Magnetic and Resonance Study of Double doped M-type hexaferrite $\text{Ba}_{1-x}(\text{Cu}_x\text{Al}_x\text{Fe}_{12-x})\text{O}_{19}$

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ABSTRACT: Nanoparticles of $\text{Ba}_{1-x}(\text{Al}_{x}\text{Cu}_{x}\text{Fe}_{12-x})\text{O}_{19}$ (x varies from 0 to 1) were prepared by solid-state reaction route method. The structural and morphological characterization of samples has been investigated by X-ray diffraction and TEM. Electron Spin Resonance (ESR), Mossbauer, VSM techniques were used to study the magnetic properties of the compounds. Change in the magnetic state of nanoparticles due to doping was also examined. Mossbauer studies suggest that the sample is ferromagnetic when x equals 0 and completely paramagnetic when x equals 1. ESR studies reveal splitting of resonance lines at certain concentrations (x) of Cu/Al. VSM and ESR studies reveal the destruction of collective magnetism when doping is increased.

Keywords: Doped barium ferrite; Eelectron spin resonance; Mossbauer spectroscopy; Magnetic properties

INTRODUCTION

Hexagonal ferrites are one of the most important materials for modern industrial world. These materials have been employed in the manufacture of simple permanent magnets, data storage and microwave devices$^1$. Ba – hexaferrite, also known as M-type [(Ba, Sr)Fe$_{12}$O$_{19}$], has attracted particular attention because of its significant properties like magnetic magnetocrystalline anisotropy, high coercive field, and Curie temperature$^2$. Being M-type hexaferrite, it has a complex structure where Fe$^{3+}$ ions have nonequivalent positions and are separated by O$^2-$ ions. The spins of opposing iron ions are linked by interactions which take place via the intermediate oxygen atom, and this is termed as super-exchange$^2$. In general, the magnitude of exchange depends on distance in Metal-Oxygen-Metal chain angle. The smallest interaction occurs at 90$^\circ$ and maximum exchange takes place at 180$^\circ$. So the distortions in iron environments (oxygen polyhedrons) result in super-exchange and hence consequently change the magnetic properties. Thus the behavior of hexaferrites is similar to substituted manganese where magnetic ions of manganese interact with each other by double exchange$^3$.

Doping is an effective method to change properties. Ba$^{2+}$ (1.34 Å) and Fe$^{3+}$ (0.64 Å) ions have different ionic radii. Accordingly, substitutions in different positions cause a different response. Also, Fe$^{3+}$ ions of barium hexaferrites are distributed over five different sub-lattices$^9$. There are many reports on substituted barium hexaferrites by different elements with different oxidation states and ionic radii$^{4,10}$. In a number of studies, it is observed that the substitution of Al$^{3+}$ in the Fe$^{3+}$-site leads to an increase in coercive force ($H_C$)$^{4,11,12}$ and change in magnetic properties. Aluminum ions replace the iron ions and reduce the volume of the unit cell since ionic radius of Al$^{3+}$ (0.54 Å) is less than that of Fe$^{3+}$ (0.64 Å). This leads to a distortion of the oxygen polyhedrons and as a consequence, the super-exchange interaction is affected. The partial substitution of Fe$^{3+}$ by Al$^{3+}$ in barium hexaferrite has been studied by varying the concentration of x (0 ≤ x ≤ 0.4)$^5$. These results suggest that saturation magnetization ($M_S$) reduces and similarly coercive field growth increases with increase of aluminum concentration. Similar, results are also reported by others$^{15}$. The increase of $H_C$ was established when concentration of Al is increased from 2.8 kOe at x = 0 to 4.7 kOe at x = 2. The change of magnetic properties with small and equal amount of either Al or In in Ba ferrite has been studied$^4$. These studies reveal that $H_C$ is 14 times greater in indium doped ferrite than that in Al doped Ba ferrite. Further the total magnetic moment of samples is reduced from 20 μ$_B$ (for pure Ba ferrite) to 19 and 16.7 μ$_B$ for Al and In doped Ba ferrites. The authors reported that in In doped sample, distortion occurs in crystal structure due to the replacement of iron by indium ions (In$^{3+}$ = 0.79 Å) which have greater ionic radii thus creating frustrated magnetic structure in the sample.

