt nitrate were dissolved in 100 ml of distilled water at room temperature and stirred for 15 min continously. During constant stirring, 100 ml of ammonia solution was added in drops form. The droppings were done so as to get a solution with pH about 12. The mixture solution/precipitate were dried in an oven at 80 - 100 ^oC for 48 hrs. The obtained dry powder were then calcined at in a furnace at 600 ^oC and 900 ^oC for 2 hrs. The annealed samples were allowed to cool naturally to room temperature.

64 *Measurement*

X-ray diffraction (XRD) of the prepared sample were recorded in a Phillips diffractometer equipped 65 with a CuK α monochromator. The indentification of the crystalline phases present in the samples were 66 achieved using the XRD profiles. Transmission Electron Microscope (TEM) JEOL JEM 1220 was used 67 to explore the sample morphology and its particle size distribution. The room temperature infrared 68 spectra of nanopowders were recorded using a Fourier Transform Infrared Spectrophotometer Perkin 69 Elmer. The dispersing of the samples were observed in the range from 380 to 4000 cm⁻¹. The 70 temperature and field dependence of magnetization were carried out in vibrating sample magnetometer 71 (VSM). While dielectric measurement were carried out using open ended coaxial probe via Agilent HP 72 85072. 73

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75 **Results and discussion**

76 XRD analysis

77 X- ray diffraction study of both samples confirmed the formation of single phase cubic spinel structure.

- The single phase spinel ferrite obtained by high temperature calcination at 900 0 C is in agreement with
- the standard JCPDS data reference (JCPDS: 22-1086).
- 80 The micropgraph of the $CoFe_2O_4$ sintered sample at 600 ^{0}C and 900 ^{0}C are presented in Fig. 1. The
- 81 average crystallite sizes were calculated by Scherer's equation (Eq.(1)) from the intensity of reflection
- of the spinel structure's of the (311) plane. The equation of Scherrer is given as [10];

$$D = \frac{k\lambda}{\beta \cos\theta} \tag{1}$$

where *k* is a constant equal to 0.9, λ is the wavelength of the X-ray radiation (all diffraction patterns shown in this paper were performed with CuKa radiation), θ is the diffraction angle, and β is the full width half maximum (FWHM).



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The lattice parameters have been computed using the d-spacing values and the respective (hkl) parameters from the classical formula given in Eq.(2) [10].

$$a = \frac{\lambda [h^2 + k^2 + l^2]^{1/2}}{2sin\theta}$$
(2)

Calculation showed that the crystallite size and lattice parameter of the powder were found to 49.8nm and 8.343 A for 600 °C and 67.7 nm and 8.362 A for the 900 °C sintered sample. This finding indicated that the synthesized powders had nano size crystallites and that the sintering temperature caused the sample to increase in the crystallite size. In addition, the sintering tempearture also led to a slight increase of 0.019 A in the lattice parameter. This is attributted to the heat coupling in the sample at higher temperature allowing the diffusion of metals into their octahedral sites [10].

The sharp increase in ctrystallinity of the sample is attributed to calcination temerature as observed for the 900 0 C heating.

99 **TEM measurement**

Figure 2 shows the TEM images of the ptreparred Cobalt ferrite nanoparticles after annealing with 100 temperatures of 600 ⁰C and 900 ⁰C respectively. The particles in both cases are spherical and exhibit a 101 homogeneous distribution, especially at higher lower temperature. The average sizes of the 102 nanoparticles were found to be 43.5 nm and 62.5 nm for the 600 ^oC and 900 ^oC respectively. The result 103 obtained for the average particle size distribution were in agreement with the values obtained using the 104 XRD spectra. An increase of particle size with increasing annealing indicated that when temperature 105 increases, more metal ions are produce which in turn increases the particle sizes. This increment in 106 number of ions and particle size is supports the enhancement of the intensity of absorption as shown in 107 the FTIR spectra for the 900 ⁰C annealed sample. It important to note that tiny amount of 108 agglomeration is observed at higher temperature due to growing distribution of particle size. A 109 summary of properties of the prepared sample is tabulated in Table 1. 110





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Figure 2: TEM images of cobalt ferrite nanoparticles calcined at 600 °C and 900 °C

Table 1. Magnetic parameters of cobalt ferrite						
Compound	T _{calc} (⁰ C)	a (Å)	<i>d</i> (nm)	H _C	M_r	Ms
				(Oe)	(<i>µ_{<i>B</i>}/mol)</i>	(<i>µ^{<i>B</i>}/mol)</i>
				12,405	2,819	3,349
CoFe ₂ O ₄	600	8.343	43.5	696	937	2,987
	900	8.362	62.5	10,640	3,080	3,569
				720	1,048	3,294

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116 FTIR Spectroscopy

117 Careful observation on the FTIR spectra shown in figure 3 shows an absorption peak at 592cm^{-1} 118 corresponding to intrinsic stretching vibrations of metal at the tetrahedral site, whereas the v2-lowest 119 band, observed in the range of 400–300 cm⁻¹, is due to the octahedral–metal stretching [11]. Co²⁺ ions 120 usually occupy octahedral-site, while Fe³⁺ ions has the tendency to occupy both octahedral and 121 tetrahedral sites [12]. Further observations shows strong absortions peaks at 3414 and 1617 cm⁻¹ 122 wavelengths.



