VARIATION OF STRUCTURAL PROPERTIES OF Al DOPED Ni-Cd FERRITES WITH SINTERING TIME


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The aim of present work is to synthesis and study the structural properties of Al doped Ni-Cd ferrites, series by ceramic method. The samples are sintered at a temperature of 1100 C with sintering time of t=9, 11, 13, 15 hrs. The samples were characterized by X-ray diffraction, scanning electron microscope, energy dispersive X-ray analysis and Fourier transform-Infrared. The single phase simple cubic spinel structure is confirmed by X-ray diffraction patterns. The lattice parameter is found to be in the range of 8.478 Å-8.481 Å. The uniform size distribution is observed in the scanning electron microscope micrographs. The elemental analysis is done by energy dispersive X-ray analysis confirming the presence of all the metal ions present in the ferrite formula. Fourier transform-Infrared spectra showed two bands υ₁(581-582 cm⁻¹) and υ₂(less than 400 cm⁻¹) which are attributed to metal oxygen bond vibrations at tetrahedral and octahedral sites respectively.

Keywords: X-ray diffraction, scanning electron microscope, energy dispersive X-ray analysis, Fourier transform-Infrared

Introduction

Ferrites are an attractive class of research materials because of their comprehensive physical properties and application in advanced technology. Modification of their structure has been shown to play an important role in the optimization of their structural, optical and electrical properties [1-2]. Cubic spinel ferrites are chemical compounds with the formula of AB₂O₄, where A and B represent various metal cat-ions, usually including iron. In the normal spinel structure, the 8 divalent metal ions go into the A sites, and the 16 trivalent iron ions have preference for B sites [1]. Among the magnetic materials that have found broad practical application in technology, ferrites deserve attention [2].

Cobalt ferrite (CoFe₂O₄) has attracted much attention owing to its excellent properties, such as its relatively high saturation magnetization (Mₛ), high coercivity (H_c), magnetic anisotropy, good chemical stability and good catalytic activity. Its physical properties are strongly dependent on the cation distribution among the tetrahedral (A) and the octahedral (B) sites in the crystal lattice. Depending upon the distribution of the various cat-ion substituents over tetrahedral and octahedral sites, its magnetic and electrical properties can be modified according to the research interest [3].

Spinel ferrite is known to be good candidate for magnetic recording and microwave applications [4]. Nano-sized grains of ferrite are being applied in permanent magnets [5], magnetic drug delivery [6], and high-density information storage [7]. Soft magnetic material have a widespread applications in recording heads, antenna rods, loading coils, microwave devices, core material for power transformers due to their high resistivity and low eddy current losses [8]. Ferrite materials can be prepared by numbers of methods such different routes like co-precipitation [9], solid state reaction [10-11], citrate precursor [12], sol–gel and auto-combustion [13], ball milling [14] hydrothermal [15].

This paper presents the influence of sintering time with the general chemical formula Ni₀.₅Cd₀.₅Al₀₁Fe₁.₉O₄ series which are prepared by ceramic method. Using X-ray diffraction (XRD), scanning
electron microscope (SEM), energy dispersive X-ray analysis (EDAX) and Fourier transform-Infrared (FT-IR) analysis, detailed structural and mechanical properties of crystalline Ni-Cd-Al spinel ferrites reported in this paper.

1. Experiments

In photo acoustic effect, interaction of electromagnetic radiation with matter generates sound. This effect consists of the absorption of incident radiation by target molecule. Photo acoustic effect is popular due to a minimal sample preparation during execution, the ability to examine scattering and opaque sample [12] along with the capability to access depth profile [13]. The ferrites with general chemical formula Ni$_{0.5}$Cd$_{0.5}$Al$_{0.1}$Fe$_{1.9}$O$_4$ were synthesized by ceramic method using AR grade Nickel oxide, Cadmium oxide, and Ferric oxide in molar preparation and mixed mechanically in agate motor in acetone medium as shown in Fig.1. Then the ferrites were presintered with temperature constant 1100ºc for different time 9 hours, 11 hours, 13 hours, and 15 hours.

