Formation of iron nitride thin films with Al and Ti additives

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(Received 10 October 2011; accepted 20 April 2012; published online 21 May 2012)

In this work, we investigate the process of iron nitride (Fe-N) phase formation using 2 at. % Al or 2 at. % Ti as additives. The samples were prepared with a magnetron sputtering technique using different amount of nitrogen during the deposition process. The nitrogen partial pressure ($P_N$) was varied between 0% and 50% (rest argon) and the targets of pure Fe, [Fe + Ti] and [Fe + Al] were sputtered. The addition of small amount of Ti or Al results in improved soft-magnetic properties when sputtered using $P_N \leq 10\%$. It was found that the thermal stability of soft-magnetic phase increases considerably when Al or Ti additives were used. When $P_N$ is increased to 50% non-magnetic Fe-N phases are formed. We found that iron mononitride (FeN) phases (N at. % $\sim$50) are formed with Al or Ti addition at $P_N = 50\%$ whereas in the absence of such addition $\epsilon = \text{Fe}_{x-x} \text{N}$ phases (N at. % $\sim$30) are formed. It was found that the overall nitrogen content can be increased significantly with Al or Ti additions. On the basis of obtained result, we propose a mechanism describing formation of Fe-N phases with Al and Ti additives. © 2012 American Institute of Physics. [http://dx.doi.org/10.1063/1.4718579]

I. INTRODUCTION

Iron nitrides are interesting compounds both from application as well as basic point of view. As nitrogen concentration increased their major phases in the order of increasing nitrogen concentration are $\text{Fe}_{16}\text{N}_2$, $\text{Fe}_2\text{N}$, $\text{Fe}_3\text{N}$, $\text{Fe}_2\text{N}$, $\text{FeN}$, and $\text{Fe}_5\text{N}_4$, of which $\text{Fe}_{16}\text{N}_2$, $\text{Fe}_2\text{N}$, $\text{Fe}_3\text{N}$ are magnetic while $\text{Fe}_2\text{N}$ and $\text{FeN}$ are non-magnetic.1 The $\text{Fe}_3\text{N}_4$ phase has been predicted theoretically but has not yet been evidenced experimentally.2 One of the most studied iron nitride compound is $\text{Fe}_{16}\text{N}_2$ as it posses a giant magnetic moment.3,4 During the last decade the $\text{Fe}_2\text{N}$ has emerged as an alternative to pure iron due to its corrosive resistance and mechanically hard surface.5 Very recently non-magnetic iron nitrides have also been explored as a possible candidate in spintronics applications.6,8

Reactive sputtering of iron with nitrogen results in nanocrystallization and amorphization of the deposited films.9–15 This happens due to incorporation of nitrogen atoms at the interstitial sites resulting in expansion and distortion of bcc-Fe unit cells. This essentially implies that reactive nitrogen sputtering plays an important role in refining the grain sizes. As the grain size decreases below the ferromagnetic exchange length (~20 nm for Fe) the averaging is done on very fine grains and magnetization follows the easy direction of each individual grain. This results in very low coercivity. Further, as the grain size decreases, there is a significant increase in the volume fraction of grain boundaries or interfaces resulting in reasonably high values of saturation magnetization. A combined effect results in good soft-magnetic properties in nanocrystalline or amorphous iron nitrides. However, incorporation of nitrogen in iron also results in loss of magnetization due to formation of non magnetic covalent Fe-N bonds. This leads to magnetic anisotropy and an increase in coercivity. In addition, it has been observed in magnetic Fe-N that Fe and N atoms tend to diffuse out at temperatures as low as 400 K.16–18 This imposes an intrinsic limitation for iron nitrides in succeeding as good soft-magnetic alloys.

Since early 1990s to recent years, various Fe-N thin films were studied by adding a tiny amount of a third element say X, it was observed that using such additives the limitations of binary Fe-N can be reduced to a great extent.19–37 The basic idea behind adding such element X was to chose an element which has more affinity to N than Fe. When a third element $X = \text{Al, Ti, Ta, Zr, etc.}$, is added in an appropriate amount it can be substitutionally dissolved into bcc-Fe unit cell.35 This addition results in increased thermal stability of the system as the binding energy of the system increases because of the heat formation of X-N is less than Fe-N.35 Therefore element X works like a trap for nitrogen diffusing out. Although many third elements have been used in Fe-X-N system, the choice of element X has been rather arbitrary. For example, earlier works were mainly focused on Fe-Ta-N systems,20,22,23,25,27 more recently other elements, e.g., Ti,26,31,35,38 Al,32,34 Zr,30,33 Rh,28 etc., have been used.