Copper doped hexagonal barium ferrite [BaFe$_{12-x}$Cu$_x$O$_{19}$] was examined$^{14}$. Magnetic properties were investigated and found that when x = 0.03, the low coercive change (∼ 0.64 %) and saturation magnetization increase sharply$^{15}$. Further increase of Cu doping increases saturation magnetization, Curie temperature and coercivity. Hexaferrite properties are sensitive to the method adopted in the synthesis$^2$. Since Ba ion has large ionic radius, doping by ion having more ionic radius (∼ 1 Å) such as Ce$^{3+}$, Pr$^{3+}$, La$^{3+}$ etc.$^{16,17}$ can replace...
Ba in hexaferrite. Double doping in hexaferrite causes unexpected structural and magnetic properties\(^5,6,18,19\). So far, dual substitution of ions with large and small ionic radii of Al and Cu ions in hexaferrite was not reported. Therefore, we propose to synthesize \(\text{Ba}_1-x(\text{Cu}_x\text{Al}_x\text{Fe}_{12-x})\text{O}_{19}\) ferrite via solid state route method by varying \(x\) from 0.00 to 1.00 in steps of 0.25 to study the influence of concentration of Cu/Al on the magnetic properties of the compound.

**EXPERIMENTAL**

Barium hexagonal ferrite \(\text{Ba}_1-x(\text{Cu}_x\text{Al}_x\text{Fe}_{12-x})\text{O}_{19}\) (\(x = 0.0, 0.25, 0.5, 0.75, 1.0\)) samples were synthesized by employing solid state route method. Generally in synthetic ferrite systems, Ba composition remains constant and is at its regular lattice site while Fe has been replaced by Cu/Al. But in the present synthesis, compositions of Ba, Cu, Al and Fe are varied.

X-ray diffractometry was performed by Philips X-ray diffractometer using Cu-K\(_\alpha\) radiation (\(\lambda = 1.54060\) Å) at room temperature. Information about morphology and particle size were obtained on Philips CM 200 transmission electron microscope operating at 200 kV and resolution 0.23 nm. ESR spectra of all the samples were recorded from 150 K to 300 K on Carl JES X3 series ESR spectrometer operating at X band frequency (\(\nu = 9.198\) GHz). \(^{57}\)Fe and \(^{55}\)Fe Mossbauer absorption spectra were recorded in transmission geometry at room temperature using a multichannel analyzer with a drive in constant acceleration mode; \(^{57}\)Co(Rh) source with initial activity of 20 mCi was used. Magnetic properties of the compounds were measured using vibration sample magnetometer (VSM). The magnetization loops (M versus H) of all samples were carried out at 5, 150, 200, 255 and 300 K using VSM in a fields up to 80 kOe.

**RESULT AND DISCUSSION**

**Micro and crystal structure**

Figure 1 presents the X-ray diffraction patterns of all barium ferrite samples of \(\text{Ba}_1-x(\text{Cu}_x\text{Al}_x\text{Fe}_{12-x})\text{O}_{19}\). The X-ray diffraction patterns of all the species are similar with very sharp peaks which indicate that all have hexagonal crystal structures. No additional phases and impurities are found.