![Flow chart of synthesis of ferrite samples using ceramic method.](image)

The pre sintered powders were subjected to hard milling process in acetone medium. One or two drops of PVA solution of 20% concentration was added as binder. All the samples were pressed into two pellets of 1cm in diameter and 0.2 cm in height by applying a pressure of about 5tons/square inch for 5 minutes. The pellets were subjected to final sintering at 1100ºc for 15 hours.

The Structural characterization of the ferrite pellets is carried out the X-ray diffraction patterns of the samples have been obtained on Bruker AXS D8 Advance Diffractometer (XRD), (with Cu-K$_\alpha$ radiation, wavelength, $\lambda$= 1.5406 Å), the SEM-EDAX spectrographs of the samples have been obtained on JEOL Model JSM - 6390LV and the FTIR of the samples have been obtained on Thermo Nicolet, Avatar 370.

2. Results and Discussion

2.1. X-ray diffraction (XRD) analysis

XRD patterns of the Ni$_{0.5}$Cd$_{0.5}$Al$_{0.1}$Fe$_{1.9}$O$_4$ (t=9, 11, 13, 15 hrs) ferrite series is shown in Fig.2. The peaks matched well with those from the Joint Committee on Powder Diffraction Standards (JCPDS) card no.00-022-1086. The reflections observed from the planes (220), (311), (400), (422), (511), (440) and (533) for the samples confirm the formation of a cubic spinel structure.

The lattice parameter of all the samples was calculated and the values are given in Table 1. The lattice parameter was found to remain constant. The hopping length; the bond length, ionic radii and the value of X-ray density ($\Delta x$) and stacking fault coefficient were calculated elsewhere [10, 11] and the values are tabulated in Table 1. The actual density is calculated using standard method. The values are tabulated in Table 1.
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Fig. 2. XRD graph of Ni_{0.5}Cd_{0.5}Al_{0.1}Fe_{1.9}O_{4} ferrite synthesized by ceramic method.

Table 1 - Calculated values of lattice parameter, volume of unit cell, hopping length, bond length, ionic radii, stacking fault coefficient, x-ray density, actual density and porosity.

<table>
<thead>
<tr>
<th>Sintering time (t)</th>
<th>9 hr</th>
<th>11 hr</th>
<th>13 hr</th>
<th>15 hr</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lattice parameter a (Å)</td>
<td>8.48</td>
<td>8.47</td>
<td>8.48</td>
<td>8.48</td>
</tr>
<tr>
<td>Volume of unit cell V (10^{-30} m^3)</td>
<td>609</td>
<td>608</td>
<td>609</td>
<td>610</td>
</tr>
<tr>
<td>Hopping length L_A (Å)</td>
<td>3.67</td>
<td>3.67</td>
<td>3.67</td>
<td>3.67</td>
</tr>
<tr>
<td>Hopping length L_B (Å)</td>
<td>3.00</td>
<td>3.00</td>
<td>3.00</td>
<td>3.00</td>
</tr>
<tr>
<td>Bond Length A-O (Å)</td>
<td>1.94</td>
<td>1.94</td>
<td>1.94</td>
<td>1.94</td>
</tr>
<tr>
<td>Bond length B-O (Å)</td>
<td>2.06</td>
<td>2.06</td>
<td>2.06</td>
<td>2.06</td>
</tr>
<tr>
<td>Ionic Radii r_A (Å)</td>
<td>0.59</td>
<td>0.59</td>
<td>0.59</td>
<td>0.59</td>
</tr>
<tr>
<td>Ionic Radii r_B (Å)</td>
<td>0.71</td>
<td>0.71</td>
<td>0.71</td>
<td>0.71</td>
</tr>
<tr>
<td>Stacking fault coefficient α (10^{-3})</td>
<td>0.43</td>
<td>0.45</td>
<td>0.36</td>
<td>0.53</td>
</tr>
<tr>
<td>x-ray density Δx (gr/cm³)</td>
<td>5.63</td>
<td>5.64</td>
<td>5.63</td>
<td>5.63</td>
</tr>
<tr>
<td>Actual density d_a (gr/cm³)</td>
<td>4.38</td>
<td>3.51</td>
<td>3.82</td>
<td>3.76</td>
</tr>
<tr>
<td>Porosity (%)</td>
<td>22.24</td>
<td>37.67</td>
<td>32.17</td>
<td>33.25</td>
</tr>
</tbody>
</table>