In this work we have chosen $X = \text{Al, Ti in Fe-X-N}$, as the atomic radii $a_{\text{Fe}} = 0.156 \text{nm}$ is larger than $a_{\text{Al}} = 0.118 \text{nm}$ but smaller than $a_{\text{Ti}} = 0.176 \text{nm}$.39 Ti addition is expected to expand the units cell of Fe when substitutionally dissolved whereas no such expansion is expected with

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Al addition. By adding similar amount of Ti and Al, i.e., 2 at. % we systematically studied formation of Fe-N phases. The addition of small amount of Al or Ti has been reported to increase the thermal stability of Fe-N.\textsuperscript{26,29,40} It may be noted that though the heat of formation of TiN and AlN is almost similar,\textsuperscript{41} their affinity to N is different.\textsuperscript{25} Therefore the mechanism by which addition of element X affects formation of Fe-N phases is not very clear. In addition, the studies with addition of element X in Fe-X-N system have been mainly focused to magnetic phases where small amount of reactive N is used during sputtering. In the present work, we also investigate the effect of Al or Ti addition in the formation of non-magnetic Fe-N phases. In our approach we have taken extreme care to deposit a series of samples in a single sputtering run so as to ensure identical deposition conditions. The nitrogen partial pressure during sputtering was successively increased from 0% to 50% (rest Ar) with a step of 5%. The structure of deposited phases was studied using x-ray diffraction (XRD) and magnetic moment of the samples was measured using polarized neutron reflectivity (PNR). The magnetization measurements were carried out using a superconducting quantum interference device (SQUID) magnetometer and conversion electron Mössbauer spectroscopy (CEMS).

II. EXPERIMENTAL

Thin film samples were prepared at room temperature using a direct current magnetron sputtering (dc-MS) deposition system. The samples were prepared simultaneously on Si (100) and float glass substrates. A mixture of argon and nitrogen gases at different ratio was used to sputter a target. The total gas flow was kept constant at 20 standard cubic centimeter per minute (sccm). The nitrogen partial pressure defined as $R_N = P_{N_2}/(P_Ar + P_{N_2})$ was varied at 0%, 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, and 50%. A base pressure of $\approx 1 \times 10^{-7}$ mbar was achieved prior to deposition. During the deposition the partial pressure in the chamber was $\approx 4 \times 10^{-3}$ mbar. The targets of pure Fe, and composite targets of [Fe + Ti], and [Fe + Al] were sputtered using a power of 50 W. Before deposition the vacuum chamber was repeatedly flushed with argon and nitrogen gases so as to minimize the possible contamination of trapped gases inside the vacuum chamber. The targets were pre-sputtered at least for 10 min in order to remove possible surface contaminations.

To minimize a variation in deposition parameters for a set of samples, i.e., sputtering of Fe, [Fe + Ti], or [Fe + Al] targets all parameters were kept constant, except $R_N$. On a 99.95% pure target of Fe, small pieces of 99.999% pure Al or 99.99% Ti were pasted using a very small amount of silver epoxy paste on the rear side. Possible contamination due to this silver epoxy was checked by measuring $^{107}$Ag depth profiles using secondary ion mass spectroscopy (SIMS) on epoxy free samples, e.g., Fe-N and that in Fe-Ti-N and Fe-Al-N samples. It was found that the $^{107}$Ag signal in samples prepared without this epoxy was similar to those prepared using silver epoxy. This indicates that contamination due to epoxy is not present throughout the depth of film.

Thin films of different $R_N$ were prepared without exposing the deposition chamber to the atmosphere. The targets were covered with a small slit of size about 10 mm and the substrate was exposed at center of the target. After deposition at a particular $R_N$, the substrate was translated using a computer controlled linear translation stage. Once deposition of a sample at a particular $R_N$ is completed, the gas flows were changed using mass flow controllers and monitored using a residual gas analyzer until a stable gas ratio is obtained. Such a procedure minimized the variation in deposition conditions which might occur otherwise. A total of 10 samples were deposited at various $R_N$ in a series on a glass substrate of 250 mm, and also on Si (100) substrates placed on the glass substrate. The thickness of the films was calibrated with x-ray reflectivity and by using different deposition times it was kept at about 100 nm for all samples prepared in this work.

The composition of deposited Fe-Ti and Fe-Al films (prepared without nitrogen) was determined using energy dispersive x-ray analysis installed in a scanning electron microscope. The obtained composition are Fe$_{98.5}$Ti$_{0.5}$ and Fe$_{98.5}$Al$_{0.5}$. The structural characterizations of the samples were carried out by XRD using a standard x-ray diffractometer (Bruker D8 Advance) equipped with Cu K-\(\bar{\alpha}\) x-rays in \(\theta\)–2\(\theta\) mode. A 1D position sensitive detector based on silicon strips technology (Bruker LynxEye) was used. By using such detector it was possible to obtained XRD data with good statistics.

SIMS measurements were performed to measure relative concentration of nitrogen in the samples using a Hiden SIMS Workstation. An O$_2$ ion beam of energy 5 keV and 400 nA was used to sputter the samples and concentration of $^{14}$N was recorded. The base pressure in the measurement chamber was $1 \times 10^{-9}$ mbar or lower and during measurements it was about $8 \times 10^{-9}$ mbar.

The magnetization measurements were carried out using a quantum design SQUID magnetometer and PNR. The PNR measurements were performed at AMOR (Ref. 42) and NARZISS neutron reflectometers at SINQ/PSI, Switzerland. The CEMS measurements were carried out at room temperature using a $^{57}$Co source embedded in a rhodium matrix. The conversion electrons were detected by a proportional counter having continuous flow of a helium-methane (5% methane balance helium) gas mixture.

