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# **Electronics Characterization of Al and Co doped Lithium Cermagnet**

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#### ABSTRACT

A new compound of Lithium hexaferrites with general chemical formula  $Li_{0.5}Fe_{0.5+x}Al_{12-2x}Co_xO_{19}$  where x = 2, 3, 4, 5 and 6, were synthesized by standard ceramic technique using stoichiometric proportion. The lattice parameters of the multicrystalline materials have been determined and characterized by X-ray diffraction technique (XRD). From XRD data shows the formation of samples of a single phase hexagonal M-type ferrites with a space group P6<sub>3</sub>/mmc (194). The electrical conduction was carried out over the temperature range from 27<sup>o</sup>C- 800<sup>o</sup> C. The electrical conduction in the ferrites was explained on the basis of Vervway models. A Seebeck study reveals that all samples are N-type semiconductors.

Keywords: Magnetoplumbite, Hexaferrites, electrical conductivity, Seebeck coefficient etc.

## INTRODUCTION

Among the family of hexagonal ferrites, the Mangnetoplumbite hexaferrites (M-Type) created much attention due to wide range of application in industries and created potential to interest in technological and scientific research due to their application importance such as high density magnetic recording, microwave device materials, in computer system etc. The application need particularly magnetic and electrical specification with the view, many attempt have been improved the properties of hexagonal ferrites using different tract of additives. The Calcium hexaferrites [1-9] have magnetic properties comparable to BaM and SrM. In calcium ferrites many attempt has been made to replace Fe<sup>+3</sup> ions with Al<sup>+3</sup>, Cr<sup>+3</sup> and Co<sup>+3</sup> etc. A compound with the combination of bivalent-tetravalent cation was also used to replace Fe<sup>+3</sup> ions such as Cu-Ti, Co-Ti, Co-Sn, Zn-Sn etc [10-12] without any appreciable change in BaM structure. When Fe<sup>+3</sup> ions are replace by non magnetic ions like Ti<sup>+4</sup> and Sn<sup>+4</sup> etc. In the same way Lithium ferrites substituted with Al and Co has been studied structurally, electrically and magnetically [13] In the present study a series of five sample with chemical formula  $Li_{0.5}Fe_{0.5+x}Al_{12-2X}Co_XO_{19}$  where x=2, 3, 4, 5 and 6 were prepared. Lithium ferrites have attracted considerable attention because of the squareness of hysterisis loop coupled with superior temperature performance the crystal structure of M-type like compound with a space group P6<sub>3</sub>/ mmc (194) can be described as superposition of two structure block namely R-block with composition BaFe<sub>6</sub>O<sub>11</sub> and S-block with composition Fe<sub>6</sub>O<sub>8</sub> [14] in the stoichiomatric ratio.

# MATERIALS AND METHODS

All the multicrystalline powder sample were synthesized by high temperature solid state reaction using A.R grade oxides of stoichiomatric ratio  $Li_2O_3$ ,  $Fe_2O_3$ ,  $Al_2O_3$  and  $Co_2O_3$  mixture.  $Li_2O_3$  oxide was carefully dehydrated before the mixing procedure. After grinding the mixture under acetone acid for six hours, the preparation of pellets of thoroughly grounded mixture in the proper molar ratio with 5% of PVA as a binder are prepared by applying 10 tone

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pressure per square inch. These pallets slowly heated in the furnace at 600  $^{0}$ C for 5-6 hours to remove binder [15].Then it was fired at 1200 0C for 120 hours continuously, after that the furnace were cooled at the rate of 20 $^{0}$ C per hours up to 1000  $^{0}$ C and then 50 $^{0}$ C per hours and then cooled in natural way the phase of final sample were verified by Philips X-ray diffrectometory using N-fillter copper radiation. The mean grain size of multicrystalline samples was in the range 150-200 Å. All the samples show single phase formation with a space group P63/mmc (194).

The d.c.electrical conductivity was measured by the methods using LCR meter. The end faces of pellets were coated with thin layer of conducting silver paste and measurement were made from room temperature to 800 K Thermoelectric power measurement were carried out after sandwiching the thick pellet between two copper rod from room temperature to  $550^{\circ}$ C

#### **RESULT AND DISCUSSION**

In the present work cobalt and aluminium substituted lithium hexaferrites were introduced with general formula  $Li_{0.5}$   $Fe_{0.5+x}$   $Al_{12-2X}$   $Co_X$   $O_{19}$  where x =2, 3, 4,5 and 6. The ions in Ba-M compounds can be replaced partly by  $Co^{+3}$  or completely  $Li^{+1}$  and combination of  $Fe^{+3}$  and  $Al^{+3}$  ions without changing the crystal lattice symmetry [16]. In all the specimens substituted ions would be chosen to keep electrical neutrality and to have similar ionic radii in these ferrites. The cobalt and aluminium play an important role in the property variation. The XRD technique is used to confirms the formation of hexagonal magnetoplumbite structure of compounds belonging to a space group  $P6_3 / mmc$  (194) A X-ray diffraction pattern of the compound Li0.5  $Fe_{6.5}$   $Co_6O_{19}$  is shown in Fig.1.