The crystallite size of ferrite samples was calculated by using Debye-Scherrer [10, 11] equation

\[ D = \frac{0.9 \cdot \lambda}{\beta \cos \theta} \]

where \( D \) is the crystallite size, \( \beta \) is pure diffraction broadening given by the full width half maximum (FWHM) of the peaks, \( \lambda \) is the wavelength of the radiation used, \( \theta \) is the angle of diffraction. The Dislocation density calculated using two different formulas \( \rho_D = 1/D^2 \) and \( \rho_D = 15\epsilon/aD \) and micro-strain calculated using \( \epsilon = \beta \cos \theta / 4 \) are also tabulated in Table 2.

Table 2 - Calculated values of crystallite size, micro strain and dislocation density using WH plots, SSP and standard formula.

<table>
<thead>
<tr>
<th>Sintering time t in hr</th>
<th>Crystallite size (Å)</th>
<th>Micro strain (10^{-3})</th>
<th>Dislocation density (\rho) (10^{14})</th>
</tr>
</thead>
<tbody>
<tr>
<td>From WH graph</td>
<td>From SSP</td>
<td>From formula</td>
<td>Micro strain = slope/4 (WH)</td>
</tr>
<tr>
<td>9</td>
<td>835</td>
<td>651</td>
<td>522</td>
</tr>
<tr>
<td>11</td>
<td>766</td>
<td>700</td>
<td>439</td>
</tr>
<tr>
<td>13</td>
<td>462</td>
<td>415</td>
<td>457</td>
</tr>
<tr>
<td>15</td>
<td>510</td>
<td>501</td>
<td>503</td>
</tr>
</tbody>
</table>
2.2. Williamson Hall (WH) plot and Size-Strain (SS) Plot Method

The distinct \( \theta \) dependencies of size and strain broadening of the diffraction lines laid the basis for Williamson and Hall method (WH). An additional component \( \beta_s = 4\varepsilon \tan \theta \) to the Scherrer equation results in WH equation \( \beta_{hkl} \cos \theta = \frac{K \cdot \lambda}{D} + 4\varepsilon \sin \theta \), [11, 16].

The WH plots for the \( \text{Ni}_{0.5}\text{Cd}_{0.5}\text{Al}_{0.1}\text{Fe}_{1.9}\text{O}_4 \) (t= 9, 11, 13, 15 hrs) are shown Fig. 3 (a-d).

![WH plots](image)

**Fig.3.** WH plots of \( \text{Ni}_{0.5}\text{Cd}_{0.5}\text{Al}_{0.1}\text{Fe}_{1.9}\text{O}_4 \) ferrite series synthesized by ceramic method.

Using the slope and intercept values of the graph of \( \beta \cos \theta \) with respect to \( \sin \theta \); the value of crystallite size and microstrain are calculated and are tabulated in the Table 2.

The SS plot is a tool to understand the isotropic nature and micro-strain contribution. The SS plots of \( \text{Ni}_{0.5}\text{Cd}_{0.5}\text{Al}_{0.1}\text{Fe}_{1.9}\text{O}_4 \) (t= 9, 11, 13, 15 hrs) ferrite series is shown in Fig. 4 (a-d). In this approximation, crystallite size and strain are having a relation given by the equation \((d_{hkl} \beta_{hkl} \cos \theta)^2 = \frac{K \cdot \lambda}{D} (d_{hkl} \beta_{hkl} \cos \theta) + \left(\frac{\varepsilon}{2}\right)^2\), where \( K \) is a constant [11, 17-18]. The calculated value of crystallite size and microstrain are tabulated in the Table 2. The microstrain is observed to be decreasing with increase in sintering time. The crystallite size is observed to be steady. All the samples showed the dislocation density in the range of \( 10^{14} \).

