

Magneto-structural properties of Ni–Zn nanoferrites synthesized by the low-temperature auto-combustion method

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Abstract. Using nickel, zinc and ferric nitrates, and glycine in a fuel-rich composition, Ni_{1-x}Zn_xFe₂O₄ nanoparticles were prepared by a simple low-temperature auto-combustion method without further sintering at high temperatures. The auto-combusted powders obtained were characterized by X-ray diffraction (XRD), Raman spectroscopy, scanning electron microscopy, energy-dispersive X-ray (EDAX) analysis and vibrating scanning magnetometer measurements. XRD confirms the formation of pure nanocrystalline spinel phases with an average diameter of about 55 nm. Raman spectra show tetrahedral and octahedral sites in the structure of Ni_{1-x}Zn_xFe₂O₄ and also imply the doping of Zn²⁺ and displacement of Fe³⁺ ions from the tetrahedral site. EDAX showed that the samples were close to the nominal compositions. The magnetic measurement shows that the saturation magnetization and remanence magnetization decreases with the increase in the zinc content.

Keywords. Ni–Zn ferrites; fuel-rich composition; combustion synthesis; Raman studies; magnetism.

1. Introduction

Nanocrystalline spinel ferrites with the common formula AFe₂O₄ (A = divalent metal ion, e.g., Ni, Zn, Mn, Co, Mg, Cu, etc.) are the most significant magnetic materials.¹ In recent years, this class of magnetic nanomaterials has elicited many interests because of their fascinating electronic, magnetic, catalytic and biomedical applications. The interest for using these materials permanently increases because of their usability under extreme conditions.² Nickel ferrite has been intensively investigated as one of the magnetic nanomaterials.^{3–6} Magnetic properties of ferrites can be changed by substituting various kinds of A²⁺ (Zn²⁺, Mg²⁺, Mn²⁺, Ni²⁺, Co²⁺, Fe²⁺, etc.) among divalent cations by introducing a relatively small amount of transition metal ions.⁷ The substitution of metal ions like Zn²⁺ into nickel ferrite has been proposed by many researchers to modify the electrical and magnetic properties. In particular, extensive studies of Zn-doped NiFe₂O₄ have been conducted, which showed that the substitution of Zn for Fe decreases the Curie temperature and magnetic anisotropy^{8–11} and the substitution of Zn for Ni affects the electrical properties.^{9,11,12}

Nickel–zinc ferrites are magnetic materials of much technological importance due to their high electrical resistivity, low magnetic coercivity and low eddy current losses. These properties depend upon the composition, microstructure and heat treatment of the samples.¹³ It is well known that properties of ferrite materials strongly depend on the preparation conditions. Many methods such as the citric acid combustion method,^{9,10,11,14} urea-assisted auto-combustion method,¹⁵ reverse micelles,⁸ hydrothermal method,¹⁶ co-precipitation method,¹⁷ solid-state method,¹⁸ etc., have been developed to prepare nanocrystalline nickel–zinc ferrite. The spinel ferrite particles synthesized by solid-state methods show an assembly of irregular shapes and agglomerations,¹⁹ while those prepared by most wet chemical methods require careful control of pH of the solution, concentration like parameters and high sintering temperature for the formation of particles. High-temperature synthesis of nickel–zinc ferrite may result in the evaporation of some of the constituents resulting in non-stoichiometry, and zinc volatilization and increased sintering temperatures can result in the formation of Fe²⁺ ions, thereby increasing the electron hopping and reducing the resistivity.¹¹ Therefore, a method that requires low-temperature synthesis will be most suitable for the synthesis of nickel–zinc ferrite. Among the various methods for synthesizing ferrites, the combustion method stands out as an alternative and highly promising method.²⁰ This method has additional

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advantages of simple preparation, formation of products with virtually any size or shape, and formation of high-purity homogeneous products. In combustion synthesis, the nature of fuel and fuel to oxidizer (metal nitrates) ratio can be used to tune the morphology, phase and magnetic properties of the final product.

In most auto-combustion processes, glycine as a fuel is preferred because of its high negative combustion heat ($-3.24 \text{ kcal g}^{-1}$) as compared to urea ($-2.98 \text{ kcal g}^{-1}$) and citric acid ($-2.76 \text{ kcal g}^{-1}$); hence, high sintering temperature is usually not required unlike with the other fuels. Also, glycine is readily available, economic and highly soluble in water. In our previous study on the combustion synthesis of cobalt–magnesium ferrite using different glycine metal nitrate ratio, a fuel-rich composition was found to produce the purest nanocrystalline ferrite with the highest saturation magnetization with no further sintering temperature compared with the fuel lean and fuel stoichiometric samples. Therefore, in the synthesis of the different compositions of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ ($x = 0.65, 0.7$ and 0.75) using glycine as a fuel, fuel-rich composition is used ($G/N = 2.22$). X-ray diffraction (XRD), Raman spectroscopy, scanning electron microscopy (SEM), energy-dispersive X-ray (EDAX) analysis and vibrating scanning magnetometer (VSM) techniques are used to study the structural, morphological, chemical composition and magnetic properties of as-synthesized Ni–Zn ferrite MNPs, respectively.