III. RESULTS

A. X-ray diffraction

Fig. 1 shows the XRD pattern of Fe-N samples prepared using $R_N = 0\%$–50\%. The obtained structural changes are similar as observed in an earlier work.\textsuperscript{13} Here, samples without Al or Ti additions were essentially prepared as a reference so that effect of alloying elements can be compared precisely. However, as we use a small slit of 10 mm to control over the deposition area there are some interesting features we observed in the XRD patterns shown in Figs. 1(a)–1(c). It may be noted that nitrogen free ($R_N = 0\%$) sample have a bcc-$\alpha$-Fe structure with preferred orientation along (200) direction. Normally pure iron thin films are
expected to align in (110) direction. The preferred orientation along (200) direction was caused by the small slit placed beneath the target. Such slit was necessary to control deposition area so as to deposit ten samples with different $R_{N_{2}}$. This slit basically reduced the number of adatoms depositing on a substrate, i.e., deposition rate. In this situation preferred orientation may change from (110) to (200) direction as the surface energy of (200) plane is lower than that of (110) plane. Similar effects on preferred orientation can also be observed by reducing the sputtering power. At sufficiently high deposition rates, large influx of adatoms may give rise to some strain which results in preferred orientation along (110) direction. Such behaviour with sputtering power has been observed in the case of Ti (Ref. 43) and with substrate biasing in the case of Cr. However, as we observe from Fig. 1(a) with reactive sputtering, the preferred orientation seems to change from (200) to (110) direction. This effect is more prominent when samples were prepared with addition of Al or Ti. As discussed later in Sec. IV, reactive sputtering with nitrogen or with Al or Ti additions, the deposited films cause strain in the Fe and in this case the strain energy may dominate over the surface free energy.

We will first analyze the XRD pattern of samples prepared without Al or Ti additions as shown in Fig. 1(a). Here the line width of the diffracted pattern can be used to calculate the grain size of the diffracting specimen in the direction perpendicular to the plane of the film using Scherrer formula, \[ t = 0.9 \frac{\lambda}{b \cos \theta}, \] where $t$ is the grain size, $b$ is an angular width in terms of 2$\theta$, $\theta$ is the Bragg angle, and $\lambda$ is the wavelength of the radiation used. Although the boundary between nanocrystalline and amorphous phases may not be sharp, the obtained grain sizes (see Table I) may be used to identify the structure as nanocrystalline or amorphous. For pure iron sample, the grain size calculated using (110) reflection is 9.0 nm which decreases slightly to 7.8 nm when sputtered using $R_{N_{2}} = 5 \%$. When $R_{N_{2}}$ is increased to 10\%, the estimated grain size is about 2.4 nm therefore at this partial pressure it may be difficult to label the structure as nanocrystalline or amorphous. It may be noted that a broad peak appearing around 2$\theta = 58.6^{\circ}$ which may be due to formation of $\varepsilon - Fe_{3-x}N$ phase. Above this $R_{N_{2}}$, the peaks are broad enough to identify them as amorphous. This amorphous phase persists up to nitrogen partial pressure of 20\% and for $R_{N_{2}} \geq 25\%$, $\varepsilon - Fe_{3-x}N$ compounds are obtained. Since the

\[ \begin{array}{|c|c|c|c|}
\hline
\text{Grain Size} & \text{Fe-N} & \text{Fe$_{0.98}$Ti$_{0.02}$N} & \text{Fe$_{0.98}$Al$_{0.02}$N} \\
\hline
0 & 9.0 \pm 0.3 & 11.3 \pm 0.3 & 10.3 \pm 0.3 \\
5 & 7.8 \pm 0.2 & 11 \pm 0.3 & 9.6 \pm 0.3 \\
10 & 2.4 \pm 0.3 & 3.8 \pm 0.1 & 4.1 \pm 0.1 \\
15 & \text{Amorphous} & \text{Amorphous} & \text{Amorphous} \\
\hline
\end{array} \]

FIG. 1. X-ray diffraction pattern of iron nitride (Fe-N) thin films (a), Fe$_{0.98}$Al$_{0.02}$N films (b), and Fe$_{0.98}$Ti$_{0.02}$N thin films (c) prepared using nitrogen partial pressure, $R_{N_{2}} = 0\%–50\%$ (rest argon). The lattice parameter obtained from XRD data for samples prepared for $R_{N_{2}} = 0\%$, 5\%, and 10\% are plotted in (d), here solid lines are guide to eye.
heat formation for \( \epsilon - Fe_{3-x}N \) phases is about -40 to -45 kJ mol\(^{-1}\) as compared with neighboring, e.g., Fe\(_2\)N (−12 kJ mol\(^{-1}\)) or Fe\(_2\)N (−34 kJ mol\(^{-1}\)), phases.\(^4\) Therefore from the energetics of binary iron nitrides at room temperature it is expected that \( \epsilon - Fe_{3-x}N \) phase should be readily formed as enough reactive nitrogen is made available which may be the case when \( R_{N_2} \) exceeds 25%.

On the basis of amount of nitrogen partial pressure used, the phases formed can be divided into three ranges: (i) \( R_{N_2} \leq 10\% \), where nanocrystalline Fe-N phases are formed, (ii) \( R_{N_2} = 10\% - 20\% \), where amorphous Fe-N phase are obtained, and (iii) \( R_{N_2} \geq 25\% \), where \( \epsilon - Fe_{3-x}N \) or \( \gamma^\prime - Fe-N \) phases (with addition of Al or Ti) are obtained. We will now discuss the effect of small amount of Al or Ti in formation of iron nitrides in these three \( R_{N_2} \) ranges.

