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著者	Mohan John Rex, Medwal Rohit, Gupta Surbhi, Gogia Kriti, Vas Joseph Vimal, Gupta Rekha, Deka Angshuman, Rawat Rajdeep Singh, Subramanian Annapoorni, Fukuma Yasuhiro	
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Nonstoichiometric FePt nanoclusters for Heated Dot Magnetic Recording Media

John Rex Mohan^{1#}, Rohit Medwal^{2,#,*}, Surbhi Gupta^{1,2}, Kriti Gogia³, Joseph Vimal Vas², Rekha Gupta⁴, Angshuman Deka¹, Rajdeep Singh Rawat², Subramanian Annapoorni⁴, and Yasuhiro Fukuma¹**

¹School of Computer Science and System Engineering, Kyushu Institute of Technology, Iizuka, Fukuoka 820-8502, Japan. ²Natural Science and Science Education, National Institute of Education, Nanyang Technological University, 637616, Singapore. ³Department of Emergency Medicine, Weill Cornell Medicine, New York, NY 10065, United States

⁴Department of Physics and Astrophysics, University of Delhi, Delhi, 110007, India.

Contributed equally Correspondence: *rohit.medwal@nie.edu.sg, **fukuma@cse.kyutech.ac.jp

Abstract

Heated-dot-magnetic-recording (HDMR) provides a path to increase the areal density of magnetic recording media beyond 4 Tb/in². HDMR based recording media requires ultra-small, non-interacting and thermally stable magnetic dots with high perpendicular anisotropy. We have synthesized non-stoichiometric $Fe_{60}Pt_{40}$ nanoclusters with and without Pt buffer layer on silicon substrates, which shows a reduction in chemical ordering temperature. The $Fe_{60}Pt_{40}$ nanoclusters retain the hard-magnetic phase up to 1023 K with the coercive field of 1.3 Tesla due to the Pt element compensation from the buffer layer. This compensation of Pt was confirmed through XRD investigations where two distinct phases of Fe_3Pt , and $FePt_3$ are observed at elevated annealing temperature. Micromagnetic simulations were performed to understand the effect of magnetic-anisotropy, dipolar interaction and exchange-coupling between the soft magnetic Fe_3Pt and hard magnetic FePt. The results imply that non-stoichiometric $Fe_{60}Pt_{40}$ with Pt buffer layer facilitates low chemical ordering temperature retaining the high perpendicular anisotropy with minimal non-interacting behavior, suitable for HDMR.

Keywords - Information storage, Heated dot magnetic recording, FePt, non-stoichiometric, magnetization reversal.

I. Introduction

Successful implementation of Heated Dot Magnetic Recording (HDMR) technology enables the increase in areal density of magnetic recording media beyond the achieved 4 Tb/in^{21–3}. To realize HDMR, media materials with high magneto-crystalline anisotropy such as FePt and CoPt are best suited due to their high uniaxial anisotropy (K_u) $\geq 10^6$ J m^{-3 2,4}. In film based granular media, $L1_0$ FePtC, (FePt)Ag-C, exchange coupled CoPt/FePt, Fe/FePt, FePt/FeRh, FePt:C/Fe are widely investigated for tuning the switching field distribution and thermal stability ²⁵⁻³¹. The high K_u of FePt and CoPt would also require a high magnetization switching field for write operations, which can be temporarily reduced by localized laser heating. The laser locally heats the isolated bit of the recording media to momentarily reduce the K_u before recording the information⁵ assisted by a near-field-transducer element. Despite these advancements, the critical challenge during this high-temperature writing is the thermal stability of the selected pattern and its magnetic interaction with neighboring bits which requires further optimization^{2,6}.