2. Experimental

2.1 Materials

All the reagents are of analytical grade and are used as-received without further purification. Nickel nitrate [$\text{Ni}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$], zinc nitrate [$\text{Zn}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$] and iron nitrate [$\text{Fe}(\text{NO}_3)_3 \cdot 9\text{H}_2\text{O}$] obtained from Sigma Aldrich, Germany, are taken as oxidants, while glycine (G, $\text{C}_2\text{H}_5\text{NO}_2$) obtained from SD Fine Chem. Ltd., Mumbai, was employed as fuel to drive the combustion process.

2.2 Synthesis

The $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ ($x = 0.65, 0.7$ and 0.75) ferrite is prepared by the low-temperature auto-combustion method. The amount of fuel used was calculated to be more than the stoichiometric amounts required for completion of the combustion process without heat exchange. Here, the glycine to nitrate ratio was $G/N = 2.22$ (fuel-rich composition). For $x = 0.65$ sample, 1.02 g $\text{Ni}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$, 1.93 g $\text{Zn}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$, 8.08 g $\text{Fe}(\text{NO}_3)_3 \cdot 9\text{H}_2\text{O}$ and 5.0 g glycine were dissolved in 20 ml of distilled water and the solutions were heated to 80°C to form a viscous gel of precursors under magnetic stirring. Secondly, the gel is transferred to a pre-heated coil (300°C). Finally, after a short moment, the solution precursors boiled, swelled,

evolved a large amount of gases and ignited, followed by the yielding of puffy black products. The powder (auto-combustion powder) was heated in a hot air oven at 200°C for about 12 h to remove any organic product or unreacted glycine. Similarly, for $x = 0.7$ sample, same procedures were followed except that 0.87 g $\text{Ni}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ and 2.08 g $\text{Zn}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ were used as precursors, while for $x = 0.75$ samples, 0.73 g of $\text{Ni}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ and 2.23 g $\text{Zn}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ were used.

2.3 Characterization methods

The X-ray diffractograms of the auto-combustion powders were recorded using an X-ray diffractometer (D8 Advance, Bruker, Germany), equipped with a $\text{CuK}\alpha$ radiation source ($\lambda = 1.5406 \text{ \AA}$) and the crystallite size (D) is calculated from X-ray line broadening of the (311) diffraction peak using the well-known Scherrer relation

$$D = \frac{0.9\lambda}{\beta \cos\theta}, \quad (1)$$

where β is the full-width at half-maxima of the strongest intensity diffraction peak (311), λ the wavelength of the radiation, and θ the angle of the strongest characteristic peak. X-ray density (D_X) was calculated using

$$D_X = \frac{8M}{Na^3}, \quad (2)$$

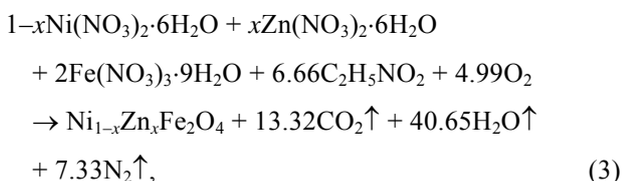
where M is the molecular weight, N Avogadro's number and a the lattice constant. The surface morphology and elemental detection of the powders were examined with a scanning electron microscope (SEM), Ametek model XL30 Lab 6. The Raman spectra were obtained using a LabRAM HR (Olympus BX41) visible single spectrometer equipped by a microscope and a Peltier-cooled CCD detector. The 633 nm He–Ne laser line was used for excitation. The power was adjusted using a set of neutral filters. The spectral slit width at the conditions used was 1 cm^{-1} . The laser beam was focused on the powders pressed on a glass surface by an $\times 50$ long working distance objective. The acquisition time for all samples was 60 s ($\times 2$ – times per scan) for all optical excitation intensities. The Raman measurements were performed at room temperature and atmospheric pressure. The magnetic characterizations were carried out with a VSM (Lake Shore cryotronics-7400 series) under the applied field of $\pm 20,000 \text{ G}$ at room temperature.

3. Results and discussion

3.1 Combustion reaction

In order to synthesize crystalline $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ MNPs with no further heat treatment, to avoid sintering of the

MNPs, the enthalpy or flame temperature was increased by using glycine with a fuel-rich composition. The possible combustion reaction is



where $x = 0.65, 0.7$ and 0.75 . The precursor mixtures resulted in a dark red solution and the combustion types were flamy combustion. Combusted products resulted in brown powders with some black patches dotting it. This might be as a result of the unreacted glycine being a fuel-rich composition. After heating in the oven at 200°C for 12 h, brown powder resulted with no black patches which were grounded and used for further characterizations.