2.3. Scanning electron microscope studies

The SEM micrographs of \( \text{Ni}_{0.5}\text{Cd}_{0.5}\text{Al}_{0.1}\text{Fe}_{1.9}\text{O}_4 \) (t= 9, 11, 13, 15 hrs) series are shown in the Fig. 5(a-d). The images show uniform distribution of the grains with uniform grain size of the order of 1 \( \mu \)m. The grains are having octahedral and tetrahedral structures showing very sharp boundary growth. The grain size increased with increase in the sintering time.
Fig. 4. SS plots of Ni$_{0.5}$Cd$_{0.5}$Al$_{0.1}$Fe$_{1.9}$O$_4$ ferrite series synthesized by ceramic method.

The elemental analysis is done by EDAX analysis and the diffraction spectra of Ni$_{0.5}$Cd$_{0.5}$Al$_{0.1}$Fe$_{1.9}$O$_4$ (t= 9, 11, 13, 15 hrs) series are shown in Fig. 6, a-d.

Fig. 5. SEM micrographs of Ni$_{0.5}$Cd$_{0.5}$Al$_{0.1}$Fe$_{1.9}$O$_4$ ferrite series synthesized by ceramic method.
2.4. Energy dispersive X-ray analysis studies

The spectra confirming the presence of all the metal ions present in the ferrite formula and the atomic proportion tabulated in Table 3.

Table 3 - Initial values of atomic proportions of metal ions taken by stoichiometric proportion and observed values using EDAX analysis of in the samples of Ni_{0.5}Cd_{0.5}Al_{0.1}Fe_{1.9}O_4 ferrite series with sintering time t=9, 11, 13, 15 hrs.

<table>
<thead>
<tr>
<th></th>
<th>Weight (%)</th>
<th>Observed atomic proportion using EDAX</th>
<th>Stoichiometric composition taken for synthesis</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>t=9 hr</td>
<td>t=11 hr</td>
<td>t=13 hr</td>
</tr>
<tr>
<td>Al</td>
<td>0.52</td>
<td>1.04</td>
<td>1.27</td>
</tr>
<tr>
<td>Fe</td>
<td>45.44</td>
<td>52.54</td>
<td>53.53</td>
</tr>
<tr>
<td>Ni</td>
<td>13.52</td>
<td>17.17</td>
<td>16.74</td>
</tr>
<tr>
<td>Cd</td>
<td>9.07</td>
<td>2.64</td>
<td>2.93</td>
</tr>
<tr>
<td>Tot</td>
<td>93.76</td>
<td>98.72</td>
<td>100</td>
</tr>
</tbody>
</table>

2.5. Fourier transform infrared (FT-IR) studies

The FT-IR spectra of Ni_{0.5}Cd_{0.5}Al_{0.1}Fe_{1.9}O_4 ferrite series with sintering time t=9, 11, 13, 15 hrs Fig. 7. The transmittance spectra showed two peaks for all the samples of the series. The first peak $\nu_1$ (568-580 cm$^{-1}$) corresponds to the tetrahedral metal oxygen bond and second peak $\nu_2$ (near 400 cm$^{-1}$) corresponds to octahedral metal oxygen bond in the ferrite structure [19].
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Conclusion

The $\text{Ni}_{0.5}\text{Cd}_{0.5}\text{Al}_{0.1}\text{Fe}_{1.9}\text{O}_4$ ferrite series with sintering time $t=9$, 11, 13, 15 hrs are successfully synthesized by ceramic method. The XRD studies confirm the single phase cubic spinel structure of the samples. The lattice parameter ($8.47$ Å - $8.48$ Å) remains almost constant with increase in sintering temperature. Stacking fault coefficient observed to be very low, affirming the XRD peak position are at the expected value. The crystallite size is calculated using three different methods showed almost constancy (~50 nm) with increase in the sintering time. The SEM images show uniform size of the grains of size of the range of 1 μm. The octahedral and tetrahedral structures are observed in morphology study. The grain size increased with increase in the sintering time. The EDAX studies confirm the presence of all the metal ions used in synthesis. The FTIR studies show two peaks $\nu_1$ (581-582 cm$^{-1}$) and $\nu_2$ (less than 400 cm$^{-1}$) confirming the presence of octahedral and tetrahedral positions inside the structure.

REFERENCES


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