Self-assembled FePt nanoparticles (NPs) are widely studied for developing the bit-per-grain pattern media⁵. The as-prepared FePt NPs exhibit disordered A1 i.e. face-centered cubic (FCC) phase and require subsequent thermal treatment above ~780 K to transform into $L1_0$ face-centered tetragonal (FCT) phase which display high K_u, required for designing of thermally stable recording media ^{7–9}. Surfactants such as oleylamine and oleic acid are commonly used for synthesising these NPs to ensure the uniform size distribution and reduced magnetic interaction among them, also facilitating self-organization. The organic coating results in a carbonaceous layer around the nanoclusters (NCs) after thermal annealing, which allows the clusters to be self-stabilized against natural oxidation^{3,10-12}. However, high annealing temperature leads to particle sintering, size increase, and agglomeration¹³⁻¹⁵. Alternatively, this indispensable sintering growth

of NPs can be controlled by reducing this ordering temperature of FCT phase by doping of materials such as Ag, NaCl, Pb, Bi, which modifies the grain growth kinetics¹⁶⁻¹⁸. For instance, Pb doping reduces the ordering kinetics and Bi doping allows for a slower diffusion which provides superior surface segregation of dopants even at low temperature annealing. The ordering temperature can also be reduced by varying the relative composition of Fe and Pt in the FePt alloy system¹⁹⁻²². Such a variation in the elemental composition of Fe and Pt in the alloy provides controlled particle size distribution but may affect the magnetic properties including the magnetic anisotropy and the saturation magnetization. The chemical synthesis route provides a superior controllability in achieving a high performance HDMR media. Choosing the composition of metal precursors for the synthesis process requires careful attention ^{23,24}.

In the present work, non-stoichiometric Fe₆₀Pt₄₀ NPs are synthesized using the chemical coreduction method and subsequently, two systems 1) Si/Fe₆₀Pt₄₀ and 2) Si/Pt/Fe₆₀Pt₄₀ are investigated in detail. The effect of annealing temperature on chemical ordering in both Si/Fe₆₀Pt₄₀ and Si/Pt/Fe₆₀Pt₄₀ systems were investigated and compared with equi-atomic FePt composition. To understand the effect of magnetocrystalline anisotropy and dipolar exchange contributions of the Si/Fe₆₀Pt₄₀ and Si/Pt/Fe₆₀Pt₄₀ systems, micromagnetic simulations were utilized using commercial LLG Micromagnetic Simulator TM. The addition of the platinum buffer layer between Si substrate and Fe₆₀Pt₄₀ NPs is expected to lower the chemical ordering temperature as well as a good thermal stability during the high-temperature processing ³².

II. Experimental and characterization methods

A non-stoichiometric amount of Fe(acac)₃ (0.65mmol) and Pt (acac)₂ (0.35 mmol) were mixed in 40 ml phenyl ether with 0.5 mmol Oleic acid, 0.5 mmol Oleylamine and 4 mmol of 1,2 hexadecanediol. This mixture was stirred for 30 min. under inert Argon gas. Thereafter, processing temperature was increased to 373 K and held for next 30 min. to dissociate the metal precursors into metal ions, followed by refluxing at 523 K for another 30 min^{7,33}. The solution was then cooled down to room temperature. Thereafter, 40 ml Ethanol was added followed by 15 min. of stirring. The resultant black precipitate of NPs was washed thoroughly with ethanol and centrifuged at 12000 rpm for 15 min. and finally dispersed in 20 mL Hexane. The dispersed NPs were drop-casted on Si and Si/Pt(20nm) substrates. The deposited NPs assemblies on both substrates were annealed at successive temperatures (T_a) from 673 K to 1023 K. A schematic diagram of the process is illustrated in Scheme 1. The annealing was performed using microprocessor-controlled furnace under forming gas (10% Hydrogen and 90% Argon) for 60 min. with a ramp rate of 5 K per minute. Hereafter, we shall discuss these two systems as (1) Si/Fe₆₀Pt₄₀ and (2) Si/Pt/Fe₆₀Pt₄₀. The first system (Si/Fe₆₀Pt₄₀) was investigated to compare the effect of non-stoichiometry with equi-atomic ratio system, while the second system (Si/Pt/Fe₆₀Pt₄₀) was studied to probe the composition compensation effect arising from self-diffusion of Pt into Fe₆₀Pt₄₀ system at high T_a.



Scheme 1. Synthesis and annealing process of non-stoichiometric Fe₆₀Pt₄₀ nanoparticles.