3.2 Phase formation and structural analysis

The powder X-ray patterns recorded for the samples of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ are shown in figure 1. They are all consistent with the standard pattern cubic spinel structure of bulk NiFe_2O_4 (JCPDS card no. 10-0325) and traces of secondary haematite ($\alpha\text{-Fe}_2\text{O}_3$) phase were observed. Secondary haematite phases were also observed in the urea-assisted combustion synthesis of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ ²¹ and sintering temperature of 1200°C was required to obtain good crystalline phase and remove the haematite phase from the urea-assisted combustion synthesis of

$\text{Ni}_{0.5}\text{Zn}_{0.5}\text{Fe}_2\text{O}_4$ ²² and the citrate-assisted combustion synthesis of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$.^{9,10} The diffractogram exhibits sharp lines, which indicates that the sample has high crystallinity. This results show that using the glycine fuel-rich composition in the solution combustion synthesis of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ without any further heat treatment at high temperatures is sufficient in the formation of the spinel ferrite phase. The X-ray patterns of $\text{Ni}_{0.3}\text{Zn}_{0.7}\text{Fe}_2\text{O}_4$ (figure 1b) recorded the sharpest and most intense XRD reflections compared with the other two diffraction patterns, indicating the highest crystallinity, while the X-ray patterns of $\text{Ni}_{0.35}\text{Zn}_{0.65}\text{Fe}_2\text{O}_4$ (figure 1a) recorded the lowest. The estimated values of various structural properties of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ MNPs are given in table 1. From Scherrer's formula, it was found that all the samples obtained are nanocrystalline with sizes ranging between 54 and 56 nm. Lattice constant decreased in $\text{Ni}_{0.3}\text{Zn}_{0.7}\text{Fe}_2\text{O}_4$ and increased in $\text{Ni}_{0.25}\text{Zn}_{0.75}\text{Fe}_2\text{O}_4$. The unit cell volume which is linearly affected by the lattice constant expectedly followed the same trend. In $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$, the smaller ionic radius of Ni^{2+} (0.69 Å) and Fe^{3+} (0.60 Å) ions are replaced by the larger Zn^{2+} (0.74 Å) ions with all the ionic radii having a coordination number of six.²³ Hence, it was expected that the lattice constant would increase because of expansion of unit cell dimension with the increase in the amounts of Zn^{2+} in obedience to Vegard's Law. However, the X-ray density (D_X) value was lowest in $\text{Ni}_{0.35}\text{Zn}_{0.65}\text{Fe}_2\text{O}_4$ (the sample with the least Zn content) which was in line with this calculation. The calculated D_X value for $\text{Ni}_{0.3}\text{Zn}_{0.7}\text{Fe}_2\text{O}_4$ sample (with the lowest lattice constant value) was the highest of all samples. This is also in line with the formula for calculating D_X (equation 2), which shows an inverse relationship between D_X and lattice constant. The same inverse variation of D_X with lattice constant has also been recorded in $\text{Co}_{1-x}\text{Mn}_x\text{Fe}_2\text{O}_4$ system.⁷

3.3 Raman studies

Room temperature Raman spectra of auto-combusted powders were recorded in the range of $0\text{--}1000\text{ cm}^{-1}$, as shown in figure 2. The popularity of Raman spectroscopy in the investigation of oxides is increasing because of its utility to probe local disorder.²⁴ The short-range disorder in oxygen octahedra induced by Jahn-Teller distortion and other interactions can be effectively probed by Raman spectroscopy, which makes it a very versatile tool.²⁵ The Raman features are assigned to the vibrational modes from the nanoparticles crystalline structure. NiFe_2O_4 crystallizes with inverse spinel structure,²⁶ described by the face-centred cubic (FCC) space group $\text{Fd-}3\text{m}$ (no. 227, $Z = 8$). In this structure the tetrahedral A-sites (8a) are occupied by half of the Fe^{3+} cations, whereas the rest of the Fe^{3+} and Ni^{2+} cations are distributed over the octahedral B-sites (16d).²⁷ According to the space group symmetry and factor group analysis, five Raman active