(i) Figs. 1(b) and 1(c) show the XRD pattern of samples with addition of Al and Ti, respectively, for different \( R_{N_2} \). For nitrogen free samples, the average grain size is about 9 nm (see Table I) which increases slightly to 11.3 nm with Ti addition and 10.3 nm with Al addition. At \( R_{N_2} = 5\% \), the grain sizes decreases slightly both with Al or Ti additions. At \( R_{N_2} = 10\% \), the grain sizes decrease appreciably to about 4 nm. The positions of Bragg peaks shift to lower-angle side when \( R_{N_2} \) is increased from 0% to 10%, although the basic structure remains same. The lattice parameter (\( a \)) can be calculated for (110) peak and is plotted in Fig. 1(d). As can be seen here with Al or Ti additions, \( a \) increases more significantly. The reasons for such behaviour will be discussed later in Sec. IV.

(ii) For \( R_{N_2} = 15\% \) and 20\%, amorphous phase are obtained and here no appreciable effect of Al or Ti addition can be seen in the XRD pattern.

(iii) For \( R_{N_2} \) between 25\% and 35\%, the results are similar to Fe-N case where \( \epsilon - Fe_{3-x}N \) compounds are obtained. However at \( R_{N_2} = 40\% \) and 50\%, the results are markedly different. With Ti addition as can be seen in Fig. 1(b) a new phase start appearing for \( R_{N_2} = 40\% \), whereas with Al addition this phase is visible only at \( R_{N_2} = 50\% \). This new phase can be indexed as \( \gamma^\prime - FeN \) (confirmed with CEMS measurements shown later) which is iron mononitride and generally formed when iron targets are sputtered using \( 100\% \) nitrogen.\(^1,11,13,48\) Formation of this high nitrogen phase with Al or Ti addition can be understood with an enhancement of nitrogen incorporation. Although \( \gamma^\prime - FeN \) phase can be obtained when sputtered only with nitrogen as sputtering gas, formation of this phase with a 50\% Ar + 50\% N\(_2\) mixture with Al or Ti addition is interesting as deposition rates will be much higher as compared to the case when sputtering is done using N\(_2\) alone as sputtering gas. The \( \gamma^\prime - FeN \) phase has lowest heat formation (−47 kJ mol\(^{-1}\)), therefore, due to enhancement of nitrogen by adding Ti or Al in Fe results in formation of \( \gamma^\prime - FeN \) phase. These results will be discussed in detail in Sec. IV.

**B. Secondary ion mass emission spectroscopy**

From our XRD results discussed in Sec. III A, we find unusual enhancement in nitrogen concentration for \( R_{N_2} = 40\% \) samples with Ti additives and \( R_{N_2} = 50\% \) with Al additive. In order to confirm this, we performed SIMS measurements in samples prepared using \( R_{N_2} = 40\% \) and 50\%. The obtained depth profile for \( ^{14}N \) is shown in Fig. 2. As can be seen in this figure, there is an enhancement in nitrogen concentration when Al or Ti additives are used while in undoped Fe-N sample the nitrogen concentration remains unaffected. These results correlate-well with XRD results and confirm nitrogen enhancement with Al or Ti additives.

**C. Polarized neutron reflectivity measurements**

PNR is a technique which is able to yield the absolute value of magnetic moment per atom in a magnetic thin film with high accuracy.\(^49\) In contrast to bulk magnetization magnetometer technique (e.g., dc extraction, VSM, or SQUID), no correction due to diamagnetic signal from the substrate has to be applied in PNR. Further, the samples dimensions and mass does not play any role in determination of magnetic moment. During the experiment, polarized neutrons with spin parallel or antiparallel to the direction of magnetization on the sample are reflected-off the surface of the sample at grazing incidence. The measurements were performed with an applied field of 800 Oe, which is sufficient to reach the saturation magnetization in all the samples.

Figure 3 shows PNR pattern of Fe-N, Fe\(_{0.98}\)Ti\(_{0.02}\)-N and Fe\(_{0.98}\)Al\(_{0.02}\)-N samples prepared using different \( R_{N_2} \). As the amount of nitrogen partial pressure is increased, the critical edge in spin down reflectivity (\( R^- \)) shows a shift towards higher q\(_z\) values, and the separation between \( R^+ \) and \( R^- \) reflectivities decreases continuously. For Fe-N samples, \( R^+ \) and \( R^- \) converge for \( R_{N_2} = 35\% \), whereas in case of Fe\(_{0.98}\) Al\(_{0.02}\)-N and Fe\(_{0.98}\)Ti\(_{0.02}\)-N samples the two reflectivities converge at 30\%.
The difference between spin up and down reflectivities is a measure of magnetic moment. The PNR profiles were fitted using a computer program based on Parratt’s formulism. The obtained values of iron magnetic moment ($l_B$) per atom are plotted in Fig. 3(d) as a function of $R_{N_2}$. As can be seen here, the magnetic moment decays as nitrogen is added in the system. Up to $R_{N_2} = 10\%$, this decay is faster when Al or Ti is not added. However, for $R_{N_2} \geq 20\%$, the decay in magnetic moment is faster with addition of Al or Ti. This result can be understood in terms of formation of Fe-N bonds which are readily formed when additives Al or Ti are not added. As discussed later in Sec. IV, with Al or Ti addition nitrogen essentially interacts with Al or Ti and therefore the magnetic moment of Fe does not decay as fast as it will be when Al or Ti is not added. When $R_{N_2}$ is further increased, the total amount of nitrogen that can be added into the system is much exceeds due to interaction of Al or Ti with nitrogen. This leads to a faster loss of magnetization in samples prepared with Al or Ti addition for $R_{N_2} \geq 15\%$. When $R_{N_2}$ exceeds 30$, the spin up and down reflectivities remain identical indicating that samples are not ferromagnetic.