The as-prepared Si/Fe₆₀Pt₄₀ sample was analysed using Rutherford backscattering (RBS) with a 2 MeV particle beam to estimate the elemental composition. A Tecnai T30 and JEOL ARM 300F GrandARM Transmission Electron Microscopes (TEM) were used to study the size and shape of the NPs dispersed on the carbon-coated copper grid. X-ray diffraction (XRD) data in the 20 range 20° –55° were collected using a Bruker (D8) diffractometer. The software TOPAS was used to perform Rietveld refinement to identify crystallographic phases. Vibrating Sample Magnetometer (VSM) Micro Sense EV9 with a maximum field of ±2.2 T was used for recording the room temperature out-of-plane M-H hysteresis curve. Air-cooled argon-ion laser Raman spectrometer from Renishaw (Invia) was used to record the Raman spectra from 0-3000 cm⁻¹ range to detect carbonaceous layer properties.

III. Results and discussion

Figure 1(i) shows the recorded RBS spectrum of as-prepared Si/Fe₆₀Pt₄₀. The black dots represent the experimental data while the red line represents the simulated spectrum using the Rutherford Universal Manipulation Program simulation code. The elemental composition of the sample is estimated to be Fe, 60% and Pt, 40%. Note that the composition was controlled by optimizing precursors in our previous study¹⁵, where 1:1 ratio of metal precursors resulted in near to stoichiometric composition of Fe₄₆Pt₅₄. The synthesized NPs are subjected to TEM studies and clearly reveal the formation of hexagonal closed packed self-organization of monodispersed Fe₆₀Pt₄₀ NPs as shown in Fig. 1(iii). The observed hexagonal packing is achieved by optimizing the Oleic acid and Oleylamine ratio as surfactants during synthesis, which establishes a protective layer around NPs and prevents them from approaching each other. The average particle size of the NPs is estimated to be ~2 nm using the ImageJ software. Figure 1(iii-vi) show the TEM images

of successively annealed NPs at $T_a = 623$ K, 823K and 1023 K. The average particle size increases up to ~ 11 nm after highest temperature treatment of 1023K. The formation of a carbonaceous overcoat due to the outgassing of hydrogen from surfactants at higher annealing temperature is confirmed by the Raman spectra as shown in Fig. 1(ii). From the Raman spectra, the characteristic peak at 1350 cm⁻¹ confirms the presence of disordered carbon, marked as "D" whereas the peak at 1590 cm⁻¹ is due to graphitic carbon formation and marked as "G". The carbonaceous layer provides two-fold advantages which are controlling the size of NPs and enabling self-stabilization by preventing unwanted natural oxidation.



Figure 1. (i) RBS spectra of as-prepared Fe₆₀Pt₄₀ NPs, (ii) Raman spectra of 1023 K annealed NPs, TEM images of (iii) as-prepared Fe₆₀Pt₄₀ NPs, (iv) TEM image of 673 K annealed NPs, (v) TEM image of 873 K annealed NPs, (vi) TEM image of 1023 K annealed NPs.

The XRD patterns of as-prepared and annealed Si/Fe₆₀Pt₄₀ and Si/Pt/Fe₆₀Pt₄₀ samples at different T_a are shown in Fig. 2(i) and (ii) respectively. The XRD pattern of the as-prepared Si/Fe₆₀Pt₄₀ sample shows a hump centered at $2\theta = 40.17^{\circ}$ corresponding to (111) plane of FePt and attributes to the disordered A1 phase. Annealing at $T_a = 773$ K results in a hump at 47.43° corresponding to (200) reflection. At $T_a = 873$ K, the emergence of superlattice peaks corresponding to (001) and (110) planes with sharp intensities at 24.13° and 33.04° respectively, provides a clear evidence of the evolution of chemical ordering in the sample. The thermal energy induced superlattice peaks are a characteristic feature of chemical ordering attributed to the alternative stacking of Fe and Pt atomic planes in the FePt unit cell. A preferred (001) orientation is observed in Si/Fe₆₀Pt₄₀, which was not observed for the equi-atomic FePt in our earlier report ³³. In addition, a split in the (200) peak into (200) and (002) peaks at $2\theta = 47.36^{\circ}$ and 49.23° respectively, reveals the induced tetragonality in the unit cell because of chemical ordering. The corresponding lattice distortion c/a value of the sample annealed at 873 K is estimated to be ~ 0.964 , indicating the degree of asymmetry in the FePt unit cell and further confirming the phase transformation from cubic to tetragonal phase. On further increase of T_a to 973 K, the relative intensity of (001) with respect to (111) peak decreases and the additional shoulder starts to appear adjacent to (111) peak resulting in asymmetry corresponding to a mixed FePt phase. At $T_a = 1023$ K, the shoulder at (111) peak becomes distinct with enhanced intensity, confirming the presence of an additional phase identified as Fe₃Pt phase (marked by *). The XRD study clearly reveals the temperature dependent structural phase transformation from A1 (Fm3m) to L_{10} (P4/mmm) phase and then to a mixed phase of L_{10} and L_{12} (Pm3m).