Using X-ray data, lattice constants are calculated and are in the range of 5.8827 Å to 5.777 Å and 23.195 Å to 23.00 Å for \(a\) and \(c\) respectively. The lattice constant \(a\) is found to decrease with increase of Cu/Al concentration (\(x\)), the lattice constant \(c\) decreases when \(x\) is increased up to 0.25 and then remains constant at concentration \(x = 0.50, 0.75\) and 1.00. This indicates that the change of the minor axis (\(a\)-axis) is larger than that of major \(c\) axis for the substitution of \(\text{Cu}^{2+}/\text{Al}^{3+}\) ions. This behavior is attributed to the substitution of \(\text{Fe}^{3+}\) ions which are of larger ionic radii (0.064 nm) compared with \(\text{Al}^{3+}\) ions (0.054 nm), and the substitution of \(\text{Ba}^{2+}\) ions which are of larger ionic radius (0.134 nm) compared with \(\text{Al}^{3+}\) ions (0.054 nm) and \(\text{Cu}^{2+}\) ions (0.078 nm). The ionic radii sum of \(\text{Al}^{3+}\) ions and \(\text{Cu}^{2+}\) ions are 0.132 nm and is almost the same as \(\text{Ba}^{2+}\) ionic radius. Thus in addition to normal doping of \(\text{Al}^{3+}\) in \(\text{Fe}^{3+}\), double doping of \(\text{Al}^{3+}\) and \(\text{Cu}^{2+}\) into Ba site takes place. Hence the noticeable value of cell size decreases and resulting low cell constant. Using Debye-Scherrer equation, the particle size is evaluated from the line broadening of the peak (203) which changes from 125 to 100 nm.

Figure 2(a-c) shows TEM images of the un-doped (\(x = 0.00\)), doped (0.50) and maximum doped (\(x = 1.00\)) samples respectively. It is noticed that the hexaferrite particles are well defined with polygonal octahedral, tetrahedral and bipyramidal shapes (shown in Figure 2(a) at \(x = 0.0\)) and are well joined. From nano-chemistry point of view, the adjacent particles which have common crystallographic orientation collide each other leading to coalescence. Adjacent primary particles attach to each other and form aggregates.
replacement of Fe$^{3+}$ ions by Al$^{3+}$ ions. Further replacement of Ba$^{2+}$ ion by both Cu$^{2+}$ and Al$^{3+}$ ions leads to decrease in radii of cations. This results in the decrease of grain size. Hence the lattice constant decreases. This suggests that adjacent particles which do not have common crystallographic orientation do not collide with each other and thus they are appearing as separate particles. Also it is seen that the particle size decreases with increase of Cu/Al concentration. From Figure 2, it is observed that the particle size is drastically affected with increase of Cu/Al composition. The particle size distribution is broader and indicates the polycrystalline nature. The particle size varies from 98 nm, 95 nm, 69 nm, 78 nm and 61 nm when x is increased from x = 0.00 to 1.00 in steps of 0.25. The particle size is large at x = 0.0 and small at x = 1.0. The ratio of particle size is about 1:6 (without doping vs maximum doping).

**Magnetic properties**

Bulk hexaferrite BaFe$_{12}$O$_{19}$ is a classic magnetic material with $T_c = 725$ K and exhibits very high uniaxial anisotropy. When the same compound is in nanoscale form, its magnetic state changes drastically.

Figure 3 shows magnetization (M emu/g) versus temperature (T/K) behavior of Ba$_{1-x}$(Cu$_x$Al$_x$Fe$_{12-x}$)O$_{19}$ (x = 0.0 to 1.0) samples under both Zero-Field-Cooled (ZFC) and Field-Cooled (FC) conditions in an applied magnetic field of 80,000 Oe. ZFC curves were obtained by cooling the sample from 300 K to 5 K without applying an external magnetic field. In the case of FC temperature of the sample after reaching 5 K, an 80,000 Oe magnetic field was applied and the magnetization was recorded as the temperature was increased. From these curves blocking temperature is known. This is the temperature at which branching of ZFC and FC curves merge together or where a maximum in magnetization in the ZFC data is seen.

In the present study nearly temperature independent magnetization behavior have been seen between 300 K to 150 K except for x = 0.50. For this compound raise in the magnetization below 250 K was observed and attributed to paramagnetic phase. Below $T_B$ magnetic susceptibility decreases with decrease of temperature. This behavior is characteristic of super paramagnetism present in the compound.

As can be seen from Figure 3, the un-doped sample (x=0.0) shows a high magnetization when compared to the doped samples; it changes slightly with increase of temperature. A similar trend is observed in doped sample when x = 0.25. In this compound the magnetization decreases by about 40% relative to the pure ferrite. In this compound $T_B$ is noticed as 150K. The magnetization is zero beyond $T \geq 150$ K. When x $\geq$ 0.5, a qualitatively different dependence inherent to super-paramagnetic state is observed. In these compounds $T_B$ is found to be approximately 25 K, 20 K and 10 K for x = 0.50, 0.75 and 1.00 respectively. These results suggest that $T_B$ decreases with the increase of Cu/Al content. At room temperature, no magnetic property is noticed with x = 0.75 and 1.0.