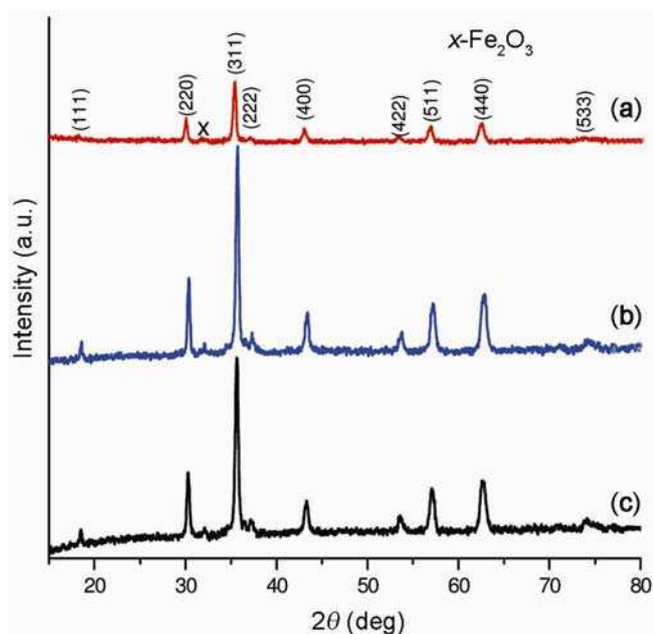
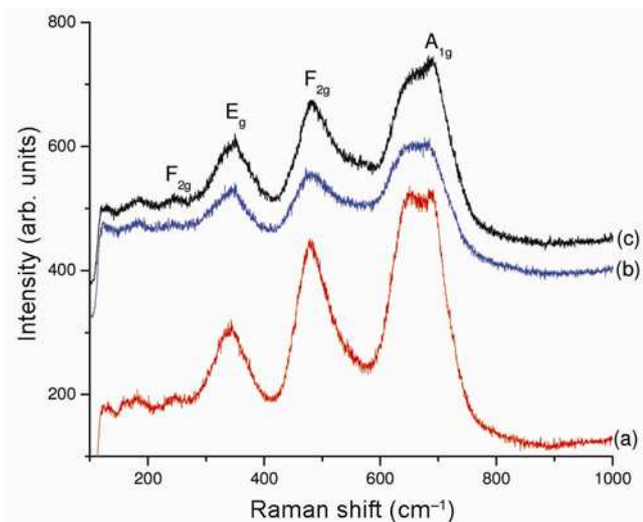


Figure 1. X-ray diffraction patterns of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ samples: (a) $x = 0.65$, (b) $x = 0.7$ and (c) $x = 0.75$.

Table 1. Structural properties of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ MNPs: (a) $x = 0.65$, (b) $x = 0.70$ and (c) $x = 0.75$.

Zn conc. 'x'	Crystallite size, D (nm)	Lattice constant, a (nm)	Unit cell volume, V (nm^3)	X-ray density, D_X (g cm^{-3})
0.65	56	0.841	0.595	5.331
0.70	55	0.833	0.578	5.495
0.75	54	0.835	0.582	5.465

**Figure 2.** Raman spectra of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ samples: (a) $x = 0.65$, (b) $x = 0.7$ and (c) $x = 0.75$.

internal modes such as A_{1g} , E_g and $3F_{2g}$ modes are predicted.²⁴ In nickel–zinc ferrite, Zn^{2+} ions are known to occupy the tetrahedral (A) site. The Zn^{2+} ion has a larger ionic radius than Ni^{2+} ion and is expected to increase the structural disorder of the oxygen sublattice. It can be seen from the Raman spectra that there is broadening of the spectra and decrease in intensity with an increase in zinc substitution. Also, all the samples gave broad A_{1g} vibrational modes. This mode has both been attributed to order–disorder effect²⁸ and the substitution of the Fe^{3+} ions by Zn^{2+} ions in the tetrahedral sites. It can be considered from the results that the vibrational modes show tetrahedral and octahedral sites in the structure of nickel–zinc ferrite with Zn^{2+} ions displacing Fe^{3+} ions from the tetrahedral site. The experimentally obtained Raman modes are consistent with those of previously reported crystalline $\text{Ni}_{0.75}\text{Zn}_{0.25}\text{Fe}_2\text{O}_4$.²⁷

3.4 Morphological and chemical composition analysis

The SEM micrographs of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ ($x = 0.65$, 0.70 and 0.75) are shown in figure 3. The images of each sample powder were taken at two different magnifications – $1000\times$ (figure 3a^I–c^I) and $30,000\times$ (figure 3a^{II}–c^{II}). It can be seen that all the samples (figure 3a^I–c^I) have

voids and pores (which decreased with the increase in the concentration of Zn^{2+}) and can be attributed to the release of large amount of gases during the combustion process. A decrease in porosity with the increase in the concentration of Zn^{2+} in Ni–Zn ferrite systems have also been reported.⁹ Densification of the powders with the increase in the concentration of Zn^{2+} can also be seen in figure 3a^I–c^I. The addition of zinc to ferrites results in the densification of the material.^{9,29} Combustion flame temperature was found to increase with the increase in the concentration of Zn^{2+} in $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ system.²¹ It was explained that the changes in the systems' compositions in response to increasing concentrations of Zn^{2+} caused the combustion flame temperature to increase, thereby favouring agglomeration and pre-sintering. This explanation can also be extended to the remarkable changes in microstructure, regarding density and porosity, observed in response to increasing concentrations of Zn^{2+} in this study. In figure 3b^{II}–c^{II}, agglomerated oval-shaped morphology was observed along with a few semi-spherulitic shapes which clearly differed from the smaller, irregular, and agglomerated morphology seen in figure 3a^{II}. This might be interpreted as the formation of a poorer crystalline ferrite phase of $\text{Ni}_{0.35}\text{Zn}_{0.65}\text{Fe}_2\text{O}_4$, corroborating the XRD results which recorded the lowest intensity XRD peaks compared with other samples.

The compositional analyses of the nanocrystalline $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ ($x = 0.65$, 0.70 and 0.75) samples were carried out by EDAX and they are shown in figure 4a–c. From the EDAX results, the presences of Ni, Zn, Fe and O in the samples were confirmed with no impurity present and the compositional molar ratio of Ni and Zn to Fe were found to be close to 0.5 (table 2).