Figure 2. XRD patterns of (i) Si/Fe₆₀Pt₄₀ and (ii) Si/Pt/Fe₆₀Pt₄₀ system annealed at T_a from 673 K to 1023 K.

The XRD patterns of Si/Pt/Fe₆₀Pt₄₀ show a similar behavior for as-prepared and $T_a = 673$ K annealed samples. However, after $T_a = 773$ K processing, a clear shift in peak at $2\theta = 41.25^{\circ}$ related to (111) peak and the presence of broaden hump at $2\theta = 46.46^{\circ}$ corresponds to the A1 phase appears. Also, onset of additional (001) and (110) peaks dictate the partial transformation into *L*₁₀ FePt at relatively lower T_a of 773 K with an additional FePt₃. On further increase of T_a to 873 K, characteristics spilt of (200) peak at $2\theta = 47.36^{\circ}$ and 49.23° correspondingly to (200) and (002) respectively confirms the induced tetragonality in the unit cell due to chemical ordering. On the other hand, at $T_a = 973$ K, asymmetry in (111) peak becomes more prominent, providing evidence of the presence of a mixed phase. At $T_a = 1023$ K, this asymmetric peak splits into two distinct peaks at $2\theta = 40.41^{\circ}$ and 41.03° corresponding to (111) of chemically ordered *L*₁₂ FePt₃ phase (indicated by +) and chemically ordered *L*₁₀ FePt phase. Interestingly, for Si/Pt/Fe₆₀Pt₄₀, the preferred orientation along (001) reflection is retained up to $T_a = 1023$ K; hence it is evident that

the Pt buffer layer is promoting the preferred orientation in Si/Pt/Fe₆₀Pt₄₀ in addition to lowering the ordering temperature compared to Si/Fe₆₀Pt₄₀.

Further to quantify respectively the Fe₃Pt and FePt₃ phase, Rietveld fitting of XRD patterns for Si/Fe₆₀Pt₄₀ and Si/Pt/Fe₆₀Pt₄₀ annealed at 1023 K are performed. Figures 3(i) and (ii) show the experimental and Rietveld refined patterns (de-convoluted) of Si/Fe₆₀Pt₄₀. Figure 3 (i) shows the simulated XRD pattern corresponds to the *L*₁₀ FePt phase. All the observed peaks in the XRD patterns are indexed with the P4/mmm space group. Figure 3(ii) shows the XRD pattern corresponding to the *L*₁₂ Fe₃Pt phase and indexed with space group Pm3m. The lattice parameter corresponding to cubic *L*₁₂ Fe₃Pt unit cell was estimated to be 3.82 Å and, for tetragonal *L*₁₀ FePt phase, lattice parameters **a** = 3.85 Å and **c** =3.73 Å. A similar exercise was also performed for the XRD pattern obtained for Si/Pt/Fe₆₀Pt₄₀. In this case, two phases corresponding to FePt and FePt₃ were considered during Rietveld refinement fitting. The experimentally observed and the de-convoluted XRD patterns corresponding to *L*₁₀ FePt₃ was estimated to be a = 3.86 Å and for the *L*₁₀ FePt phase, lattice constant a = 3.86 Å and c = 3.73 Å. Hence, Rietveld analysis clearly shows the evidence of an additional phase corresponding to FePt₃, which may be due to the diffusion of platinum from the buffer layer in the annealing process.



Figure 3. Deconvolution to (i) FCT L_{10} -FePt phase and (ii) FCC L_{12} -Fe₃Pt phase in experimentally obtained XRD pattern of 1023 K annealed Si/Fe₆₀Pt₄₀ system.