In the present case, when x = 0.75, the magnetization is almost zero at $T \geq 200$ K. Previous reports indicate that substitution of Al even up to x = 4.0 in the compound results in weak magnetization. Using Mossbauer spectroscopy, they explained that Al ions preferentially occupy the octahedral site 4f2, 2a,12k and 4f1 at x $\leq$ 2.0.

The magnetization measurements (M verses H) of all Ba$_{1-x}$(Cu$_x$Al$_x$Fe$_{12-x}$)O$_{19}$ samples are carried out at 5, 100, 200, 255 and 300 K using vibrating sample magnetometer (VSM) in a field of 80000 Oersted. The measured hysteresis loops for Ba$_{1-x}$(Cu$_x$Al$_x$Fe$_{12-x}$)O$_{19}$ samples as a function of the applied magnetic field are shown in Figure 4(a-b). Magnetization is normalized to the maximum value for each curve at 30 kOe for better data comparison. From Figure 4(a), we notice that there are three different cases: For undoped

Figure 4. Ba$_{1-x}$(Cu$_x$Al$_x$Fe$_{12-x}$)O$_{19}$ hysteresis loops for different temperatures.

Figure 3. Temperature dependence of $M(T)$ of substituted Ba-ferrite.

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BaFe_{12}O_{19} sample (x=0.0), M(H) curve shows no hysteresis and its residual magnetization is an indication of superparamagnetic state. Completely doped sample (x = 1.0) M(H) show typical behavior of paramagnetic nature and its magnetization linearly increases with increase of magnetic field. In all other cases, the compound shows combined behavior. Thus, on one hand, the symmetric loop of hysteresis is clearly visible and on the other hand, the magnetization does not reach saturation even at high field. When temperature increases to 300 K, there are no qualitative changes in M – H curves. When x = 0.50, there is a decrease in residual magnetization. Further when x = 0.25, the residual magnetism Hc is 0.3 kOe at 100 K and it decreases to 0.15 kOe at 300 K.

In BaFe_{12}O_{19} nano ferrite system, there are five non-equivalent sub-lattices, of which three are octahedral (2a,12k, and 4f), one tetrahedral (4f), and one trigonal bipyramidal (2b). In the Ba_{1-x}(Cu_{x}Al_{x}Fe_{12-x})O_{19} nano compounds, Fe^{3+} ions are partially replaced by an equal amounts of Cu^{2+} and Al^{3+} ions, similarly Ba^{2+} ions are replaced by both Al^{3+} and Cu^{2+} ions. Now when the diamagnetic Al^{3+} ions are substituted by Fe^{3+} ions, it results in decrease of the magnetic saturation (M_s) value because of the super exchange interaction (Fe^{3+}-O-Fe^{3+}) since the Al^{3+} ions preferentially occupy the octahedral 2a site followed by 12k and 4f sites. Introduction of Cu^{2+} and Al^{3+} into the tetrahedral site does not change the net magnetization of the ferrite compound but replacement of Cu^{2+} ions in octahedral sites resulted in the variation of the net magnetization of the materials. Thus all the magnetic properties are found to decrease with the increase of Cu/Al contents, which is due to the occupation of the doped cations at the octahedral sites (12k and 2a) having a spin of electrons in an upward direction.

Previous reports indicate that substitution of Cu/Al in tetrahedral and octahedral sites in cobalt ferrite lowers the magnetization values. In our compounds, similar trend is noticed suggesting that Cu/Al doping enters both tetrahedral and octahedral sites which results in decreasing of the magnetic properties.

It is well known that the size of nanoparticles has a strong influence on the magnetic properties. In the case of our samples, it has been found that substitution affects the size of nanoparticles. This gives the possibility of partially eliminating the size effect in magnetic properties. For further investigation of the magnetic state, the method of electron spin resonance is used. This powerful method, in some cases, allows us to separate the magnetic phases of magnetic materials including complex oxides.