Fig. - 1: X-ray Diffraction pattern of Li 0.5 Fe 6.5 Co<sub>6</sub>O<sub>19</sub>

Due to the resemblance of ionic radii of  $Fe^{+3}$  with  $Co^{+3}$  and  $Al^{+3}$  ions, the ferrites ions will replace by cobalt and aluminium. It is seen that former ions are very easily replaced at any substituted variation in all specimens .The hexagonal lattice parameters 'a' and 'c' decreases linearly with the substitution variation  $Co^{+3}$  and  $Fe^{+3}$  concentration in all specimens. The decreases in lattice parameters due to close packing of lattices in the materials [17]. The decrease in lattice parameter and cell volume agree with result for Ba and Sr ferrite [18], the numerical values of compositional data such as lattice constant, cell volume and X- ray density are tabulated in table -1. The observed value of electrical conductivity, and activation energy for specimens are also tabulated in table -2.

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Compounds	Lattice parameters		CallWoluma $(\hat{A})^3$	Mol.Wt	X-Ray Density
	a (Å)	c (Å	Cenvolume (A)	gm	gm/cc
Li <sub>0.5</sub> Fe <sub>2.5</sub> Al <sub>8</sub> Co <sub>2</sub> O <sub>19</sub>	5.859	22.323	663.81	780.89	3.9065
Li <sub>0.5</sub> Fe <sub>3.5</sub> Al <sub>6</sub> Co <sub>3</sub> O <sub>19</sub>	5.818	21.879	641.48	841.79	4.3578
Li <sub>0.5</sub> Fe <sub>4.5</sub> Al <sub>4</sub> Co <sub>4</sub> O <sub>19</sub>	5.807	21841	637.90	902.66	4.6992
Li <sub>0.5</sub> Fe <sub>5.5</sub> Al <sub>2</sub> Co <sub>5</sub> O <sub>19</sub>	5.794	21.807	633.96	93.54	5.0473
$Li_{0.5}Fe_{6.5}Co_{6}O_{19}$	5.695	21.262	597.22	1024.42	5.6963

 Table – 1 Consolidated structural data of lithium ferrites

Table –	2 Electric	al conductivity	of lithium	ferrites
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Compounds	Electrical Resistivity at room temperature. ( $\Omega$ cm)	Activation energy E in (ev)	Electrical Conductivity at room temperature. $(\Omega \text{ cm})^{-1}$
Li <sub>0.5</sub> Fe <sub>2.5</sub> Al <sub>8</sub> Co <sub>2</sub> O <sub>19</sub>	4.559x10 <sup>11</sup>	0.77	2.193 x10 <sup>-12</sup>
Li <sub>0.5</sub> Fe <sub>3.5</sub> Al <sub>6</sub> Co <sub>3</sub> O <sub>19</sub>	3.21x 10 <sup>9</sup>	0.63	3.11 x10 <sup>-10</sup>
Li <sub>0.5</sub> Fe <sub>4.5</sub> Al <sub>4</sub> Co <sub>4</sub> O <sub>19</sub>	4.65x10 <sup>7</sup>	0.52	2.148 x10 <sup>-8</sup>
Li <sub>0.5</sub> Fe <sub>5.5</sub> Al <sub>2</sub> Co <sub>5</sub> O <sub>19</sub>	6.84x10 <sup>6</sup>	0.48	1.46 x10 <sup>-7</sup>
Li <sub>0.5</sub> Fe <sub>6.5</sub> Co <sub>6</sub> O <sub>19</sub>	1.72x10 <sup>5</sup>	0.39	5.78 x10 <sup>-6</sup>

The plot of  $\ln \sigma vs (1/T) x 10^{-3} \text{ K}$  for the entire sample was almost linear. The electrical conductivity of these ferrites increases with increasing ferrite ion concentration. The electrical conductivity of sintered specimens varies from 2.193 x10<sup>-12</sup> ( $\Omega$  cm)<sup>-1</sup> to 5.78 x10<sup>-6</sup> ( $\Omega$  cm)<sup>-1</sup> of these ferrites [20], the electrical conductivity values obtained by Reddy et al 1981 is 2.3 x 10<sup>-6</sup> ( $\Omega$ cm)<sup>-1</sup> cm<sup>-1</sup> and 2x10<sup>-2</sup>( $\Omega$ cm)<sup>-1</sup> Rezlesen et al 1974 for lithium ferrite.

In the present work the electrical conductivity value obtained for the compounds are  $2.193 \times 10^{-12} (\Omega \text{cm})^{-1}$  to  $5.78 \times 10^{-6} (\Omega \text{cm})^{-1}$ . The value of the conductivity may be partly attributed to the low evaporation of lithium from the sample prepared different from these of Rozlescu etal 1974 and Venugopal Reddy 1981. The variation of activation energy with the substitutional variable parameters x -may be explain on the basis of Verway model [21] a small number of ferrous ions (Fe<sup>+2</sup>) are generally developed during sintering process which lead the conductivity in ferrites suggesting the hopping mechanism according (Fe<sup>+2</sup>-Fe<sup>+3</sup>+e<sup>-</sup>) [23-24], However these transition take place for a very small interval of time and are not detectable by the ordinary method. This valence exchange mechanism of Verway may be considered for these ferrites as general applicable to M-type ferrites.

#### CONCLUSION

In the present work of lithium ferrites, is to check the formation of ferrite containing  $AI^{+3}$ , and  $Co^{+3}$  ions along with Fe<sup>+3</sup>ions. All these compounds have M-structure through the site distribution changes. No changes occur in the charge distribution but the site distribution is change due to strichiometric changes.

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