Figure 4. Deconvolution to (i) FCT *L*₁₀-FePt and (ii) FCC *L*₁₂-FePt₃ phase in experimentally obtained XRD pattern of 1023 K annealed Si/Pt/Fe₆₀Pt₄₀ system.

Figures. 5(i) and (ii) show the normalized M-H hysteresis behavior of as-prepared and annealed Si/Fe₆₀Pt₄₀ and Si/Pt/Fe₆₀Pt₄₀ samples. All M-H curves are recorded at room temperature in outof-plane geometry. The as-prepared Si/Fe₆₀Pt₄₀ sample exhibits neither a square curve nor a tendency to saturate up to H = 2 T. The sample annealed at $T_a = 673$ K exhibited only small coercive field $\mu_0 H_c$ of 0.03 T. The value of $\mu_0 H_c$ increases from 0.23 T for $T_a = 773$ K to 1.17 T for $T_a = 873$ K. On the other hand, equi-atomic FePt attained lower $\mu_0 H_c \sim 0.58$ T after $T_a = 873$ K, which is reported earlier ¹⁵. Hence, the deviation from equi-atomic stoichiometry promotes the chemical ordering at relatively low $T_a = 773$ K compared to $T_a = 873$ K processing. In Fig. 5(i), the presence of a small shoulder near low magnetic field indicates the existence of soft magnetic Fe₃Pt phase, which is also evident from XRD studies. Further improvement in the coercivity $\mu_0 H_c = 1.33$ T is obtained for $T_a = 973$ K. However, for $T_a = 1023$ K, a decrease in the coercivity to 0.60 T with a pinched curve shows a decoupled behavior of soft and hard magnetic phases. For Si/Pt/Fe₆₀Pt₄₀, the as-prepared sample shows a soft magnetic behavior, similar to as-prepared Si/Fe₆₀Pt₄₀. However, the sample annealed at $T_a = 673$ K, shows higher value of $\mu_0 H_c \sim 0.12$ T as compared to $\mu_0 H_c \sim 0.03$ T for Si/Fe₆₀Pt₄₀, which further increases to 0.23 T after T_a = 773 K process. For $T_a = 873$ K, a sudden rise of $\mu_0 H_c \sim 1.24$ T confirms the formation of hard magnetic FePt L1₀ phase. The shoulder near the low magnetic field is due to the existence of the FePt₃ phase. For $T_a = 973$ K, a slight reduction in the coercivity can be observed. This reduction is due to the exchange spring composite behavior of magnetically hard L_{10} FePt phase and magnetically soft L1₂ FePt₃ phase. However, with further annealing at 1023 K, the sample retains such high value of coercivity and shows a stability in the magnetic hardness. This observed stability is attributed to the diffusion of Pt atoms to Fe rich FePt. The variation of coercivity for the Fe60Pt40 samples and Fe46Pt54¹⁵ is plotted in Fig. 5(iii). The coercivity of Fe46Pt54 monotonically increases with Ta up to 1023 K. However, for Si/Fe₆₀Pt₄₀, the coercivity increases with increasing T_a up to 973 K and later decreases at $T_a = 1023$ K. In the case of Si/Pt/Fe₆₀Pt₄₀, the coercivity increases with an increase of T_a up to 873 K. Thereafter, the coercivity becomes stable even at high temperature treatment, which is one of the essential requirements to improve the intense laser induced change in the magnetic properties while writing information in HAMR and HDMR media. The calculated magnetic anisotropy for different annealing temperatures for the Fe₆₀Pt₄₀ samples is shown in Fig.5(iv). The values are calculated based on the relationship $H_c = 2K_1/\mu_0 M_s$, where $\mu_0 M_s = 1.43$ T for bulk FePt NPs is considered. The presence of hard and soft FePt phases within the NPs have been ignored for this formulation.



Figure 5. Out-of-plane M-H curves of (i) Si/Fe₆₀Pt₄₀ and (ii) Si/Pt/Fe₆₀Pt₄₀ samples at different annealing temperatures. (iii) coercive field as a function of annealing temperature for Si/Fe₆₀Pt₄₀, Si/Pt/Fe₆₀Pt₄₀ and equiatomic composition¹⁵ (with permission) (iv) Calculated magnetic anisotropy of NPs for different annealing temperature.