3.5 Magnetic studies

Figure 5 shows the magnetic hysteresis curves of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ samples recorded at room temperature. The effects of Zn doping on the magnetic properties are given in table 3. NiFe_2O_4 exhibits ferrimagnetism originating from the magnetic moment of anti-parallel spins between Fe^{3+} at the tetrahedral sites and Ni^{2+} at the octahedral sites.³⁰ The saturation magnetization (M_s) and remanence magnetization (M_r) of $\text{Ni}_{0.35}\text{Zn}_{0.65}\text{Fe}_2\text{O}_4$ sample is 36 and 22.5 emu g^{-1} , respectively, while M_s and M_r of $\text{Ni}_{0.25}\text{Zn}_{0.75}\text{Fe}_2\text{O}_4$ sample is 25 and 15 emu g^{-1} , respectively.

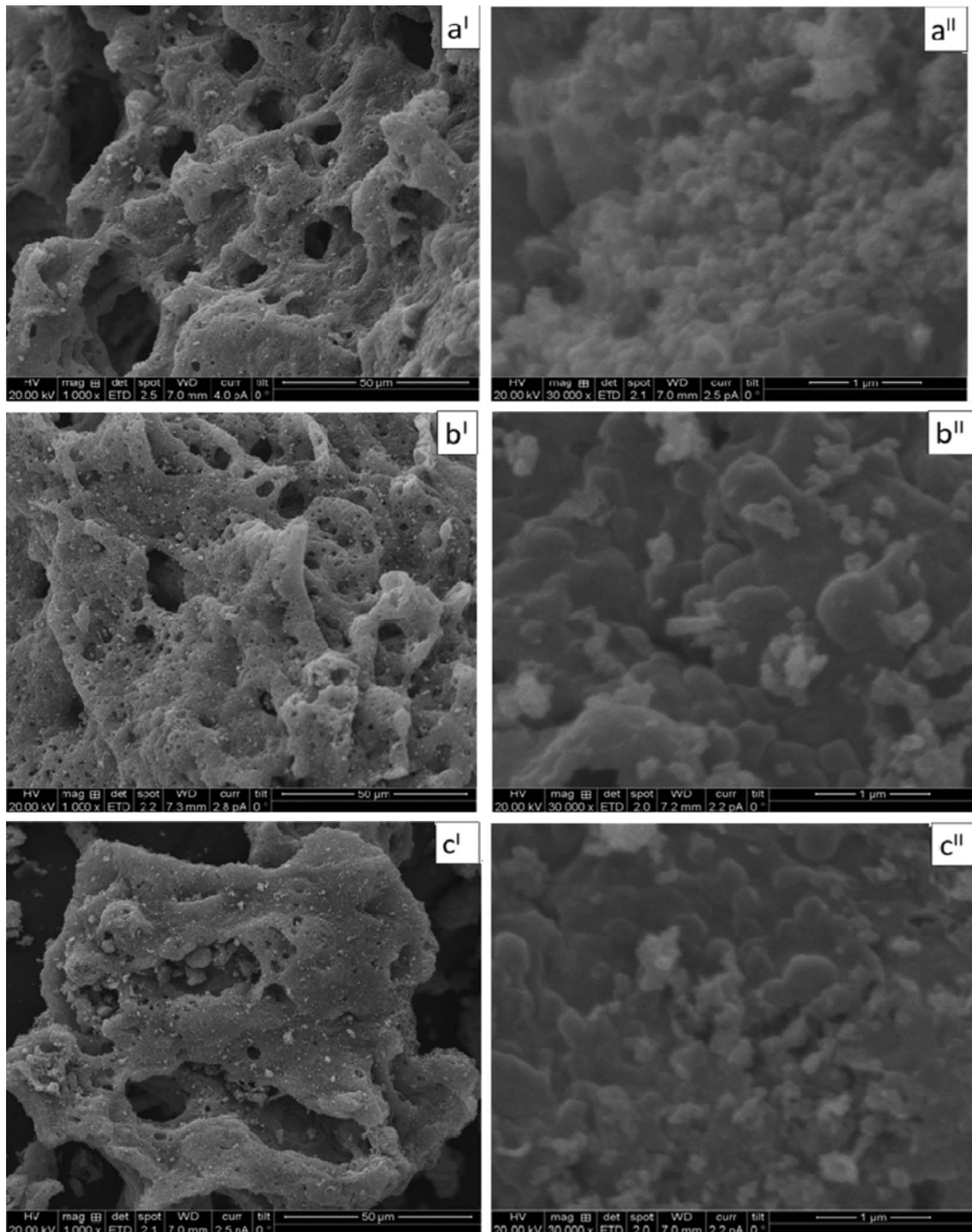


Figure 3. SEM micrographs of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ samples: (**a**^{I&II}) $x = 0.65$, (**b**^{I&II}) $x = 0.7$ and (**c**^{I&II}) $x = 0.75$.