D. Magnetization measurements

The magnetization measurements on different samples were carried out using a SQUID magnetometer. Fig. 4 shows normalized M-H loops for Fe-N, Fe$_{0.98}$Al$_{0.02}$-N and Fe$_{0.98}$Ti$_{0.02}$-N samples prepared using $R_{N_2} = 0$, 5, and 10. The obtained values of coercivity ($H_c$) are plotted in Fig. 4(d). For nitrogen free samples, the value of $H_c$ for Fe thin film is about 30 Oe, which decreases to about 17 Oe, when Al or Ti added. When sputtered using 5$\%$ nitrogen, $H_c$ decreases to about 10 Oe. It may be noted that when sputtered using 5$\%$ nitrogen the average grain decreases (see Table I). A reduction in grain size below ferromagnetic exchange length allows exchange coupling between the neighboring grains and results in a reduced effective anisotropy. The results obtained for the samples which were prepared using $R_{N_2} = 10\%$ are very interesting. For the sample prepared without Al or Ti additions, $H_c$ increases significantly to 170 Oe whereas the samples prepared using Al or Ti it further decreases to about 5 Oe. As can be seen from Fig. 3(d), the magnetic moment decreases as nitrogen is added in the system. Up to $R_{N_2} = 10\%$, this decay is faster when Al or Ti is not added. However, for $R_{N_2} \geq 20\%$, the decay in magnetic moment is faster with addition of Al or Ti. This result can be understood in terms of formation of Fe-N bonds which are readily formed when additives Al or Ti are not added. As discussed later in Sec. IV, with Al or Ti addition nitrogen essentially interacts with Al or Ti and therefore the magnetic moment of Fe does not decay as fast as it will be when Al or Ti is not added. When $R_{N_2}$ is further increased, the total amount of nitrogen that can be added into the system is much exceeds due to interaction of Al or Ti with nitrogen. This leads to a faster loss of magnetization in samples prepared with Al or Ti addition for $R_{N_2} \geq 15\%$. When $R_{N_2}$ exceeds 30$, the spin up and down reflectivities remain identical indicating that samples are not ferromagnetic.

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decreases rather slowly with Al or Ti additions. Therefore with Al or Ti additions, soft-magnetic properties can be improved. The obtained results can be understood in terms of structural changes caused by reactive sputtering. When Fe was sputtered with 10% nitrogen, formation of non-magnetic $\epsilon$-Fe-N covalent bonds may introduce large anisotropy in the system as reported in literature.\textsuperscript{13} Whereas in presence of Al or Ti alloying elements, nitrogen predominantly interacts with these alloying elements and due to formation of finer grain nanocrystalline structure, the coercivity decreases further.

E. Thermal stability of soft-magnetic thin films

From our XRD, PNR, and magnetization measurements, we find that nanocrystalline soft-magnetic phases are obtained when sputtered at $R_{N_2} = 10\%$ using Al or Ti additives. It has been observed in the literature that addition of impurity element $X$ leads to thermally stable Fe-N phases\textsuperscript{29} and therefore it will be interesting to observe the effect of Al and Ti additives on the thermal stability. We annealed three samples prepared without any addition and with Al and Ti additions in a vacuum furnace at different temperatures between 423 K and 573 K with a step of 50 K. A background pressure of typically $5 \times 10^{-7}$ mbar was achieved prior to thermal annealing and these three samples were annealed together for 2 h at each temperature. The annealing of the samples was done isochronaly and x-ray diffraction patterns were recorded after each annealing treatment. Fig. 5 shows XRD pattern of samples in the as-deposited state and after annealing at different temperatures. The structure of the as-deposited samples was already presented in Sec. III A and here the effect of thermal annealing will be discussed. As can be seen from this figure, the sample prepared without any additive start showing additional peaks around $2\theta = 47.7^\circ$ and $65^\circ$ after annealing at 423 K which becomes more pronounced as the annealing temperature is increased. The phase formed after annealing is $\gamma$-Fe$_2$N. However when Al or Ti is added, we find that the XRD pattern remains almost similar to as-deposited samples even after annealing at 573 K which clearly shows that the thermal stability improves significantly with addition of Al or Ti. In a recent study it was observed that in an amorphous Fe$_{85}$N$_{15}$ alloy, N diffusion is orders of magnitude higher as compared to Fe diffusivity.\textsuperscript{18} Therefore it is expected that N atoms will be diffusing rapidly involving jumps through the vacant interstitial sites in the Fe network. The addition of Al or Ti elements works like a trap for these N atoms to diffuse and therefore they remain within their respective positions leading to enhanced thermal stability. Such mechanism has been discussed in the literature.\textsuperscript{30} From observed results we find the Al or Ti addition provides iron nitrides with excellent thermal stability.