Micromagnetic simulations were carried out to gain further insight of interparticle distance relationship with the exchange-coupled soft and hard FePt structures. Owing to the strong exchange coupling and interaction between the hard and soft layers, exchange spring magnets have added advantages in reducing the coercive field and maintaining the thermal stability. In order to have a meaningful comparison of experimental results and theoretical simulations, we considered the following assumptions. We modeled Si/Fe₆₀Pt₄₀ and Si/Pt/Fe₆₀Pt₄₀ systems as an array of 4×4 cylinder-shaped bits consists of an exchange-coupled Fe₃Pt/FePt and FePt₃/FePt bilayers, respectively. Comparing XRD patterns of the samples annealed at 973 K and 1023 K, a clear dissociation of Fe₃Pt and FePt₃ phases is observed only in 1023 K annealed samples. This indicates that after $T_a = 973$ K processing, soft layers are strongly exchanged coupled with the hard FePt layer. Moreover, from the M-H curve of 973 K annealed samples, further confirmation of the exchange coupling behaviour can be seen with the absence of kinks or two-phase reversal. Hence the experimental M-H curves of 973K annealed samples was compared with the simulated curves for analyzing the exchange coupling behavior. The total thickness and diameter of each bit was fixed to 12 nm since TEM analysis of the sample annealed at 1023 K exhibited spherical shaped particles with an average particle size between 11-12 nm. Despite not having any direct thickness measurements of each layer, we assumed the soft layer thickness should be within the exchange length of the hard layer in accordance with the reversal mechanism to be the coherent rotation³⁴.

Material	Parameter	Value
FePt	M_s	$1.14 \times 10^{6} \mathrm{A}\mathrm{m}^{-1}$
	A _{ex}	$1.13 \times 10^{-11} \text{J m}^{-1}$
	K _{u1}	$0.8 - 2.6 imes 10^6 \text{ J m}^{-3}$
Fe ₃ Pt	M_s	$1.43 \times 10^{6} \mathrm{A}\mathrm{m}^{-1}$
	A _{ex}	$2.5 \times 10^{-11} \text{J m}^{-1}$
	Kui	0 J m ⁻³
FePt ₃	M_s	$0.56 \times 10^{6} \mathrm{A}\mathrm{m}^{-1}$
	A _{ex}	0 J m^{-1}
	K _{u1}	0 J m ⁻³

Table 1: Parameters used for the Micromagnetic simulation

We performed set of preliminary simulations with fixed 12 nm thickness, varying the soft layer thickness in an increment of 2 nm, and keeping the magnetic anisotropy of $K_{u1} = 1.8 \times 10^{6} \text{ Jm}^{-3}$. By mimicking the experimental M-H curves with the simulation, we first optimized the thickness of soft and hard layers to be 4:8 aspect ratio. Each bit was then modeled as a bilayer of 4 nm soft ferromagnetic Fe₃Pt or FePt₃ bottom layer and 8 nm top hard ferromagnetic FePt layer. The simulation parameters are given in table 1. The discretization size for the simulations was chosen to be 1 nm × 1 nm. The magnetic easy axes of FePt were set to have a minimum orientation along in-plane direction and maximum along out-of-plane direction to imitate the misorientation of easy axes in NPs. The easy axes of Fe₃Pt were set in the in-plane direction with

cubic anisotropy while for FePt₃, the easy axes were adjusted to be equally oriented in all three directions (33% of X, Y, Z) to mimic a room temperature paramagnetic behavior³⁵.



Figure 6. Simulated out-of-plane M-H curves (i) Fe₃Pt/FePt and (ii) FePt₃/FePt for different FePt magnetocrystalline anisotropies (iii) coercivity as a function of FePt anisotropy.