Table 2. Compositional concentration (%) of the constituent elements of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ ferrite system by EDAX.

x	Ni	Zn	Fe	O
0.65	4.29	10.20	30.38	55.13
0.70	2.70	11.70	30.61	54.99
0.75	2.66	12.33	30.41	54.60

The results show that the M_s and M_r decreases with the increase in the zinc content. A marked reduction in M_s has been reported for $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$, as the concentration of Zn^{2+} was increased from $x = 0.5$ to 0.7 .²¹ This behaviour was attributed to the spin canting effect that occurs when BB interactions are comparable to AB interactions.³¹ It is important to state that due to antiparallel

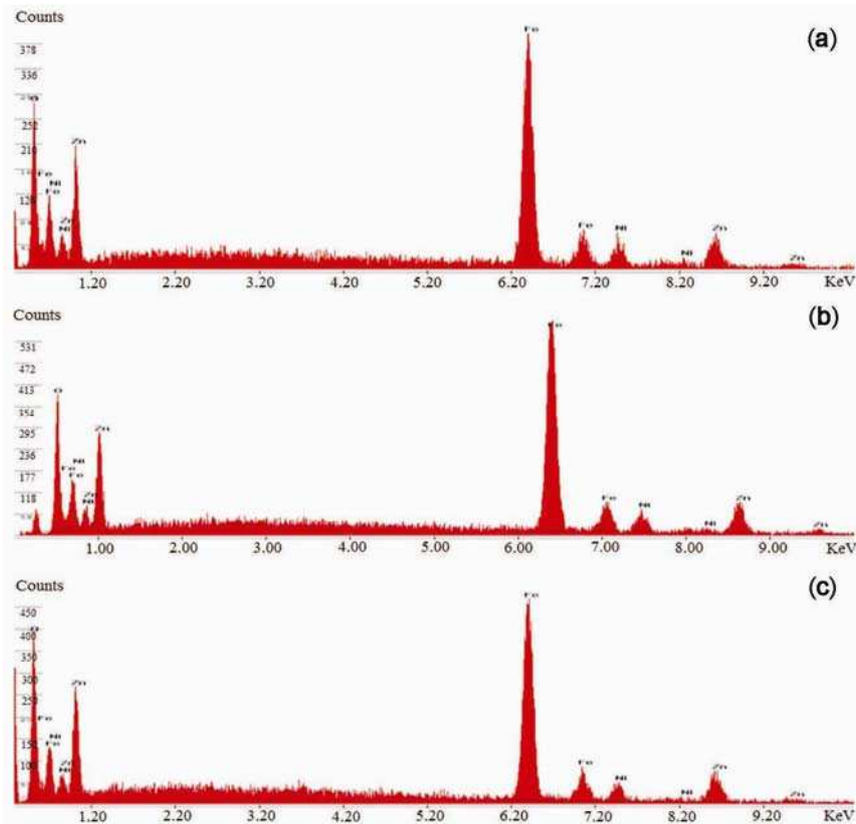


Figure 4. EDAX spectra of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ samples: (a) $x = 0.65$, (b) $x = 0.7$ and (c) $x = 0.75$.

Table 3. Effects of Zn doping on the magnetic properties of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ MNPs: (a) $x = 0.65$, (b) $x = 0.70$ and (c) $x = 0.75$.

Zn conc. ' x '	Saturation magnetization, M_s (emu g^{-1})	Remanence magnetization, M_r (emu g^{-1})	Coercivity, H_c (Gauss)	M_r/M_s
0.65	36	22.5	239	0.63
0.70	32	19.0	224	0.59
0.75	25	15.0	234	0.60

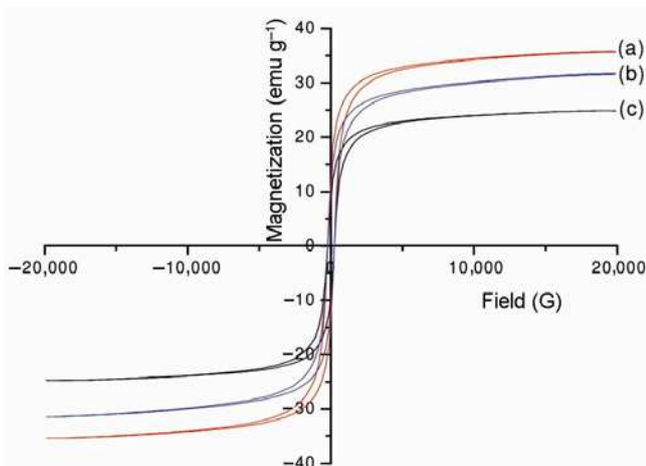


Figure 5. Magnetic hysteresis curves of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ MNPs measured at room temperature for (a) $x = 0.65$, (b) $x = 0.70$ and (c) $x = 0.75$.