F. Conversion electron Mössbauer spectroscopy measurements

From XRD measurements we observed that at the highest nitrogen partial pressure used in this work, new phases of Fe-N are obtained with Al or Ti addition. Therefore in order to understand the phase formed with Al or Ti addition, we performed CEMS measurements in the samples prepared using $R_{N_2} = 50\%$. The measurements were first performed with velocity of the drive between $\pm 8.5$ mm s$^{-1}$, which is sufficient to cover all resonance lines of magnetic Fe. Fig. 6 shows the CEMS pattern corresponding to Fe–N, Fe$_{0.98}$Al$_{0.02}$–N and Fe$_{0.98}$Ti$_{0.02}$–N samples prepared with 50 at.\% of $R_{N_2}$. The CEMS pattern of the Fe-N samples was found to have asymmetric doublets and no magnetic lines can be seen. In order to get precise information about the samples, we prepared Fe-N and Fe$_{0.98}$Ti$_{0.02}$–N samples using $^{57}$Fe enrichment and the CEMS measurements were performed with a reduced drive velocity between $\pm 2$ mm s$^{-1}$ to get better CEMS data. The inset of Fig. 6 shows CEMS pattern of $^{57}$Fe-N and $^{57}$Fe$_{0.98}$Ti$_{0.02}$–N samples. The patterns with Ti addition are clearly different. From XRD results we find that the Fe-N phase formed has $\epsilon$–Fe$_{3-\nu}$N type structure. Therefore in order to fit the observed symmetric doublets we deconvoluted it into two doublets corresponding to Fe-III

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig5.png}
\caption{X-ray diffraction pattern of Fe-N (a), Fe$_{0.98}$Al$_{0.02}$–N (b), and Fe$_{0.98}$Ti$_{0.02}$–N (c) thin films prepared using $R_{N_2} = 10\%$, in the as-deposited state and after annealing at different temperatures.}
\end{figure}
and Fe-II sites. The fitted parameters are given in Table II. These values match well with reported values for $\epsilon$-Fe$_{3-x}$N. With the obtained relative area (RA) of Fe-III and Fe-II sites, nitrogen concentration can be calculated. As discussed by Schaaf et al., and references therein) the probability ($p$) of occupation at Fe-III and Fe-II sites is given by

$$
p(\text{Fe-III}) = \frac{6}{z} - 2,
$$

$$
p(\text{Fe-II}) = \frac{3}{z} - \frac{6}{z}.
$$

With known relatives of Fe-III (RA1) and Fe-II (RA2) sites as given in Table II, we obtain the value of $z = 2.28 \pm 0.02$. This gives the composition of the sample as $\epsilon$-Fe$_{2.8}$N.

On the other hand, CEMS pattern of samples prepared with Al or Ti addition was different as compared to a sample prepared without such additions. Here it was found that the CEMS pattern is rather more asymmetric. The XRD pattern also revealed that the crystal structure is completely different and correspond to ZnS-type fcc structure. The phase identified was a monoatomic iron nitride having 1:1 atomic ratio Fe and N. As described in the literature, fitting of such patterns can be done considering two singlets. The fitted parameters are given in Table II and match well with the reported values. Here the singlets with almost zero value of isomer shift (IS) corresponds to Fe coordinated tetrahedrally with four N atoms and singlets with higher value of isomer shift corresponds to defects or vacancies. Therefore it can be seen that with Al or Ti addition Fe-N phase having more nitrogen can be obtained.

In an earlier work Liu et al. investigated CEMS pattern of Fe-Al-N thin films prepared using 2 at.% Al in a wide $R_N$ range. In their XRD pattern they also observed a peak around $2\theta \approx 34^\circ$, but they identified it as a “signature of Si$_3$N$_4$.” The CEMS pattern of this phase not presented and most probably they overlooked the formation of iron mononitride phase.

### IV. DISCUSSION

Combining our results obtained from different experimental techniques, we investigate a mechanism leading to formation of iron nitride phases with small addition of Al and Ti. We will first focus on the samples prepared with highest $R_N$, i.e., 50%. Here XRD and CEMS results clearly show that $\epsilon$-Fe$_{2.8}$N having about 30 at. % N is formed when additional element X is not added. Whereas with Al or Ti addition $\gamma$-Fe-N phase having about 50 at. % N is obtained. Clearly with Al or Ti addition more nitrogen can be incorporated in iron nitride even though the amount of reactive nitrogen used during sputtering process remains the same. It can also be seen from XRD pattern that high-N $\gamma$-Fe-N phase already start appearing at $R_N = 40\%$ when Ti is added where with Al addition only $\epsilon$-Fe$_{2.8}$N is obtained. This indicates that in Ti addition is more effective than Al in increasing nitrogen content in Fe-N. This behavior can also be observed in the PNR data where the magnetic moment falls-off more rapidly with Ti addition as compared to Al addition for $R_N \geq 15\%$.

As mentioned before, the atomic radii of Fe is larger than Al but smaller than Ti. Therefore if small amount of Al or Ti is getting substitutionally dissolved in Fe, it is expected that Ti addition should expand the unit cell of Fe while Al addition may not be able to do so. From the XRD measurements performed on samples prepared without reactive nitrogen, we find that the lattice parameter ($a$) with Ti addition is 2.853 Å whereas with Al addition it is 2.849 Å. The experimentally obtained value of $a$ for pure Fe sample is 2.849 Å. Therefore it appears that Ti addition is expanding the unit cell slightly while Al addition is shrinking it slightly. Such a small change has been reported in literature with addition of third elements Al (Ref. 29) or Ti. A schematic representation of this scenario can be understood from Fig. 7. With this argument alone we expect that Ti addition should result in incorporation of more nitrogen atoms due to expansion in

![Graph showing conversion electron Mössbauer spectroscopy measurements on Fe-N](graph)