**Resonance study**

The results of VSM measurement are confirmed by Electron Spin Resonance spectroscopy. ESR spectra recorded for barium ferrite powder Ba_{1-x}(Cu_{x}Al_{x}Fe_{12-x})O_{19} in the range 150 K to 300 K are shown in Figure 5. Fe^{3+}, a d^{5} ion with S_{O}=5/2 spin state splits into three Kramer’s doublets, separated by nearly 1.0 cm⁻¹. In this case, only the lowest doublet is populated and one can observe resonances at g = 9.0. On the other hand, if the middle Kramer’s doublet is also populated, a g value of 4.29 is expected. If third doublet is also populated, g values ranging from 0.3 to 4.29 can be seen.

ESR spectra recorded for nano powder (BaFe_{12}O_{19}) in the range 150 K to 300 K is shown in Figure 5(a). It is seen that only a resonant signal with strongly symmetric resonance at all temperatures with same g value of 2.12, varying line intensity and width is noticed. It is attributed to the ions Fe(III) distributed in octahedral coordinate sites. It is ascribed to the ferrimagnetic resonance exhibited by Fe(III) cations in the ferrite at different interstitial sites that couple anti-ferromagnetically due to a super exchange interaction. The line-width of EPR signal may be due to magnetic dipole interactions among the particles or super exchange interactions between the magnetic ions through the oxygen ions. If dominant dipole interactions are present in the compound large line-width and g-factor occurs, whereas super exchange interactions are dominant small-line-width and lower g-values are observed. Occurrence of narrow peak-to-peak line-width at room temperature in the sample may be due to dominant super exchange interaction. As Cu/Al doping progresses, line splitting is observed. The similar results were observed by the authors. EPR spectra of nano (Cu_{0.75}Ba_{0.25})(Al_{0.75}Fe_{11.25})O_{19} recorded in the range 150 to 300 K is shown in Figure 5(b). The spectrum is characteristic of Fe(III) and Cu(II). The calculated g values are 4.0, 2.25 are ascribed to Fe(III) and Cu(II) respectively. In particular at 250 K the observed g values are 2.74 and 2.18 which may be due to the superimposition of Fe(III) resonant signal with g ≈ 2.18.

EPR spectra of nano (Cu_{x}Ba_{1-x})(Al_{x}Fe_{12-x})O_{19} recorded in the range of 150 K - 300 K is shown in Figure 5(c). The spectrum is resolved with characteristic of Fe(III) and Cu(II) signals. At 250 K, the observed resonant signals give g values 5.67, 3.65 and 2.25 which are ascribed to Fe(III) and Cu(II) respectively. This suggests that as Cu/Al concentration increases the nano compound reducing its ferromagnetic character and slowly converting into super paramagnetic nature. Further we notice that the Cu is entering into tetragonal distortion.