coupling between the A-sites and B-sites in spinel structured materials, the net magnetization (M) is given by $M = \sum M_{\text{Bsites}} - \sum M_{\text{Asites}}$. Zn^{2+} ions are known to occupy the tetrahedral (A) site; Ni^{2+} ions have a preference for the octahedral (B) site while the Fe^{3+} ions are distributed over both sites in mixed Ni–Zn ferrites. When the concentration of Fe^{3+} ions in the A sublattice is diluted by low concentrations of diamagnetic substitutions (such as Zn^{2+} ions), the net magnetization increases.³² However, magnetization decreases at higher levels of doping as seen in this study with $x = 0.65$ – 0.75 . The reason for this is that low Zn concentrations reduce the number of spins occupying the A sublattices, causing the net magnetization to increase. As the Zn content increases, the exchange interactions are weakened and the B spins are no longer held rigidly parallel to the few remaining A spins. The decrease in the B-sublattice moment, interpreted as a spin departure from colinearity, causes the

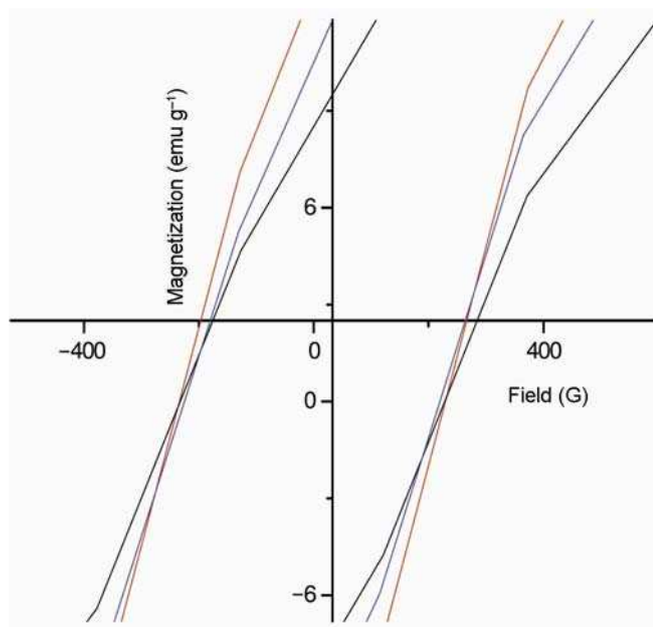


Figure 6. Magnified view of hysteresis loops of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ samples to show their ferrimagnetism.

effect known as canting.²¹ Also, the decrease in M_s with the increase in the zinc content can be explained from the crystallite sizes of the samples shown in table 1. The crystallite sizes (and by extension, the M_s) of the samples clearly show a slight decrease with the increase in the zinc content. It is known that M_s gradually increases with crystallite size,^{33,34} which might be due to decreased domain walls displacement as the crystallite sizes increases in the multi-domain range.³⁵ The M_r is known to vary linearly with M_s in ferromagnetic (or ferrimagnetic) materials as observed in this study.

The variation in coercivity (H_c) with Zn doping is clearly observed in the magnified view of the hysteresis curves in figure 6. It shows hysteresis loops that are typical of soft ferrimagnetic materials. The H_c of $\text{Ni}_{0.35}\text{Zn}_{0.65}\text{Fe}_2\text{O}_4$ sample recorded the highest values, while $\text{Ni}_{0.3}\text{Zn}_{0.7}\text{Fe}_2\text{O}_4$ sample recorded the lowest. It can be seen that H_c decreased in $x = 0.7$ and increased in $x = 0.75$. The H_c of the samples shows a nonlinear relationship with zinc doping and the crystallite size. According to literature,^{7,36–38} there is no firm relationship between H_c and grain size thereby establishing that the influence of the crystallite size of different alloy systems on H_c is undetectable. The squareness ratio (M_r/M_s) values of all samples were higher than 0.5, highlighting the potential applications of these materials for recording medium. These results show that the magnetic properties of the material are dependent on the amount of Zn present. All samples recorded a fairly high saturation magnetization (25–36 emu g^{-1}) with the increase in M_s and M_r with the decrease in x (Zn content).

4. Conclusion

A fuel-rich glycine–nitrate mixture in the combustion synthesis of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ was sufficient for the formation of pure nanocrystalline spinel ferrite phase without any further heat treatment at high temperatures. Raman spectra show tetrahedral and octahedral sites in the structure of $\text{Ni}_{1-x}\text{Zn}_x\text{Fe}_2\text{O}_4$ and also imply the doping of Zn^{2+} and displacement of Fe^{3+} ions from the tetrahedral site. EDAX showed that the samples were close to the nominal compositions. The magnetic measurement shows that the saturation magnetization and remanence magnetization decreases with the increase in the zinc content, while coercivity showed a nonlinear relationship. The squareness ratio (M_r/M_s) values of all samples were higher than 0.5, highlighting the potential applications of these materials for recording medium.