**TABLE II.** Conversion electron Mössbauer spectroscopy parameters: IS, quadrupole shift (QS), and RA for Fe-N, Fe$_{0.98}$Ti$_{0.02}$N and Fe$_{0.98}$Al$_{0.02}$N samples prepared with $R_N = 50\%$ nitrogen.

| Parameter | Fe-N | Fe$_{0.98}$Ti$_{0.02}$-N | Fe$_{0.98}$Al$_{0.02}$-N |
|-----------|------|------------------------|------------------------|
| IS1 (±0.02) | 0.45 | 0.01 | 0.1 |
| IS2 (±0.02) | 0.38 | 0.64 | 0.66 |
| QS1 (±0.04) | 0.25 | – | – |
| QS2 (±0.04) | 0.50 | – | – |
| RA1 (±3\%) | 63 | 54 | 62 |
| RA2 (±3\%) | 37 | 38 | 46 |
the unit cell as evidenced in earlier works. However formation of nearly equiatomic iron mononitride phase with Al addition cannot be understood with this argument as the unit cell of Fe is actually shrinking with Al addition. In addition, the expansion in unit cell is very small with Ti addition to cause such an increase in N at. %. Therefore, distortion in the unit cell caused by Al or Ti addition may not give rise to such a prodigious changes as observed in $R_N = 50\%$ samples.

The heat of formation of stoichiometric nitrides TiN is $-337 \text{kJ mol}^{-1}$ and that of AlN is $-320 \text{kJ mol}^{-1}$. These values are much larger as compared to FeN which is $-47 \text{kJ mol}^{-1}$. Therefore it is expected that Ti or Al should be attracting more nitrogen as compared to iron. Although the heat of formation for TiN and AlN is almost equal, the affinity of Al or Ti to N is different. In ternary Fe-X-N systems, the affinity between element X and N has been defined with an interaction parameter $\epsilon_{XN}$. The reported values of values $\epsilon_{XN}$ for Ti and Al are ($-0.63$) and ($+0.0025$), respectively. It has been reported in the literature that a lower value of $\epsilon_{XN}$ signifies high nitrogen incorporation. Our result that with Ti addition higher nitrogen incorporation is achieved as compared to Al addition can be understood with this argument.

V. CONCLUSION

In the present work, we have systematically studied formation of magnetic and non-magnetic phases of iron nitride using Al and Ti as additives. By carefully choosing these additives which either have a larger or smaller size than Fe, we have studied the process by which such additives result in enhanced nitrogen content in iron nitrides. An enhancement in nitrogen results in excellent soft-magnetic properties in magnetic iron nitrides. Here it was also seen that the thermal stability of the soft-magnetic phase improves considerably when Al or Ti additives are added. On the other hand in non-magnetic iron nitride, equiatomic iron mononitride phases can be deposited using Al and Ti which are not possible otherwise. By studying the phase formation using different experimental methods, we investigate a mechanism leading to formation of iron nitride phases in presence of Al and Ti additives. It was found that rather than the atomic size or heat of formation for nitrides of additives, their affinity to nitrogen is deterministic in enhancement of nitrogen in iron nitride phases.

ACKNOWLEDGMENTS

We acknowledge DST, Govt. of India for providing financial support to carry out NR experiments under its scheme “Utilization of International Synchrotron Radiation and Neutron Scattering facilities.” A part of this work was performed at the Swiss Spallation Neutron Source, Paul Scherrer Institute, Villigen, Switzerland. We would like to thank Dr. A. Banerjee for SQUID measurements and Dr. V. R. Reddy for CEMS measurements. We are thankful to Dr. P. Chaddah for continuous support and encouragement.

FIG. 7. Schematic illustration of bcc-Fe unit cell with Al and Ti additives getting substitutionally dissolved. The Fe-Fe, Fe-Al, and Fe-Ti distances are obtained from XRD measurements.