Figures 6(i) and (ii) show the simulated M-H curves for Fe₃Pt/FePt and FePt₃/FePt with different K_{u1} (FePt). The coercivity increases monotonically with the anisotropy. The reversal mechanisms of both systems confirms the strong exchange spring behavior in such a way that the bilayers reverse together. The K_{u1} values obtained from the simulation for the corresponding experimental

coercivity values are 2.4×10^6 J m⁻³ and 1.2×10^6 J m⁻³ for Fe₃Pt/FePt and FePt₃/FePt, respectively. Note that there is a reduction of K_{u1}(FePt) value in FePt₃/FePt compared to Fe₃Pt/FePt. This is due the variation in the *M_s* of the soft layers, for FePt₃/FePt, high *M_s* of both layers result in a higher anisotropy than with FePt₃/FePt layers with much lower *M_s*²⁸. In addition, FePt₃/FePt with higher K_{u1}(FePt) = 1.6×10^6 J m⁻³ shows a small kink. A similar kink can also be seen in the experimental M-H curve obtained for the Si/Pt/Fe₆₀Pt₄₀ sample annealed at Ta = 1023 K in Fig. 5(ii), indicating a decoupled behavior in the bilayer. Although there is no variation in the interlayer exchange coupling, an increase of K_{u1}(FePt) leads to magnetic decoupling. This reduction of anisotropy can be attributed to the formation of graded interface as reported earlier^{34,37} and would require a more detailed study. In either case, the presence of exchange spring behavior in Si/Pt/Fe₆₀Pt₄₀ is considered to be responsible for the reduction of K_{u1} (FePt) which would be beneficial for lowering the writing temperature. In addition, the decoupled behavior with higher K_{u1}(FePt) indicates that even for Si/Pt/Fe₆₀Pt₄₀ sample annealed at T_a = 1023 K, there is no degradation of hard phase at an elevated temperature.



Figure 7. Simulated M-H curves by varying inter-bit distance from 1-10 nm for (i) Fe₃Pt/FePt (ii) FePt₃/FePt. Insets show the magnified image of hysteresis for obtaining coercivity values (iii) simulated coercivity as a function of inter-bit distance.

The values of K_{u1} (FePt) obtained from these coercivity comparisons are utilized further to investigate the effect of dipolar interaction. The dipolar interaction is studied by varying the interbit distance by increasing the total dimension of the system and fixing the 4×4 arrays of 12 nm bits. Figures 7(i) and (ii) respectively show the simulated M-H curves of Fe₃Pt/FePt and FePt₃/FePt as a function of inter-bit distance. Both, FePt₃/FePt and Fe₃Pt/FePt show a nominal inter-bit interaction effect i.e. almost constant coercivity values for large inter-bit distance from 4 to 10 nm and minimal increase in coercivity for 1-4 nm, as can be seen in Fig. 7(iii). The insets in Fig. 7(i) and (ii) show the obtained coercivity values for comparing both layer schemes. In Fig.7 (i), the M-H curves for FePt₃/FePt system with 1 nm interparticle distance shows a clear step that corresponds to the late reversal of corner bits, which is not seen in other simulated hysteresis. The late reversal of corner bits in the case of the inter-bit distance of 1 nm for FePt₃/FePt is due to an increase of the demagnetizing field in the overall system. When the system is modeled with a total thickness of 12 nm and 1 nm inter-bit distance, the system more or less behaves like a continuous film. Also, it implies that at least 2 nm inter-bit distance is necessary for properly modeling dimensions with cylinder-shaped bits. Nevertheless, the minimal increase of coercivity can be attributed to negligible dipolar interaction between the inter-bit separated by 2-4 nm carbonaceous overcoat, beneficial for HDMR application.

Conclusion

Hexagonal closed packed self-organization of non-stoichiometric Fe₆₀Pt₄₀ NPs has been successfully synthesized using the chemical co-reduction method. The structural investigation of Si/Fe₆₀Pt₄₀ and Si/Pt/Fe₆₀Pt₄₀ system as a function of annealing temperature clearly revealed the transformation from chemically disordered phase to chemically ordered phase at low annealing temperature as compared to a system with nearly equi-atomic composition. The addition of the Pt buffer layer stabilizes the *L*1₀ phase due to stoichiometric compensation at higher annealing temperature. Hence, the Pt buffer layer not only promotes the phase transformation at low temperature and but also helps in retaining a high coercivity over a wide range of annealing temperatures. Micromagnetic simulations were performed and compared with Fe₃Pt/FePt and FePt₃/FePt with respect to magnetic anisotropy and dipolar coupling. The results presented in the manuscript demonstrate the low ordering temperature using self-organization of chemically ordered magnetic nanoparticles which is sustained at high temperatures for HDMR based highdensity recording media.

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Conflict of interest declaration

The authors declare that there is no conflict of interest.

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