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References

1. Sugimoto M 1999 *J. Am. Ceram. Soc.* **82** 269
2. Lazarević Z Ž, Jovalekić Č, Sekulić D, Slankamenac M, Romčević M, Milutinović A and Romčević N Ž 2012 *Sci. Sinter.* **44** 331
3. Sivakumar P, Ramesh R, Ramanand A, Ponnusamy S and Muthamizhchelvan C 2011 *Mater. Res. Bull.* **46** 2208
4. Sivakumar P, Ramesh R, Ramanand A, Ponnusamy S and Muthamizhchelvan C 2011 *Mater. Res. Bull.* **46** 2204
5. Kasapoglu N, Baykal A, Toprak M S, Koseoglu Y and Bayrakdar H 2007 *Turk. J. Chem.* **31** 659
6. Sepelak V, Baabe K, Mienert K, Schultze K, Krumeich F, Litterst F J and Becker K D 2003 *J. Magn. Magn. Mater.* **257** 377
7. Salunkhe A B, Khot V M, Phadatar M R, Thorat N D, Joshi R S, Yadav H M and Pawar S H 2014 *J. Magn. Magn. Mater.* **352** 91
8. Morrison S A, Cahill C L, Carpenter E E, Calvin S, Swaminathan R, McHenry M E and Harris V G 2004 *J. Appl. Phys.* **95** 6392
9. Krishna K R, Kumar K V and Ravinder D 2012 *Adv. Mater. Phys. Chem.* **2** 185
10. Vermaa A, Goela T C, Mendirattaa R G and Kishan P 2000 *J. Magn. Magn. Mater.* **208** 13

11. Vermaa A, Goela T C, Mendirattaa R G and Gupta R G 1999 *J. Magn. Magn. Mater.* **192** 271
12. Mohan G R, Ravinder D, Reddy A V R and Boyanov B S 1999 *Mater. Lett.* **40** 39
13. Ajmal M and Maqsood A 2007 *Mater. Sci. Eng. B* **139** 164
14. Sileo E E, Rotelo R and Jacobo S E 2002 *Physica B: Condens. Matter* **320** 257
15. Costa A C F M, Tortella E, Morelli M R and Kiminami R H G A 2003 *J. Magn. Magn. Mater.* **256** 174
16. Cabanas A and Poliakoff M 2001 *J. Mater. Chem.* **11** 1408
17. Song H J, Oh J H, Choi S C and Lee J C 2002 *Phys. Status Solidi A* **189** 849
18. Akther Hossain A K M, Mahmud S T, Seki M, Kawai T and Tabata H 2007 *J. Magn. Magn. Mater.* **312** 210
19. Deng H, Chen H and Li H 2007 *Mater. Chem. Phys.* **101** 509
20. Costa A C M, Tortella E, Morelli M R, Kaufman M and Kiminami R H G A 2002 *J. Mater. Sci.* **37** 3569
21. Costa A C M, Silva V J, Cornejo D R, Morelli M R, Kiminami R H and Gama L 2008 *J. Magn. Magn. Mater.* **320** e370
22. Costa A C M, Diniz A P, Silva V J, Kiminami R H, Cornejo D R, Gama A M, Rezende M C and Gama L 2009 *J. Alloys Compd.* **483** 563
23. Shannon R D 1976 *Acta Crystallogr.* **32** 751
24. Ahlawat A and Sathe V G 2011 *J. Raman Spectrosc.* **42** 1087
25. Dubey A, Sathe V G and Rawat R 2008 *J. Appl. Phys.* **104** 113530
26. Carta D, Casula M F, Falqui A, Loche D, Mountjoy G, Sangregorio C and Corrias A 2009 *J. Phys. Chem.* **113** 8606
27. Ivanov V G, Abrashev M V, Iliev M N, Gospodinov M M J and Aroyo M M I 2010 *Phys. Rev. B: Condens. Matter* **82** 024104
28. Wang Z, Lazor P, Saxena S K and O'Neil H S C 2002 *Mater. Res. Bull.* **37** 1589
29. Kishan P, Sagar D R and Swarup P 1985 *J. Less Common Met.* **108** 345
30. Kambale R C, Shaikh P A, Kamble S S and Kolekar Y D 2009 *J. Alloys Compd.* **478** 599
31. Costa A C F M 2002 PhD Thesis (Sao Paulos, Brazil: Department of Engineering of Materials, Federal University of Sao Carlos)
32. Bercoff P G and Bertorello H R 2000 *J. Magn. Magn. Mater.* **213** 56
33. Hu P, Yang H, Pan D, Wang H, Tian J, Zhang S, Wang X and Volinsky A A 2010 *J. Magn. Magn. Mater.* **322** 173
34. Kumar E R, Jayaprakash R and Kumar S 2014 *Mater. Sci. Semiconduct. Process.* **17** 173
35. Zhao L, Yang H, Yu L, Cui Y, Zhao X, Zou B and Feng S 2006 *J. Magn. Magn. Mater.* **301** 445
36. Shirsath S E, Jadhav S S, Toksha B G, Patange S M and Jadhav K M 2011 *J. Appl. Phys.* **110** 013914
37. Herzer G 1990 *IEEE Trans. Magn.* **26** 1397
38. Zeng Q, Baker I, McCreary V and Yan Z 2007 *Mater. Res. Soc. Symp.* **980** 0980-II06-09