1N. Kordonina, A. Yurovskikh, and A. Kolpakov, Metr. Sci. Heat Treat. 52, 457 (2011).
2Y. W. Ching, Y. N. Xu, and P. Rulis, Appl. Phys. Lett. 80, 2904 (2002). T. K. Kim and M. Takahashi, Appl. Phys. Lett. 20, 492 (1972).
3M. Komuro, Y. Kozono, M. Hanazono, and Y. Sugita, J. Appl. Phys. 67, 5126 (1990).
4E. L. P. y. Blancl, J. Desimoni, N. E. Christensen, H. Emmerich, and S. Cottencier, Phys. Status Solidi B 246, 909 (2009).
5C. Navio, J. Alvarez, M. J. Capitan, F. Yndurain, and R. Miranda, Phys. Rev. B 78, 155417 (2008).
6L. de Witt, T. Weber, J. S. Custer, and F. W. Saris, Phys. Rev. Lett. 72, 3835 (1994).
7M. Gupta, A. Tayal, A. Gupta, R. Gupta, J. Stahn, M. Horisberger, and A. Widera, J. Appl. Phys. 110, 123518 (2011).
8N. Hellgren, M. P. Johansson, E. Broitman, L. Hultman, and J.-E. Sundgren, Phys. Rev. 59, 5162 (1999).
9D. Babonneau, M. Jaouen, M.-F. Denanot, P. Guérin, and F. Petroff, Appl. Phys. Lett. 82, 30565 (2003).
10M. Gupta, A. Gupta, S. Rajagopal, and A. K. Tyagi, Phys. Rev. B 65, 214204 (2002).
11A. Gupta, M. Gupta, B. A. Dasanmacharya, S. Kikuta, Y. Yoda, and M. Seto, J. Phys. Soc. Jpn. 73, 423 (2004).
12R. Gupta and M. Gupta, Phys. Rev. B 72, 024202 (2005).
13R. Dubey, A. Gupta, and J. C. Pivin, Phys. Rev. B 74, 214110 (2006).
14R. Gupta, A. Gupta, W. Leitenberger, and R. Rüffer, Phys. Rev. B 85, 075401 (2012).
15C. Navio, M. J. Capitán, J. Álvarez, R. Miranda, and F. Yndurain, New J. Phys. 12, 073004 (2010).
16J. L. Menendez, G. Armelles, A. Cebollada, F. Briones, F. Peiro, F. Guell, A. Cornet, M. L. F. Guibieda, J. Gutierrez, and C. Meyer, J. Appl. Phys. 89, 6314 (2001).
17S. Chakravarty, M. Gupta, A. Gupta, S. Rajagopal, A. Balamurugan, A. Tyagi, U. Deshpande, M. Horisberger, and T. Gubler, Acta Mater. 57, 1263 (2009).
18N. Hasegawa and M. Saito, J. Magn. Soc. Jpn. 14, 313 (1990).
19N. Ishiwata, C. Wakabayashi, and H. Urai, J. Appl. Phys. 69, 5616 (1991).
20H. Ono, M. Fujinaga, T. Yonemoto, and T. Miyagawa, J. Appl. Phys. 73, 2438 (1993).
21Y. Takeshima, N. Ishiwata, T. Korenari, and H. Urai, J. Appl. Phys. 73, 6576 (1993).
22G. Qiu and J. A. Barnard, J. Appl. Phys. 75, 6934 (1994).
23F. Roozeboom and F. W. A. Dume, J. Appl. Phys. 77, 5293 (1995).
24B. Viala, M. K. Minor, and J. A. Barnard, J. Appl. Phys. 80, 3941 (1996).
25H. Y. Wang, E. Y. Jiang, H. L. Bai, P. Wu, Y. Wang, and F. F. Gong, J. Phys.: Condens. Mater. 9, 8443 (1997).
26L. Varga, H. Jiang, T. J. Klemmer, W. D. Doyle, and E. A. Payzant, J. Appl. Phys. 83, 5955 (1998).
27Y. Chen, C. Qian, C.-Y. Hung, and M. Miller, J. Appl. Phys. 87, 5864 (2005).
28Y.-K. Liu and M. H. Kryder, Appl. Phys. Lett. 77, 426 (2000).
29A. Chezan, C. Craus, N. Chechenin, L. Niesen, and D. Boerma, Phys. Status Solidi A 189, 833–836 (2002).
30J. Rantschler, Y. Ding, S.-C. Byeon, and C. Alexander, J. Appl. Phys. 93, 6671 (2003).
31Y.-K. Liu, M. H. Kryder, D. H. Ryan, and Z. Altounian, J. Appl. Phys. 93, 6471 (2003).
32N. G. Chechenin, A. van Veen, H. Schut, A. R. Chezan, D. O. Boerma, T. Vystavel, and J. T. M. D. Hosson, J. Phys.: Condens. Mater. 15, 7663 (2003).
34 A. Kamzin, F. Wei, Z. Yang, and S. Kamzin, Tech. Phys. Lett. 31, 461 (2005).
35 J. Das, S. S. Kalarickal, K.-S. Kim, and C. E. Patton, Phys. Rev. B 75, 094435 (2007).
36 S. S. Kalarickal, P. Krivosik, J. Das, K. S. Kim, and C. E. Patton, Phys. Rev. B 77, 054427 (2008).
37 F. Xu, S. Li, and C. K. Ong, J. Appl. Phys. 109, 07D322 (2011).
38 Y. Ding and C. Alexander, J. Appl. Phys. 91, 7833 (2002).
39 E. Clementi, D. L. Raimondi, and W. P. Reinhardt, J. Chem. Phys. 47, 1300 (1967), ISSN 00219606.
40 Y. Ma, X. Li, T. Xie, F. Wei, and Z. Yang, Mater. Sci. Eng., B 103, 233 (2003).
41 M. Kopcewicz, J. Jagielski, G. Gawlik, and A. Grabias, J. Appl. Phys. 78, 1312 (1995).
42 M. Gupta, T. Gutberlet, J. Stahn, P. Keller, and D. Clemens, Pramana, J. Phys. 63, 57 (2004).
43 Y. C. Feng, D. E. Laughlin, and D. N. Lambeth, J. Appl. Phys. 76, 7311 (1994).
44 B. D. Cullity, Elements of X-Ray Diffraction (Addison-Wesley, MA, 1978).
45 U. Holzwarth and N. Gibson, Nat. Nanotechnol. 6, 534 (2011).
46 F. Tessier, A. Navrotsky, R. Niewa, A. Leineweber, H. Jacobs, S. Kikawa, M. Takahashi, F. Kanamara, and F. J. DiSalvo, Solid State Sci. 2, 457 (2000).
47 M. Gupta, A. Tayal, A. Gupta, V. R. Reddy, M. Horisberger, and J. Stahn, J. Alloys Compd. 509, 8283 (2011).
48 S. J. Blandell and J. A. C. Bland, Phys. Rev. B 46, 3391 (1992).
49 Y. C. Feng, D. E. Laughlin, and D. N. Lambeth, J. Appl. Phys. 76, 7311 (1994).