Figure 3: FTIR spectra of cobalt ferrite nanoparticles calcined at (a) 600 °C and (b) 900 °C Magnetic measurement

Figure 4, represents the ZFC and FC curves of magnetization of both temperatures. Both ZFC and FC 127 magnetization decreases by increasing the temperature. At low temperatures in the presence of a 128 magnetic field (FC), the magnetization direction of each particle is frozen in the field direction. While 129 the ZFC magnetization is below the magnitude for FC. Both ZFC and FC magnetization is increasing 130 by increasing the temperature. However, at higher calcination temperature, the magnetization in both 131 cases is lower than that compared at low temperature. In both cases, the ZFC and FC curves are 132 seperated given an indication that there is a non-equilibrium magnetization below the separation 133 temperature. For the ZFC case, the separation represents the irreversibility temperature, (T_{irr}) . 134

In general, T, represents the blocking–unblocking process of the particles magnetic moment when thermal energy is changed [13]. As the relaxation time of a magnetic particle decreases with increasing temperature, below a certain characteristic temperature called the blocking temperature (T_B) the particles moment remains blocked with respect to the time scale of the experiment. The difference between T_B and T_{irr} corresponds to the width of the blocking temperature distribution which gives an indication of the narrowness of the anisotrop yenergy barrier distribution of the particles [13,14]. Below the Tirr, the ZFC and FC curves significantly diverge and the sample is in the ferromagnetic

state. The divergence in the ZFC–FC magnetization curves below T_B is attributed to the existence of magnetic anisotropy barriers [14].





Figure 4: Temperature dependent magnetizations of sample

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This behavior is confirmed by the isothermal magnetization curves obtained at 5 K and 320 K shown in Fig. 5. These results are in agreement with analysis obtained for the ZFC/FC discussed in Figure 4. The saturation temperature, coercive was highest at 5 K for both annealing tempeartures as shown in Table 1. Whilst the magnetic hysteris at 5K showed very large hysteresis with coercive fields of 12405 and 10640 Oe for 600 0 C and 900 0 C annealing respectively.



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Figure 5: Hysteresis loops of the samples at (a) 600 ⁰C and (b) 900 ⁰C

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Further observation showed that in Fig. 5 the coercitivity of the nanoparticles decreases as the 155 temperature increases toward the blocking temperature. The inserts in Fig. 5 show the part of the M(H) 156 loops in the region of small applied field. The saturation magnetization is dependent on the thermal 157 treatment where saturation magnetization obtained were, $M_s = 3.349 \mu_B/mol$ for sample calcined at 600 158 ${}^{0}C$ and M_S= 3.569 μ_{B} /mol for sample calcined at 900 ${}^{0}C$ at 5K. In order to verify the obtained 159 inversion degree in the cobalt ferrite prepared in this work, we used The highest saturation 160 magnetization: $M_s = 3.569 \mu_B/mol$ was used to verify inversion degree obtained in this work. The 161 inversion degree obtained was $\lambda = 0.85$. 162

163 Dielectric measurement

Open ended coaxial probe (Agilent 85702B) was used in the measurement of complex permittivity of 164 both samples. The measurement was carried out at microwave frequency (8.2 GHz to 12.4 GHz). 165 Careful observation on Figure 6(a) shows that dielectric constant decreases with the increase of 166 frequency in both samples. The dependence of dielectric constant on frequency can be explained using 167 interfacial polarization as predicted by Maxwell-Wagner [15]. It is expected that the dielectric structure 168 of a ferrite is made up of a conductive layer that consists of large ferrite grains and other grain 169 boundaries that are poor conductors. Polarization is mainly due to electron exchange between Fe⁺² and 170 Fe⁺³ in the direction of applied field. Measurement result showed that the mean directly constant were 171 4.46 and 5.05 for the 600 ^oC and 900 ^oC annealed samples respectively. 172





Figure 6: Complex permittivity of samples annealed at 600 ⁰C and 900 ⁰C

As shown in figure 6(b), the large value of loss factor at lower frequency might be attributed to the dominant species like Fe^{+2} ions, oxygen vacancies, grain boundary defects, etc. The loss factor at high frequency are due to particles which have low resistivity. The mean loss factor obtained for both samples are 0.26 and 0.29 for the 600 $^{\circ}C$ and 900 $^{\circ}C$ calcinations respectively.



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Figure 7 depicts the loss factor obtained from the ratio of loss factor to the dielectric constant. The loss tangent (tan δ) decreases with the increase of frequency and then begins to stagnate at higher frequencies for both samples. The mean magnitude of the loss tangent obtained are 0.058 and 0.057 for the 600 0 C and 900 0 C calcinations respectively. This results indicates that the 600 0 C annealed sample would absorb higher than the 900 0 C annealed sample. The loss (tan δ) depends on factors, such as

material stoichiometry, Fe^{+2} content and material dispersion, which in turn depends on the composition and synthesis methods. The decrease in loss tangent with frequency may be attributed to the Maxwell-Wagner polarization and conduction mechanism in ferrites [16].

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190 Conclusions

The thermal heat caused an increase in the average crystallite size, magnetization and the coercive field 191 of the sample. Similarly, infrared spectroscopy studies confirms the presence of metal oxide in the 192 prepared nanoparticles. The main characteristic of the field cooled (FC) and zero field cooled (ZFC) 193 curves is its irreversibility below room temperature suggesting the presence of a superparamagnetic 194 behavior. In addition, the magnetic properties exhibit some dependence on the calcination temperature. 195 The M_s values increases with the increase of calcination temperature whilst the maximum value of H_c 196 is about 12405 Oe which is much higher than the coercitivity of bulk material. The increase in the 197 value of lattice parameter with increase in temperature suggests the expansion of the unit cell. 198 Crystallinity and the crystallite size are observed to increase with increase in temperature. The average 199 particle sizes estimated from TEM micrographs were 43.5 nm for sample calcined at 600 °C and 62.5 200 nm for sample calcined at 900 0 C. 201

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