EPR spectrum of completely doped Cu(AlFe_{12})O_{19} nano compound recorded in the temperature range of 150 K to 300 K is shown in Figure 5(d). The spectrum is well resolved with five resonant signals ranging from low to high field. These five resonant signals indicate that the compound may have five Fe(III) sites. These signals are due to Fe^{3+} and Cu^{2+}. Thus the compound is super-paramagnetic in nature. This is explained as follows: It is assumed that Ba^{2+} is totally replaced by Fe^{3+} and only one Fe^{3+} is replaced by diamagnetic Al^{3+} ions in the unit cell. Thus paramagnetic behavior of Cu^{2+} is dominated. From the spectrum, the g values obtained for Cu^{2+} are g_{11} = 2.70 and g_{⊥} = 2.27 which indicate that Cu^{2+} is in tetragonal distortion. Tetragonal cupric complexes with D_{4h} generally have g_{11} (corresponding to the magnetic field along the Z axis of the complex) > g_{⊥} > 2.04 have the ground state d_{x^2-y^2}. In the present case g_{11} ≥ g_{⊥} > 2.00 (2.70 > 2.27 > 2.0036) and hence the ground state is B_{1g} (d_{x^2-y^2}). Using this data, a new parameter G = (g_{11} - g_{⊥})/(g_{11} - 2) is defined and calculated as g ≈ 2.61. If G value falls in between 3 and 5, the unit cell contains magnetically non-equivalent ions. If G value is less than 3, the exchange coupling among the magnetically non equivalent Cu(II) ions in the unit cell is not very strong. If G is greater than 5, a strong exchange coupling takes place among the magnetically non equivalent Cu(II) ions in the unit cell. In the present case G is less than 3.0 (observed values is 2.61), which indicates that the unit cell contains magnetically non-equivalent ions and the coupling is not very strong. As the
ferrite contains a very high percentage of iron, the resonant signals due to Fe$^{2+}$ are also noticed with g values at 9.0, 4.96, 2.04. Thus the EPR results indicate that the CuAlFe$_{11}$O$_{19}$ ferrite nano-crystal is in distorted octahedron structure. The resonant signals observed at higher magnetic field shown in Figure 5(d) have g values of 1.20 and 0.90. Such a low g values may not be due to Cu(II). Since the compound contains Cu, Al, Fe and O$^{2-}$. Generally a free radical O$^{2-}$ gives a resonant signal around DPPH. Also Al is diamagnetic. Hence the resonant values are not due to Al, O$^{2-}$ and Cu(II). Thus these are assigned to Fe(III) which is occupied by third Kramer’s doublet given in the theory earlier. Thus the compound with $x=1.0$ is completely paramagnetic. This result confirmed the magnetic measurements shown in Figure 4. The sample with complete substitution has zero magnetization at 200 K and as the concentration of Cu/Al increases, saturation magnetization ($M_s$) also increases.

From Figure 4(b) ($x=0.5$), the temperature evolution of the magnetic state is observed. At a low temperature, the almost symmetric resonance line is observed. However, as temperature progresses the appearance of an additional signal is observed. At 300 K, two clear lines are observed. This suggests that the samples exhibit a weak ferromagnet-
the quadrupole moment of $^{57}$Fe nucleus and the reduction of interaction between Fe ions and octahedral Al$^{3+}$ ions. The analysis of the data shows that quadrupole splitting (0.66 mm/s) increases with the increase of doping. The quadrupole doublet pattern clearly shows that the sample exhibits paramagnetism. The results obtained are in consistent with the results of the vibrational sample magnetometer and electron paramagnetic resonance spectroscopy.

CONCLUSIONS

Nano-size particles of Cu/Al substituted barium hexaferrite $\text{Ba}_{1-x}(\text{Cu}_{x}\text{Al}_{x}\text{Fe}_{12-x})\text{O}_{19}$, $x = (0.0 – 1.0)$ have been synthesized by solid-state route method. X-ray diffractometry and TEM showed that the size of synthesized nanoparticles is about 100 nm and the doping by Cu/Al leads to a large decrease in their sizes. Magnetic properties are found to decrease with the increase of Cu/Al content which is due to the occupation of the doped cations at the octahedral and tetrahedral sites. The magnetization of the un-doped sample is high and is very slightly dependent on temperature. At the same time, magnetization of maximum-doped ($x = 1.0$) sample is weak. The study of $M(H)$ dependence of $\text{BaFe}_{12}O_{19}$ showed that there is no residual magnetization in the sample and shows almost no hysteresis. When $x = 1.0$ the sample is completely paramagnetic. This is confirmed by the linear dependence of magnetization and EPR studies which has a low value. At intermediate concentrations ($0 < x < 1$), they show a small loop of hysteresis where coercivity ($H_C$) was up to 300 Oe, but they do not reach saturation in large fields. This suggests the coexistence of strongly (ferrimagnetic) and weakly (paramagnetic) behavior which is confirmed by ESR. Mossbauer spectroscopy further confirmed that nanoparticles of un-doped samples have strong-magnetic properties (ferrimagnetic) and completely doped ($x = 1.0$) samples have weak magnetic properties (paramagnetic). Considering the results and literature data, we assume that the double substitution greatly changes the magnetic properties of individual nanoparticles. This leads to the fact that the ensembles of these particles change their behavior (Super PM – PM/ FM – PM).

Notes

The authors declare no competing financial interest.

References


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